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# How to navigate without maps: The power of taxon-like navigation in ants 

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#### Abstract

Rodents are said to have two different navigational systems, a map-like locale system and a route-based taxon system consisting of sensorimotor routines such as beaconing and turns at appropriate stimulus conditions (motor routines). Ants on the other hand are not known for map-like navigation, and seem to get by with a repertoire of taxon-like strategies. I review how this repertoire serves ants in making up for the lack of a locale system. Path integration - keeping track of the straight-line distance and direction from the starting point - operates continuously in the background, and can be called upon as necessary, or relied on in habitats in which no useful visual cues are available. Crucial to the power of a taxon-like repertoire is using the full panoramic visual context, both to guide the operation of strategies (context-modulated servomechanisms) and to guide navigation directly. The entire repertoire is backed up by systematic search strategies. I end with some reflections on the power of taxon-like strategies.


Key words: ant, navigation, views, vectors, taxon

## Introduction

Echoing philosopher Immanuel Kant, O'Keefe and Nadel wrote at the end of the first chapter of their influential (1978) book:
there must be a pre-existent spatial framework in order for organisms to experience the world coherently. Objects could not be identified, nor localised, nor even seen as extended in the absence of this framework. (p. 59).

They go on to argue for map-like representations in rats, called the locale system. This quote can be taken to state the functional advantages of having a locale system coding unitary, absolute space. Objects and their locations may be placed in such a unitary overarching framework. The spatial extent of objects may also be coded. Objects may be identified by spatio-temporal continuity. The overarching frame-

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work reduces the problem of the re-identification of objects, or perceptual aliasing (Cheng, 2008, Nehmzov, 2008). Too many objects in the world, from doors to trees, look similar, so that they need to be distinguished in part by the location on a map at which they are found.

Yet among insects whose navigation has been much studied, the existence of a map-like representation is debated in the case of honeybees (Apis mellifera; Cheng, in press; Cruse \& Wehner, 2011; Menzel et al., 2005), or doubted in the case of ants (T. S. Collett \& M. Collett, 2002; Wehner, Boyer, Loertscher, Sommer, \& Menzi, 2006). Navigational problems have many common elements. The geometric laws governing space over a local region (as opposed to a global scale of travel) are the same for all animals. For instance, in local space, three objects define a triangle whose angles add up to $180^{\circ}$. Trigonometric relations are useful for making spatial inferences based on map-like representations. How could navigating ants get by without any map-like representations, perhaps without experiencing the world coherently? Answering this question forms the major theme in this paper. Hopefully, the answers will also shed some light on comparisons of different strategies in different animals for solving common spatial problems.

The answer in short is that insects, and especially ants, rely
much more on strategies akin to those found in the second houn, 1963).
major system of navigation that O'Keefe and Nadel (1978) posited, the taxon system. The taxon system comprises routines and procedures not based on map-like representations but procedures akin to route instructions, to be detailed shortly. The theme that I explore here is that a taxon-like system can be very powerful in the real world with its rich array of cues including trees, bushes, rocks, and perhaps distant hills. Ants seemed to have carved out a taxon-like niche for navigation.

I will first elaborate briefly on locale and taxon systems in rats, without approaching anything like a review. This sets the stage for exhibiting the multitude of taxon-like strategies in a few species of ants (whose navigation has been well studied), with occasional references to honeybees. I end with some speculative reflections regarding navigational strate gies in a comparative context.

## Locale and taxon systems in rats

The taxon system takes on philosophically-empiricist notions of space, being built out of sensory data and motor routines. It includes path integration, called dead-reckoning or internal navigation in O'Keefe and Nadel's (1978) book, guidance, and orientation

Path integration means keeping track of the straight-line distance and direction of travel, so that an agent can compute the vector home at any point during the journey (M.-L. Mittelstaedt \& H. Mittelstaedt, 1980). In rats, a known compass for keeping track of the direction is internal, based on the vestibular system (Etienne \& Jeffery, 2004), although it is possible that external sources such as visual landmarks may also be used.

Guidance is focused on some aspects of the stimulus situ ation surrounding the navigator as a basis for navigation. As ne example, some object is identified from a distance, and he animal heads toward that; beaconing could be another term for this kind of guidance-based navigational behavio (Shettleworth \& Sutton, 2005). Beaconing generally refer to identifying a single visual object (which might consist of one or more physical objects), and heading toward it. But more extended stimuli may also guide animals. Rats might follow well worn tracks (Calhoun, 1963), or honeybee might follow the shore of a lake (von Frisch \& Lindauer 1954)

Orientation, on the other hand, is focused on the motoric aspects of navigation. Turning a particular angle at some point on the journey, and then heading in the turned direc tion would be one example. Together, behaviors based on guidance and orientation can be used to build routes, which might characterize much of rat navigation in the wild (Cal-

Locale navigation, in contrast, can be considered flexible, map-based behavior, to which place cells contribute a hy pothesized neurophysiological basis in O'Keefe and Nadel' (1978) treatment, and in modern updates (Sheynikhovich Chavarriaga, Strösslin, Arleo, \& Gerstner 2009). Place cell in the rat's hippocampus fire when the rat is in a particu lar place in an experimental arena (Jeffery, 2010; O'Keefe 1976; O'Keefe \& Dostrovsky, 1971). The firing rate is in general independent of the direction that the rat is facing, or how the animal got to the place. The key characteristic of the map-based locale system is that places are located in a framework of metric properties and relations, linked to othe places and stimuli (e.g., landmarks) in a map-like fashion. This allows the rat to learn to find an unmarked place, fo example, a fixed place in a swimming pool at which a plat form just under the surface may be found, and to find the arget readily when starting from varying and arbitrary loca tions in the pool (Morris, 1981; Morris, Garrud, Rawlins, \& O’Keefe, 1982).

A quarter of a century after O'Keefe and Nadel's (1978) formulation of the cognitive map as a locale system, a significant "unpacking" was presented (Jacobs \& Schenk, 2003). The unpacking finds two integrated map systems. An volutionarily more ancient bearing map system is based on gradients of various kinds (odor, magnetic cues, wind, etc as well as distant landmarks, providing coarse positiona information and directional compass. A finer but smaller caled system called the sketch map is topographic in na ure, encoding the positions of objects serving as landmark and the navigator's position in the local space. The entire map system is an integrated map formed by combining thes wo kinds of maps, all systems being instantiated in differ ent subsystems of the hippocampal formation. In the integration, the linking of sketch maps with bearing maps is crucial both for identifying the correct sketch map to use and for linking different distant locations

Taxon-like ant navigation
Insects have long been known to use landmarks for navi gation, and much classic work featured ants and bees. For example, von Frisch and Lindauer (1954) trained honeybees (Apis mellifera) to travel along some prominent large-scale feature during an afternoon. The feeder at which the bees were feeding was moved further and further from the hive over the afternoon. The large-scale landmarks might be hore of a lake or a long line of trees (Figure 1). At the end of he day when the bees had stopped foraging, von Frisch and Lindauer closed the hive and moved it to another location a which similar landmark features were found (another shor of a lake or another line of trees) running in a different com


Figure 1. Illustration of an experiment by von Frisch and Lindauer (1954). A. The bees were trained over an afternoon o forage from a feeder beside a line of trees. The feeder was moved further and further along the line of trees ove he course of the day until it reached the location shown. B. A test set up the next day at a different location with a line of trees running in a different compass direction from the line of trees encountered in training. One of the feeders F2) was located along the line of trees. Other feeders (not shown) were placed in other compass directions from the hive. Note the similarity in skyline contour between the test and training situations. Adapted from "Himmel und Erde in Konkurrenz bei der Orientierung der Bienen [Sky and Earth in competition in the orientation of bees]," K. von Frisch and M. Lindauer, 1954, Naturwissenschaften, 41, p. 249. Copyright 1954 by Elsevier. Reprinted with permission.
pass direction. Different feeders were set up before the bees came out foraging the next morning. The bees preferred to follow the landmarks rather than the learned compass direc tion, thus showing that they were utilizing large-scale landmarks to strike a direction of travel.

Ants move on the ground, and on a small enough scale hat experimental landmarks can be provided and manipulated to demonstrate that they use landmarks. Santschi (1913)
reported experiments in the 19th century by John Lubbock showing the use of landmarks in ants (Lasius niger). The ants moved back and forth between their nest and a feede over a circular disk. A candle off to one side was provided as a landmark. Rotation of the circular disk, while leaving the candle in its place, had no notable effects on the ants behavior. Rotating the disk with the candle on it, however changed the course that the ants took, showing that the were using the candle to strike a navigational course. Thi early experiment heralded modern experiments manipula ing such moveable landmarks and showing their effects on nt navigation (Cataglyphis fortis: T. S. Collett, Dillmann Giger \& Wehner 1992. Formica rufa: Durier, Graham, \& Collett, 2003; Gigantiops destructor: Macquart, Garnie Combe, \& Beugnon, 2006; Melophorus bagoti: Wystrach Schwarz, Schultheiss, Beugnon, \& Cheng, 2011)
Santschi himself (1911) showed that ants (of the genus Cataglyphis) used the sun as a source of direction by reflect ing an image of the sun in a mirror. He reported that the ants would reverse their direction of travel upon viewing th sun in the mirror. This has led to a successful program char acterizing the nature of the sky compass in hymenopteran and other insects (Rossel \& Wehner, 1986; Wehner 1994 Wehner \& Müller, 2006)
Among the studied hymenopterans, map-like navigation is not known in ants, and has not been proposed by anyone to my knowledge. A spectacular failure to exhibit map-like behavior was found in the Australian desert ant M. bagoti (Wehner et al., 2006). The ants were trained to forage at a feeder, but a direct route either to the feeder or back home was prevented, so that the ants had to detour and complete a loop as they foraged at the feeder. The ants learned such a loop readily. Wehner et al. then asked whether the traine ants could recognize on their way home a place encountered on their usual outbound route, a route that they had traversed many times on the way to the feeder. Ants on their way home with food were displaced just before reaching their nest to some place on their outbound leg to the feeder. Such ant displayed search behavior and failed to head either directly home or in the reverse direction of their outbound route ap parently not recognizing the place.

A number of well-studied ant species exhibit plenty of tax-on-like navigation, and can accomplish many feats withou resorting to map-like navigation (reviews: Cheng, in press Cheng, Narendra, Sommer, \& Wehner, 2009; T. S. Collett \& Collett, 2002; T. S. Collett, Graham, \& Harris, 2007; Wehne 2003, 2008). The most studied genera, Cataglyphis, Formi $a$, and Melophorus, all come from a segment of the formi cine tribe in the ant family (Figure 2). While closely relate given the breadth of the hymenopteran order (Hunt, 2007) these genera have nevertheless had 50-75 million years of
volutionary separation (Moreau, Bell, Vila, Archibald, \& Pierce, 2006). The hymenopteran heritage stems from fly ing wasps (Hunt, 2007), so that ants are derived somehow golved from wasps. The derived character and the phylovolved from wasps. The derived character and the phyloenetic narrowness should be kept in mind as ant naviga-

## Some formicine ants



Figure 2. A part of the clade of formicine ants, derived from information contained in Moreau et al.'s (2006) phylogeny of ants. A selected number of genera, including those whose navigation has been most studied (Cataglyphis, Formica, and Melophorus), are shown. Formicine ants and myrmecin and Melophorus), are shown.Formicine ants and myrmecine
ants are two large tribes of ants. The branches do not indicate durations of times since the genera have separated but the times of separation ranged from 50 million years ago to 75 million years ago. Adapted from "Phylogeny of the Ants: Diversification in the Age of Angiosperms," by C. S. Moreau, C. D. Bell, R. Vila, S. B. Archibald, and N. E. Pierce, 2006, Science, 312, p. 102. Copyright 2006 by the American Association for the Advancement of Science. Reprinted with permission.
2 , regarding the comparison of species). Nevertheless, these genera provide great illustrations of the power of taxon-like strategies of navigation.

Path integration
The navigational toolkit includes path integration, keeping rack of the straight-line distance and direction to the starting point, the nest (Wehner 2003; Wehner \& Srinivasan, 2003). This system keeps operating even when the ant is engaging in other navigational strategies (Andel \& Wehner, 2004; Knaden \& Wehner, 2005), ready to act as a backup when other mechanisms fail. Path integration is compass based to the extent that a celestial compass is required as a component of keeping track of the path (Figure 3). The celestial compass is based on the pattern of polarized light primarily (Wehner 1994) and the position of the sun secondarily (Wehner \& Müller, 2006). The primary mode of distance estimation dif-
fers between ants and bees, probably because one walks and the other flies. In ants, a mechanism based on step counting orms the mainstay of odometry (estimation of distance trav eled on a trip; Ronacher, 2008; Wittlinger, Wehner, \& Wolf 2006, 2007), while in honeybees, the measure of optic flow forms the mainstay for odometry (Srinivasan, Zhang, Altwein, \& Tautz, 2000; Srinivasan, Zhang, \& Bidwell, 1997) What is communicated as an odometric measure in the honeybee's dance (von Frisch, 1967) is actually a measure of the amount of optic flow. Bees flying a short physical distance in a narrow channel nevertheless report a long distance in their waggle dance, and Srinivassan et al. (2000) have calibrated waggle dance, and Srinivasan et al. thang $^{\circ}$ of optic flow for each millisecond of waggling. Functionally, optical-flow cues in

Route mechanisms


Legend

Figure 3. Schematic illustration of route mechanisms in ants. In path integration, the straight-line distance and direction to the starting point (typically home) is kept track of during the entire journey. Navigation based on path integration consists in running off the global vector calculated up to the point of homing Such a vector contains both distance and directional information. The mechanism uses the sk compass to determine directions. The local vector contains set of instructions to travel in a direction acoarding to the sy

 next servomechanism are met. In a panoramic compass or visual compass, the ant travels in a direction determined according to the panoramic visual cues
fight are more reliable indicators than internal measure such as energy expenditure or number of wing beats. That is because air is a volatile medium that can blow the flier about and wreak havoc with the internal cues. For animals mostly anchored to the ground (excepting unusual circumstance in which the wind blows the animal off course), a measure based on gait forms a reliable odometric cue.

The ability at path integrating differs across desert ant spe cies, in the direction in accord with intuitive predictions. At least this is true for distance estimation or odometry. Beween two species that have been explicitly compared, the one living in the more landmark-free habitat, Cataglyphis fortis in North Africa, does better than the one living in landmark-rich habitat, Melophorus bagoti in Central Australia (Cheng, Narendra, \& Wehner, 2006; Narendra, Cheng, \& Wehner, 2007). The thermophilic C. fortis (R. Wehner \& S. Wehner, 2011) lives on the salt pans of North Africa, with at most a few low bushes serving as possible landmarks. Most nests are located in the open, a good distance from us able landmarks, making the habitat virtually landmark free. M. bagoti, also a highly thermophilic desert ant (Christian \& Morton, 1992; R. Wehner \& S. Wehner, 2011), is wide spread in Central Australia and inhabits a cluttered envi onment filled with grass tussocks, bushes, even tall trees and in some areas, distant hills (Cheng et al., 2009; Muser Sommer, Wolf, \& Wehner, 2005; Schultheiss, Schwarz, \& Wystrach, 2010). The paucity of visual cues means that path integration is more important for $C$. fortis than it is for $M$. bagoti, as it is typically the only means of navigating home.

The ants' ability to estimate the outbound distance to a feeder was measured. The direction of travel was constrained by keeping the ants running in narrow channel open above to the sky. The measure was unsystematic error, or variance in the odometric estimate of a group tested at an outbound distance. The group variance was bigger in M. bagoti than in $C$.fortis, indicating a species difference This pattern has recently been replicated on the open field (Bühlmann, Cheng \& Wehner 2011). An unusual M bagoti Bühman, in balloon launching field devoid of plagot est field was compared with $C$. fortis in their typical open bita. Perps the most telling comparison toplan habitat. Pe haps the the thace on he training field itself, withou any capture or displacement on the part of experimenters. The ants came to a feeder, and heir return paths home from the feeder were recorded after being trained for two days. In this case, the visual conditions encountered on the homebound journey matched the conditions encountered on training runs for both species. Even in this familiar situation, M. bagoti performed worse than C. fortis. Their distance estimates had larger variance, their paths were more tortuous, and far fewer of them found their way directly to the nest without the need for searching ( 2 out of 18 vs. 18 out of 20 for $C$.fortis). This species differ-
ence suggests adaptive specialization for path integration in C. fortis, but of course, the hypothesis of adaptive special ization needs empirical support from more experiments and a lot more than two species (see Shettleworth, 2010, ch. 2) It would also be informative to compare salt-pan dwelling species (hence, living in an open habitat) with semi-desert dwelling species (hence, living with cluttered scenery) with in each genus.

The directional estimate, on the other hand, is comparable between the two species. Ziegler and Wehner (1997) tested the memory decay of the directional component of the home vectors in C.fortis ants after 12 or 15 m outbound distance. At zero delay, they obtained a circular distribution of headng directions with vector length $r=0.97$ (with 1.00 indicat g perfect concordance between all ants). Narend (2007 ing per $M$. bound dit 12 m Wehned The ver for Wehn tribution of heading directions measured $r=0.98$. M. bagot from the same study, however, only achieved $r=0.89$ at 6 m outbound distance. M. bagoti might have used panoramic andmarks on their outbound trip to help in directional es timation, with increasing travel distance adding such land mark-based compass information. Testing on an open fiel or both species, Bühlmann et al. (2011) also found simi lar directional scatter in path integration in the two species under a variety of conditions. M. bagoti had a surrounding scene of distant trees, which might have provided them som compass information to keep track of the direction traveled

A functional explanation for the lack of species differenc es in determining a direction based on the celestial compas is that the sky compass functions in other mechanisms than path integration. That might make it equally important for the two species. One such mechanism is the local vector described next

## Local vector

The celestial compass is also used to execute local vector often called sensorimotor vectors in bees; Cheng, in press) The local vector can be thought of as a servomechanism that uses the celestial compass, and is triggered under appropriate contextual conditions (Figure 3). Travel instructions for ocal vector are based on the sky compass, and not terrestrial objects. obje w. Ti, vector. in er wods, a ravel diection that is the vector. according to the sky compass is linked to appropriate trig gering conditions, such as coming out of a reeder in an ex perimental situation, or rounding an obstacle in natural navi gation. The ant heads in that compass direction until suitable conditions for engaging the next servomechanism are met

A human example might be: when you come to the gas staion, turn north and go until you come to a traffic light. In the first example in ants in which the term was coined, M. Col lett, T. S. Collett, Bisch, and Wehner (1998) trained deser ants $C$. fortis to make a two-legged journey to obtain food at a feeder. They had to head north over open ground to the mouth of a narrow channel, and then turn $90^{\circ}$ (to the west) to go to the end of the channel. On the return journey the ant had to reverse the two legs. On some tests. M. Collett et al ad the direction of the chanel on the retum trip, for changed the direction of the channel on the return trip, for xample making the ants head southeast instead of the usual ast. They observed that upon exiting the channel, the ants till headed south for an initial segment. This was the loca vector, the normally appropriate compass direction to take upon the contextual situation of exiting from the channel This direction is in conflict with the direction dictated by path integration (also called the global vector), which should point the ants directly toward their nest. After a short seg ment traveling south, however, global path integration then took over, and the ants headed in the fictive nest direction.
C. fortis lives in a saltpan habitat largely devoid of surounding landmarks. It is likely that for ants living in andmark-rich habitat, a panoramic view can act as a contex or executing local vectors. The use of the local vector was demonstrated in a recent study on the Australian desert ant M. bagoti (Legge, Spetch, \& Cheng, 2010). The ants were trained to enter a cylindrical arena and go to the middle of it for food. The arena had high enough walls that few objects were visible over the top of the walls. And another study showed that objects at high elevations have little influence on navigation anyway (Graham \& Cheng, 2009b). Legge et al.'s ants had to exit in a particular direction out of the arena. An obvious set of landmarks (to the experimenters anyway) was set at the exit point to provide what the authors thought was a powerful beacon. For example, one landmark was a prominent yellow diamond against the dark wall of the area. But tests with the beacon rotated to a different direction showed, surprisingly, that it had no control over the orientaion of the ants. The ants relied on heading in a compass direction for the exit, executing a local vector. Importantly, in one experiment, the exit direction differed from the bee ine direction to the nest, meaning that the direction of the ocal vector conflicted with the global vector as calculated from path integration. Presumably, the context of being surrounded by high walls acted as the contextual trigger for the local vector.

Ants also use landmarks encountered along their route of travel, and something akin to local vectors seems to be a play in these journeys as well. In the course of repeated journeys to and from a feeder, they typically develop stereotypical routes in a cluttered habitat (wood ants, genus Formica Rosengren, 1971; C. fortis: Wehner, Michel, \& Antonsen

1996; M. bagoti: Kohler \& Wehner, 2005; Sommer, von Beeren, \& Wehner, 2008). Experimental landmarks are used for steering a route (C. fortis: T.S. Collett et al., 1992; M bagoti: Wystrach, Schwarz et al., 2011). How might ants use such landmarks? Some mechanistic details are provided in an elegant study by M. Collett (2010) on C. fortis in North Africa. The landmark conditions were the simplest possible a single cylindrical landmark stood on the route between feeder and the nest, a little off to the side of the straight line connecting feeder and nest. In such situations, ants end up learning to steer a route, and do not require information from path integration (Kohler \& Wehner, 2005; Narendra, 2007b Sommer et al. 2008; Wystrach, Schwarz et al., 2011). Th ants typically took a route between feeder and nest that curved gently on the side opposite the landmark. By capturing ants as they approach the nest, but before they entere the nest, M. Collett could then release the ant for another trip home from a variety of locations, and compare such paire journeys. Results suggest that the ants steered a continuou course based on the direction to the landmark. I would de scribe the model as a continuous series of local vectors, each riggered by viewing the landmark in a particular compas direction. The view of the landmark in a particular direc tion, or perhaps on a particular part of the eye (the two being typically confounded) triggers travel in a particular compass direction determined according to the sky compass.

Panoramic context and visually based guidance
On the use of landmarks encountered en route, anothe species difference between M. bagoti and C. fortis can be pointed out, this time in favor of $M$.bagoti. The two specie were tested in simple discrimination learning in two-choice boxes (Schwarz \& Cheng, 2010). The decision box had two potential exits side by side. One side was black while the other was white, and the ants had to choose the black side to exit the box and get home, whether it was on the right or eft side. $M$. bagoti learned the task readily, but $C$. fortis on the whole failed to learn the task even with repeated train ing. The pattern again suggests an adaptive specialization Shettleworth, 2010, ch. 2) in M. bagoti, the ant living in cluttered habitat, for using landmarks. But again, more ex periments need to be done on more species to confirm the hypothesis.
Legge et al. (2010) thought that the panoramic surround provides contextual cues because the visual panorama ha been shown to act as a contextual cue in other situations in hymenopterans. For example, panoramic contextual cues an make honeybees tolerate (generalize across) large dis crepancies in landmark characteristics. T. S. Collett and Kel ber (1988) trained honeybees to forage from two differen platforms placed outdoors 40 m apart. With the surrounding buildings and trees, it meant that the panoramic cues were
ery different on the two platforms. One platform had two yellow cylinders serving as landmarks, with the food on one side of the two cylinders. The other platform had a blue trian ge standing upright as a landmark, base on the ground, with the food on the opposite side. To ensure that the bees had to rely on the landmarks, the entire configuration of landmarks and food was shifted around on the platform from trial to tial. Honeybees learned the tasks readily, as they do in othe dual-task situations (Cheng Collett, \& Wehner 1986). The arched around the target location even when the sugar wa arched arour war er and tiny ring indicating the food source were removed landmarks were offer that is, either patform-2 se of landmarks were offered, that is, either platform- 2 land marks on platform 1 or else platform-1 landmarks on platform 2. The bees searched most at the side appropriate for he platform, as if treating yellow cylinders as an aberran blue triangle or vice versa, and tolerating the considerable mismatch in landmark characteristics. The interpretation is that the contextual cues, presumably the panorama, drove he bees to retrieve the appropriate landmark memory for the platform.
Subsequent research has upheld this kind of finding (Col born, Ahmad-Annuar, Fauria, \& Collett, 1999; T. S. Collett Fauria, Dale, \& Baron, 1997; review: T. S. Collett, Fauria \& Dale, 2003). As perhaps the best evidence for the modulatory role played by the panorama, T. S. Collett et al. (1997) trained honeybees to enter a cylindrical arena placed outdoors. The location of the cylinder, and hence the panorama around the cylinder, served to modulate the choices that the bees made inside the arena, now with the view of the panorama cut off. Thus, at the point when the bees were making decisions, the natural panorama was gone; the bees could not be using those panoramic cues directly for guidance.

Similar results can be found in ants. In lab experiments on wood ants (Formica rufa), Graham, Durier, and Collett (2004) trained them to search at the middle between two black cylinders for food (sugar water). One cylinder was larger than the other, so that at the goal, the two landmarks subtended different retinal angles. After training, a crucial fest was given in which the two landmarks were the same intermediate size. If the ants could interpret which test landmark corresponded to the big one, and which to the small one, then they should search closer to one landmark (the supposed big one) than the other. They failed to do this when white curtains surrounded the platform on which they were oraging. Instead, they searched at the middle between the two cylinders, as if unable to distinguish which of the landmark to treat as the large one. Results were different, however, when the ants were given strong panoramic contextual cues, in the form of black shapes covering one wall. They then behaved as if they interpreted one of the landmarks as the large one, and searched closer to it. The contextual
cues were thought to disambiguate the identity of confus able landmarks. The authors pointed out that when the ant turned to face one landmark, the patterned wall would fal on one eye, whereas when they faced the other landmark, the patterned wall would fall on the other eye.

Results such as these have led to the theoretical view that contextual cues act as powerful triggers for navigational be havior, forging "associative links between long-term memories" (1. S. Collett \& Collett, 2002, p. 542) or serving occasion setters for the operation of servomechanisms (Figure 1 in Cheng, in press). It is important to execute these axon-like mechanisms at the right time and place. Contex ual cues of which panoramic cues are an important subset provide powerful modulators. The disambiguating role of he panorama featured in T. S. Collett and Kelber (1988), T. S. Collett (1997), T. S. Collt and Kelber (1988), T . Colltt et al. (1997), and Grahan et al. (2004) points to ne solution to the problem of re-identificaion or perceptua Other Other contextual cues can add to he power of running servomechanisms or taxon-like routines. They include time of day Koltermann, 1971; Prabhu \& Cheng, 2008; Wahl, 1932) motivational status (outbound to forage vs. homebound with ood; Dyer, Gill, \& Sharbowski, 2002), and time-place com binations of circadian-timed episodic-like memory (Pahl Zhu, Pix, Tautz, \& Zhang, 2007).

## Using the whole scene for navigation

If the panorama can act as a powerful contextual cue, is it also good for directly guiding navigation? Scene analysis in natural settings suggests that panoramic cues are very usefu (Stürzl \& Zeil, 2007; Zeil, Hoffmann, \& Chahl, 2003). Von Frisch and Lindauer's classic (1954) work suggests guidance by large-scale landmarks. In fact, von Frisch and Lindauer's (1954) study showed that the bees were not inclined to head to clumps of landmarks such as an isolated group of trees, when they were in the wrong compass direction. Mor recently, Dyer and Gould (1983) and Towne and Moscrip (2008; Towne, 2008) replicated findings on the use of largescale landmarks in honeybees. Even on cloudy days, ruling out the use of the sky compass, the bees managed to fol ow the large-scale landmarks that they were trained to use Towne and Moscrip (2008) put forth a mechanistic hypoth sis based on a serendipitous finding from a failed contro condition. The interesting finding and ensuing hypothes more than made up for the failure to control for unsuspecting ues. As a control, trained bees were displaced to a novel lo cation that supposedly did not contain anything resembling the landmarks that the bees had been trained to use. The bee were supposed to be lost and head off in random directions. The control failed in that the bees headings were directed. Comparing panoramic photos at the training and test site (Figure 4), Towne and Moscrip suggested that the honeybee means is the one-dimensional circular representation of how high the tops of terrestrial objects are. It is the line where ground objects such as trees and bushes meet the sky, strip ping away characteristics of the ground objects such as thei

Further manipulations on the natural scene surrounding eeder showed that not all of the panorama is of equal importance (Graham \& Cheng, 2009b). When the higher el the ants were still oriented. But ants relying only on the high


Figure 4. An experimental control condition conducted by Towne and Moscrip (2008). A. A panoramic photo of the training setup, with the hive at $H$. The bees were trained to fly toward $D$ to a feeder. B. The test situation the next day in a completely different environment, with the tested hive at $H$. The bees were expected to be random in orientation because the features of the scenery mismatch, but were instead oriented toward F. Towne and Moscrip noted an uncanny similarity between the of the scenery mismatch, but were instead oriented toward F. Towne and Moscrip noted an uncanny similarity between the
skylines of the two situations. Reprinted from "The connection between landscapes and the solar ephemeris in honeybees," by W. F. Towne and H. Moscrip, 2008, Journal of Experimental Biology, 211, p.3734. Copyright 2008 by The Company of Biologists. Reprinted with permission.
colors and patterns.
elevations were not oriented

A photo from a serendipitous control condition hardly constitutes solid evidence, and it is hard to control the sky line for honeybees, which can attain quite a height in flying. Testing whether the skyline might be used is far more ame nable in ants, which walk on the ground. Skylines can be re produced in artificial arenas (Graham \& Cheng, 2009a). The produced in artificial arenas (Graham \& Cheng, 2009a). The
skyline or the tops of the terrestrial objects as viewed from the feeder was measured directly every $15^{\circ}$, and an artificial kyline was created using black cloth and a couple of tall black landmarks (Figure 5). Linear interpolation was used between the measurements. Such an arena captured a rough approximation of the skyline at the feeder, but nothing else of the visual surround. The colors in the surrounding scene were stripped away and rendered all black, and distance to objects were equalized. Trained ants that had grabbed piece of food at the feeder were placed in the middle of the rena with the artificial skyline, and their initial directions of avel were noted on a goniometer (a circle divided into sec The aren was at some distance from the training site, ors). The ar bas so vibe and nothing but sky was visible above its walls. The ants
concentrated their directions of headings in the nest direconcentrated their directions of headings in the nest direc
ion according to the artificial skyline, and they did so even ion according to the artificial skyline, and they did so eve when the arena was rotated with respect to the true compa direction to the nest. Giva how degraded the atificial sky line was, it seems to be a readily usable cue.

Much remains to be researched on the use of skyline, from both mechanistic and comparative, evolutionary perspectives. On the mechanistic side, how is the skyline extracted The sky is brighter than ground objects. But using bright hess contrast as a cue requires adjusting the threshold valu separating sky from ground. Everything is brighter when th sun is out than when the sun is behind a cloud. Möller (2002) showed that the contrast between ultraviolet (UV) light and green light provides a theoretically attractive dimension. Th sky contains more UV wavelengths relative to ground ob jects, which reflect more wavelengths that look green to us An opponent-processes channel that measures UV - green in some way could be used to segregate sky from ground. Such an opponent-processes channel would provide constancy in the face of changes in overall illumination. The theory is at tractive, but the empirical evidence that any insect uses such a channel is currently lacking. On the comparative side, how widespread is the use of skyline information? Is it common to all hymenopterans, or has it evolved independently i multiple lineages, in those for whom the trait would be useful? Only comparative research can tell.

Work has only just begun to model how well skyline in ormation can guide journeys. Using panoramic photo taken at the habitat where M. bagoti live (our field site at


Figure 5. An experiment by Graham and Cheng (2009a) showing the use of the skyline contour by desert ants Melophoru bagoti. A. A panoramic view from the feeder of the surrounding scenery. B. A panoramic view from the middle of the test arena, at which trained ants were released for a test. The arena matched the elevation of the actual scenery at the feeder every $15^{\circ}$. C. An actual photo of the test arena, with a goniometer at the center. Ants were oriented in the nest direction according to the skyline no matter how the test arena was oriented in compass direction. Adapted from "Ants use the panoramic skyline as a visual cue during navigation," by P. Graham and K. Cheng, 2009, Current Biology, 19, p. R936. Copyright 2009 by Elsevier. Reprinted with permission.

Alice Springs), Graham and colleagues have made a first ing not much worse than pixel-by-pixel matching, which of attempt (Philippides, Baddeley, Cheng, \& Graham, 2011). course encompasses much more information. The size of The photos were taken along a number of straight transects, catchment areas (Zeil et al., 2003) is an important measure unwarped, and then used for modeling in silico. The sky- of how well a strategy works. This represents the distance at line proved useful for view-based matching, often perform- which the procedure would actually succeed in bringing the
gent to the goal. Under suitable conditions, using a skyline can produce catchment areas with 4 m to 7 m radius, useful distances for M. bagoti's foraging trips. Thus, Muser et al (2005), reported that $90 \%$ of foraging runs were confined within a maximum distance of 20 m from the nest, with the average maximum ant-nest distance measuring 10.6 m . A small number of panoramic matches then, can in principle guide a homeward journey.

In fact, multiple memories might not even be necessary in some circumstances. In a cluttered lab setting, robots could navigate a curved S -shaped route based on a holistic representation (Baddeley, Graham, Philippides, \& Husbands, 2011). Such a representation picks out (learns to use) the most useful features for distinguishing locations on vs. off e familiar route It is also likely that other parameters than he familiar rous. I by ants in panoramic matching. Much he skyline are used by ants in panoramic matching. Muc

Panoramic matching, no matter what parameters or fea ures are used for matching, works like a panoramic com pass based on terrestrial cues (Graham, Philippides, \& Baddeley, 2010; Wystrach, Beugnon, \& Cheng, 2012; Wystrach Cheng, Sosa, \& Beugnon, 2011; see Figure 3). One reviewer suggested that it is akin to beaconing, except that the entire panoramic scene acts as a beacon. In traditional beaconing, he identified object defines the direction of travel: towar t. Unlike a traditional beacon strategy in which the target object needs to be picked out, in the panoramic terrestria compass strategy, a remembered direction of travel according to the encoded panorama must be learned. As the entire panorama is used, no single object needs to be segregated and identified. The panoramic terrestrial compass can also be thought of as an oriented panoramic view. It might, for example, be defined in terms of a retinal code, for example how high the skyline should be on each part of the eye. In finding the correct direction of travel, the ant would rotate on the spot until it finds the best match between the current panorama and the remembered panorama. The direction in which she is facing is then the direction of travel (Wystrach t al., 2012). The traveler might match as many remembered anoramas as are necessary to negotiate all the segments of route. Such a mechanism strikes me as a form of guidance sensu O'Keefe and Nadel (1978). Evidence from ants suggests that entire scenes, or at least visual information over a arge area of the eyes are used for navigation (Graham, Fauria, \& Collett, 2003; Wystrach, Beugnon, \& Cheng, 2011) and that the panoramic terrestrial compass works especially well for travel along a familiar route (Wystrach et al., 2012)
In lab conditions, or in open saltpan habitats, landmark tand out readily against the uniform background. In som cases, such as M. Collett's (2010) research on C.fortis, only a single landmark is used. The landmark in this case consiti
par of the parama, for instance, ring the skyline. It is possible that M. Collett's ants wer a sequence of skylines to guide their behavior.
In wood ants tested in the lab, a single beacon (black cyl inder) can stand out against the background of the room and serve to attract foragers (Graham et al., 2003). The ant naturally approached the beacon on their way to a feeder, even though the beacon was off to one side and required detour. They did this when starting from different locations, so that they did not appear to be executing a motor routine (such as turn so many degrees and walk). But did they learn to approach a beacon object? A most interesting manipulation on the trained ants was to remove the beacon. The ant still approached the location of the absent beacon enroute to he feeder! They apparently did not need the beacon object itself. One They appar. I like to offer is the obe obect not approaching an extracted and identified object, but a salient part of the panorama. In the process, they learned to ap proach a direction based on the rest of the panorama as well They could then approach that part of the panorama eve when the object was missing, thus showing the robustness of the panoramic terrestrial compass
But what about ants traveling in natural habitats with com plex visual information? The Australian desert ant M. bagoti makes an ideal subject in this case, and an attempt was mad to train the ants to use a large landmark in cluttered condi ions (Wystrach, Beugnon, et al., 2011). The landmark wa black cloth 3 m wide and 2 m high, held up just behin the nest (Figure 6). The training field was cleared of clut ter, so that the beacon and the panorama were clearly visible hroughout the journeys between feeder and nest. Other objects being a good distance away, the high-contrast edges of the beacon provided dynamic cues on approach: the beacon expanded in azimuthal size (width) from $54^{\circ}$ to $118^{\circ}$ in the ast 2 m of approach to the nest. Surely, it specified the nest's location well. To the primate visual systems of humans do ing research there, the beacon looked obvious and stood out But to the ants, it did not

The ants were trained to run repeatedly between their nes and a feeder 10 m from their nest. Then a number of different tests were conducted, with each individual ant only participating in a single test. In one test, the beacon was set up at a distant test field with an unfamiliar panoramic view, excepting the beacon. The ants failed to home in on the beacon, in ffect showing that they did not treat the beacon as a beacon. They failed to do so even when released just 2 m from th fictive nest position. Apparently, mismatch in the rest of th cenery mattered. Other tests were conducted at the trainin field with the beacon removed, with only the tested ant hom ing (and the others trapped temporarily at the feeder). Ant also failed to home on this test, showing that the beacon did


Figure 6. Photos of the beacon setup used by Wystrach Beugnon, et al. (2011), supplied by Antoine Wystrach. A. A substantial beacon in the form of a black cloth 3 m wide and 2 m high was set up just behind the tested nest of desert ants Melophorus bagoti. To human (primate) eyes, the beacon stands out as an obvious object to aim for in navigation B. The view as blurred to $5^{\circ}$ resolution, approximating the visual acuity of this species of ants ( $\sim 4^{\circ}$; Schwarz, Narendra \& Zeil, 2011)
form an important part of the scenery for the ants. Yet othe tests were conducted on the training field with the beacon displaced. When it was displaced $32^{\circ}$ to either side, homing again failed, the ants finding neither their nest nor the fictive nest position in front of the beacon. Even having much of the familiar panoramic context did not drive the ants to head the beacon. When the beacon was displaced $16^{\circ}$ to eith the beacon. Whid the on the no 16 to either side, ants agan id engaged in systematic searching.

But were the ants beaconing in, in the sense in which the erm is usually understood? That is, did they pick out the beacon as an object to head to, and then head straight toward it, using the identified beacon in a guidance strategy sensu O'Keefe and Nadel (1978)? Closer examination of their be havior showed otherwise. If the ants had identified the beacon as a beacon, they should have headed straight toward it at some point in the journey. Uncertainties in extracting and identifying the object should be found, if at all, near the beginning of the journey. The last few meters should be straight run, with the sizeable beacon identified by that stage of the journey. Instead, numerous ants turned around, mean-
dered, and searched before ending up in front of the beacon. Many of these turns first took place $6-8 \mathrm{~m}$ into the journey ( $2-4 \mathrm{~m}$ from the goal). Such behavior is inconsistent with guidance strategy of beaconing in.

It turns out that a panoramic view-matching strategy ex plains this kind of searching behavior as well as other hom ing behaviors not discussed here. At the beginning of the journey, much of the scenery matches, with just the beacon out of place. A view-matching strategy would drive the ant traight ahead, toward their real nest, a pattern found empiri cally. At the real nest, with the beacon missing, the match is very poor, apparently too poor for the ants to search there. In front of the beacon, however, is a partial match, a local mini mum in mismatch, so that some ants might end up searchin here Scene analysis also shows that in the middle, especial here. Sc $6-8 \mathrm{~m}$, is $y$ in the $6-8 \mathrm{~m}$ range, is , The it is oul ing, and it is out of pace. Tha, a global view-matching strategy, such as the matching or skyline pattrr (Grahaw Cheng, 2009a, 2009b) explains the searching and meande ing behavior of ants in this zone, without the need to invoke object segregation and then beaconing.
Using the panoramic scene not only gets the ants to head in a homeward direction along a regular route, but also when hey have been displaced a small amount (review: T. S. Col lett et al., 2007). We have observed ants being blown of course, so that small displacements of a few meters are eco ogically realistic. Wood ants (F. rufa) displaced centimeter from their usual starting point can find their way to an in conspicuous feeder, using landmarks set up in the lab (Du rier et al., 2003). Wood ants of another species (F. japonica) can also cope with displacements in the field over a larger distance (meters, Fukushi, 2001; Fukushi \& Wehner, 2004) The Australian desert ant M. bagoti can cope with displacements of up to 10 m sideways from their usual starting point (feeder, Narendra, 2007b), as well as smaller displacement Graham \& Cheng, 2009b; Kohler \& Wehner, 2005; Nar ndra 2007b). With smaller distances of displacement $M$ agoti $M$ ) 1 , 1 , ote follo
 , A W, No personal observations).A ert ant C.fortis located in cluttered terrain also managed to home after displacements (Wehner et al., 1996). Desert ant often search for a while near the release point, before heading off in a direction, something that displaced honeybee have also been observed to do (Menzel et al., 2005). How ants (and bees for that matter) manage to compute a home ward direction from the view at a displaced location poses an interesting research question to be explored. But some initial analysis suggests that navigational mechanisms on and of the route may differ.

In another study on M. bagoti, barriers were used to con- bered panoramas were often similar. Another mechanism fine ants of two different nests to experimentally defined areas for traveling and foraging (Wystrach et al., 2012). Furhermore, by confining the ants in this way for a number of days before the experiment began, only naïve ants were allowed to take part. The nature of their traveling experience was thus carefully controlled. Some ants were confined to an area of 1.1 m radius around their nest. Others got to forage along a 10 m route that was gently curved. Ants were then ested after different amounts of training at locations on and ff the route A zero-vector ant (captured after she had an off the route. Azero-vector an (captured after she had nearly returned to her nest) was placed successively on a goniomer at Ants typically turned on the spot initially and the oted. Ants typically tumed on the spot initially, and then headed off unhesitatingly in one direction (for an example of his kind of turning, see the supplementary movie). Impor tantly, panoramic photos from the experimental sites were taken so that models based on the actual sceneries could be evaluated against the ants' behavior.
The already described panoramic terrestrial compass model accounted for the ants' behavior on the familiar route very well, but it performed poorly for test locations off the route. The trouble arose because to perform well off the route, the most suitable remembered panorama needed to be chosen, and mistakes were often made because remem-
however, had better success in explaining the orientation of the ants off their route. And that was to compare the skyline heights of remembered and current panoramas (Figure 7) These panoramas must already have been oriented according to the sky compass (or in principle, any other compass such as a magnetic compass). In that case, if the skyline in one region of the panorama appears too high (higher than the corresponding part in the remembered panorama), then the ant was probably displaced in that direction, and should turn away from that region. Conversely, if a part of the skyline appears too low, compared to what height it should be according to the remembered view, then the ant has drifted too far from that region, and should be attracted toward that region. Such a model worked reasonably well in accounting for orientation off the route. Ants in general aimed in a di rection toward a part of the route. It would be interesting to manipulate skyline heights directly in experimental arena to test this model.
In sum, the ants seem to use different strategies on and off their familiar route. They might decide whether they are on or off the route by the quality of the match between the cur rent scene and remembered views (Wystrach et al., 2012) The calculations based on the panoramic photos show size able differences in the level of matching on and off the route


Figure 7. An illustration of the skyline-height-comparison model proposed by Wystrach et al. (2012). A hypothetical remembered skyline is compared with the currently viewed skyline. Both skylines are oriented in compass direction. That is, the $x$ axis codes absolute compass directions, based on the sky compass. The ant would be attracted by parts of the skyline that look too low in comparison with the remembered skyline (blue arrow), and repelled by sections of the skyline that look too high in comparison with the remembered skyline (green arrow). In Wystrach et al.'s (2012) model, only the attraction process was assumed. More research is needed to determine which of these processes (or both) is at work.

## Searching as taxon-like behaviour

View-based navigation and especially path integration are not perfect strategies, so that in homing, ants often need to search for their nest on natural foraging trips. I have observed such searching behavior on natural foraging trips in both C. fortis and M. bagoti. The two cases are different in that $M$. bagoti has a rich panoply of surrounding visual cues - the scenery around her nest - to guide the search, whereas $C$. fortis usually has none. The search of M. bagoti in a situation resembling that facing $C$. fortis can be studied by displacing the homing ant to a distant site so that the visual scene provides no useful cues. I now discuss searching with and without visual cues in turn.

In landmark-based search, behavioral records have been obtained on tests in which the target of search was missing Artificial landmarks defining the goal are provided in many cases. In contrast to searching in the absence of predictive visual cues (to be discussed shortly), the details of search patterns have not been documented. But search distribution that have been obtained indicate tight, concentrated search ing around the fictive goal, in honeybees searching for a missing feeder (Cartwright \& Collett, 1982, 1983), in desert ants searching for the fictive nest at a test site far from thei real nest (C. bicolor: Wehner \& Räber, 1979; C. fortis: Åkes son \& Wehner, 2002; M. bagoti: Narendra, Si, Sulikowski, \& Cheng, 2007b). Most recently, the use of landmarks around the nest has also been found in the Namibian desert ant $O c y$ myrmex robustior, a myrmecine ant (Müller \& Wehner, 2010 Wehner \& Müller, 2010). In the desert ant studies, plastic cylinders served as landmarks. For M. bagoti, the displacement of a set of landmarks meant that the rest of the scenery mismatched, a point whose significance was not fully appreciated by the authors when the research was conducted It took quite a number of trials of training, at the training site returning to their nest, before the ants learned to use the experimental landmarks at the test site. In contrast, Wystrach et al. (2012) found that even naïve ants arriving at a feede ear their nest were significantly oriented when displaced ear their nest were significantly oriented when displaced few meters away. The panoramic scenery is learned quickly ome recent advances in this topic include the matching of optic flow patterns created by motion parallax, in honeybees haaf, 2010), and turning behavior in wood ants that are finely tuned to anticipated positions of a feeder serving as a goal Lent, Graham, \& Collett, 2010).
This line of research has led to a proliferation of models on this topic, without firm agreement (Cartwright \& Collett, 1983; Möller, 2001; Narendra, Si, et al., 2007; Nicholson Judd, Cartwright, \& Collett 1999). These models of image matching differ in specifying what is matched, but what is
common to them indicates a taxon-like strategy rather than an insect locale system or a sketch map. Various kinds of parameters are thought to drive the matching in a servomechanistic fashion. The insect moves so as to reduce the discrepancy between parameters in the current image, and the remembered parameters. None of the models propose the encoding of distances to multiple landmarks or to distant sites, characteristics of map-like representations.

Detailed search paths of ants have been recorded for searches in the absence of landmarks, sometimes in onedimensional narrow channels (C. fortis: Cheng \& Wehner 2002; M. bagoti: Narendra, Cheng, Sulikowski, \& Wehner 2008), other times in the open field (Cataglyphis: Merkle Knaden, \& Wehner, 2006; Merkle \& Wehner, 2009, 2010; Müller \& Wehner 1994. Wehner \& Srinivasan 1981; bagot bagot. Schurs \& Chenlat in in

 1985a, b). An example of a search in M. bagori is shown in Figure 8. The looping area-restricted search patterns found in such ants have been thought to be an evolutionary pre cursor to cognitive searching (Hills, 2006). The pattern of searching are thought to be close to optimal (Hoffmann, 1983b).
Systematic searching in the absence of predictive visua cues was not featured in O'Keefe and Nadel's (1978) book but clearly, it is a taxon-like strategy. It is what an anima engages in when nothing familiar instructs the navigator as to a defined direction of travel. Its goal is to find, in the mini mum of time, something familiar. This would be the nest or he cues emanating from it in the case of animals search ing without landmarks (C.fortis, H.reaumuri), or else som familiar scenery from which instructions for directed trave may be derived (M.bagoti)

## Putting it all together: The power

 of a taxon-like repertoireA repertoire that I have called taxon-like has considerable utility for ants in their navigation. It suffices to steer ants traveling on habitual routes. The use of panoramic informaion also allows ants to cope with some displacements, a job hat a locale system is supposed to serve in rats. That is
 litelf in loca ions. The han, instructions for getting back to a familiar route, based o panoramic information, but does not inform the ant of its lo cation in the world. Harking back to the quote from O'Keef and Nadel (1978, p. 59) at the beginning of this paper, suc a system does not let the animal experience a "coherent


Figure 8. An example of a search pattern in a desert an Melophorus bagoti) from Schultheiss and Cheng's (2011 study, supplied by Patrick Schultheiss. The desert ant wa trained to run back and forth between a feeder and her nest On a test, she was captured just before she reached her nest on returning from a foraging trip, and placed at a distant test field marked with a grid. The circle shows the release point of the ant on the test field. Typical search pattern. began with tight loops around the release point, expanding to much larger loops as the search went on.
patial world. It delivers a set of incoherent but functional instructions based on available visual and idiothetic (based on path integration) cues. Its only sense of place is the vector delivered by path integration connecting, approximately and with cumulating errors, the traveler's current location to its home.

The taxon-like repertoire is much expanded compared with the motor routines and beaconing that form the main tay in O'Keefe and Nadel's (1978) characterization. Hence, the addition of the suffix "like" expresses some similarity, but also indicates possible differences. The ant can learn to head in a direction defined by a panorama, which is another way of describing a panoramic terrestrial compass mechanism. The taxon-like repertoire includes path integration, intructions linking context to compass-based vectors of travel (local vectors), possibly the comparison of skyline heights, and systematic searching. Aside from path integration, these strategies have not received much attention in vertebrate animals.
I have given some hints that the entire concept of beaconing, taken to be an obvious and easy routine by most of us
humans, may be doubted when it comes to ant navigation For us, beaconing means identifying an object associate with a goal and heading toward it. Beacons may also be as sociated with route instructions, such as when we turn left at the corner with the Brand X gas station. It is possible that w are beguiled by our own evolved primate visual system into thinking that that is easy and widespread. We are blessed with high-acuity foveas in the eyes, but more than that, we primates possess an entire stream devoted to object identification, called the ventral stream (Goodale \& Milner, 1992; Milner \& Goodale, 1995; Mishkin, Ungerleider, \& Macko, 1983). Most animals do not have this dedicated machinery for extracting objects. As a consequence, beaconing may be harder for many than it is taken to be. In ants, at least the ide needs serious revisions. Beacons might only attract ants in so far as they form some salient characteristic of the panorama with the full set of such attractors requiring much researc to unravel. The entire panorama might also be considered as a beacon, with some direction in it defined as the targe direction. But it is clearer to use instead the term panorami errestrial compass to distinguish this mechanism from tradi tional beaconing in which a single object is identified as the target to head toward.

I suspect that the navigational repertoire in ants, if not in all insects, might have evolved in the absence of any visual object identification. It is possible that landmark-based navi gation may not be based on landmarks in this sense. In one recent formulation regarding the honeybee, a landmark is defined as "a coincidence of several different cues in a local region of the eye" (Horridge, 2009, p. 2728). The retino topically-defined cues are in turn parameters from a limited palette that the bee processes. A thesis to explore is that in hymenopterans, the use of large-scale cues of panoramic ter restrial scenery and celestial patterns of polarized light by passes any need for navigational object extraction, whos mechanisms would prove too costly to evolve. The exten f visual cognition in hymenopterans is debated (Avarguès Weber, Deisig, \& Giurfa, 2011; Horridge, 2009). But object identification, whatever its form, might only have evolve in insects if some other function than navigation drove it volution. Thus, honerbees, which need to determine whic llowers are the most profitable, may possess some sophisticated skills of visual cognition (Avarguès-Weber et al., 2011 Giurfa, 2007).

## Map-like integration?

While the systems or modules of path integration, loca vectors, and different matching routines, each operating in isolation, do not amount to a system that we would cal cognitive mapping, integrating them by operations comput ing and combining their outputs would result in a map-lik system. Tolman (1948) expressed the idea metaphorically as being "worked over and elaborated in the central contro
oom" (p. 192). Cruse and Wehner (2011) took this metapho on board and specified that integrative processes are what makes a system a cognitive map. Thus, linking vectors to panoramic views gives a map-like flavor: from view X, the vector $y$ would bring me to location Y. As another example, the comparison of skyline heights gives a gradient in two dimensions with characteristics of a bearing map. Combining this system with representations of vectors between loations, and it looks like calibrating a map. In contrast,
 with the only integration being competition over access to a common command system that drives action.

Cruse and Wehner's (2011) modular, taxon-like system accounts for extant navigational performance in ants and bees, including performance that has been taken to show cognitive mapping abilities by some. Menzel et al. (2005) displaced bees from a feeder before their homebound journey. The bees first flew the vector, according to path integration, tha would have taken them home. Not finding themselves at thei hive, they engaged in some searching behavior. Then, mos bees found the way home. The authors interpret the homing behavior as showing reliance on a map-like representation the bees searched to locate themselves on the map, and then flew home. Yet Cruse and Wehner's (2011) model solves the case by having the agent search until it finds a landmark with which a home vector (a local vector) is linked. And in displacement experiments on $M$. bagoti (not modeled by Cruse and Wehner), the operations in turn of a skyline-height comparison routine off the route (driving the ant to the route) and a panoramic terrestrial compass routine on the route (driving the ant home) solves the case (Wystrach et al., 2012). Map like integration has so far been unnecessary for explaining hymenopteran navigation.

## Panorama and context

Recent research gives some indications of the power of using the entire visual panorama in direct guidance of navi gation. Using the entire surround serves one of the functions of a locale system, to reduce perceptual aliasing. Perceptual aliasing arises from the similarities of isolated objects such as individual bushes or trees. The locale system helps to reduce confusions by locating objects approximately on a map. A similar object at a different region on the map is treated as a different object. The taxon-like panoramic strategy reduce perceptual aliasing by taking on the maximum amount of inormation defining a place, given the constraints imposed by he insect's perceptual systems. Even the limited information provided by skylines does this job well. Wystrach et al (2012) measured the mismatch levels on and off the familia route of the ants, and the gap is sizeable: the mismatch was more than 5 times bigger at the experimental off-route tes ocations than at the on-route test location. It would be inter-
esting to compare this performance formally (quantitatively) with how a locale + taxon dual system would do
The desert isopod $H$. reaumuri attempts to deal with the re-identification problem even in the absence of externa contextual cues (Hoffmann 1985a, b). This arthropod has no vision to speak of, but builds a ring of feces around its nes as landmarks. The ring facilitates the job of finding the nest. But this means that the homing isopod needs to distinguis its ring from the rings of other families of $H$. reaumuri. How much time the isopod searches to determine whether the ring is the correct ring is determined in part by the expected prob ability (in the statistical rather than the cognitive sense) that the ring is in fact correct. Thus, a ring far from the expected location of home according to the isopod's olobal vector is deemed unlikely, and the arthropod might linger little if at al at the ring. In short, path integration is used to solve, albei only partially, the re-identification problem.

## Locale and taxon systems revisited

If a taxon-like repertoire suffices for certain kinds of navigation, such as the tasks faced by desert ants, then what kinds of ecological pressures might have driven the evoluion of locale systems, and in what animals? The taxon-lik ystem is likely evolutionarily more ancient. The rodent neurobiological literature contains hints of building a locale sys tem using components serving the taxon system. Thus, path integration might be crucial for building up metric map (Gallistel, 1990; Sheynikhovich et al., 2009). Grid cells in rats (Fyhn, Molden, Witter, Moser, \& Moser, 2004; Hafting Fyhn, Molden, Moser, \& Moser, 2005) might be a key component serving both path integration (McNaughton, Battaglia, Jensen, Moser, \& Moser, 2006) and as a "metric for the cognitive map" (Jeffery \& Burgess, 2006). A recent model of ocale and taxon spatial learning in rats starts out with view as the basis for both systems (Sheynikhovich et al., 2009) The views are panoramic, and consist only in an array of the features processed early in the visual system, edge orientations. The taxon system links views directly to behaviors much in the spirit of the view-based taxon-like procedures of ants. The locale system links path integration and views, vi he neurophysiology of place cells (O'Keefe, 1976; O'Keef \& Dostrovsky, 1971). This integrative process linking taxon components makes the system map-like

Asa final speculation, the scale of travel might dictate the heed for a locale-like system. I suspect that views of pan orama, together with the rest of the armamentarium of the axon-like repertoire, copes readily with the scale of tens of meters that desert ants deal with. A small number of view ogether with path integration can probably suffice to cope with navigational needs. On the largest scale of global travel any landmark-based systems, with or without a locale sys
m, fails. Long migrations in the dark, or over or in the sea, re not conducive to the use of landmarks of any kind. Unde such circumstances, some Earth-based locale system or bearing map needs to be used. I mean here what others call true navigation (Bingman \& Cheng, 2005; Boles \& Lohmann, 2003; Griffin, 1952). This is the ability to estimate the approximate longitude and latitude on Earth from some Earth based cue, of which magnetic cues are the best candidate date. Thus, sea turtles can estimate latitude (Lohmann, , Fhman, Ehrhart, Bagley, \& Swing, 2004) and longitude (Putman, Endres, Lohman, \& Lohmann, 2011) by using Putman, Endes. Bingman and I (Bingman \& Cheng using magnetic cues. Bingman and (Bingman \& Cheng, 2005) rgued that in volatile fluid media (water and especially air) here one may be driven off course, such an ability is neces sary for long global-scale voyages
Rats travel at an intermediate scale. In one recent study Russell, McMorland, \& MacKay, 2010), rats roamed over hundreds of meters, covering up to 8 ha of range over a week of exploring a new island on which they had been released ingly. The rat-free island was limited in size, so that with larger space to colonize, they might move even further. Is it at this scale of travel, rather than the scale found in a lab, that vectors linking different sets of views characterizing a place or region become necessary? Is this when a traveler might want to plan and design routes between locations that are not based on learned routes and path integration, something hat a map-like representation can help with? Is it at this scale when integration of bearing and sketch maps (Jacob \& Schenk, 2003) becomes necessary? In hymenopterans, a good test case is the honeybee, which, like rats, also roam over hundreds of meters or more. Although Menzel et al.'s (2005) data can be explained by the taxon-like procedures in Cruse and Wehner's (2011) model, it is worth investigating Cruse and Wehner's (2011) mode, it is worth investigating oneybee performance ot al (2005) studied bees on anple fld fled devor sklin in tor far way to define ny skyln all haracteristic

Much remains to be discovered about the evolution of navigational abilities. It is a sphere that promises to reveal much about the origins of intelligence in animals. I hope that have shown here that a repertoire of taxon-like procedures without any map-like representation, may go a long way toward solving many navigational problems.

## References

Åkesson, S., \& Wehner, R. (2002). Visual navigation in desert ants Cataglyphis fortis: are snapshots coupled to a celestial system of reference? Journal of Experimental Biology, 205, 1971-1978.

PMid:12089203
Andel, D., \& Wehner, R. (2004). Path integration in desert
ants, Cataglyphis fortis: how to make a homing ant run away from home. Proceedings of the Royal Society B-Biological Sciences, 271, 1485-1489
doi.org/10.1098/rspb.2004.2749
PMid:15306320 PMCid:1691744
Avarguès-Weber, A., Deisig, N., \& Giurfa, M. (2011). Visual cognition in social insects. Annual Review of Entomology, 56, 423-443.
doi.org/10.1146/annurev-ento-120709-144855
PMid:20868283
Baddeley, B., Graham, P., Philippides, A., \& Husbands, P. (2011). Holistic visual encoding of ant-like routes: Navi gation without waypoints. Adaptive Behavior, 19, 3-15. doi.org/10.1177/1059712310395410
Bingman, V. P., \& Cheng, K. (2005). Mechanisms of animal global navigation: Comparative perspectives and enduring challenges. Ethology, Ecology and Evolution, 17, 295 318. doi.org/10.1080/08927014.2005.9522584

Boles, L. C., \& Lohmann, K. J. (2003). True navigation and magnetic maps in spiny lobsters. Nature, 421, 60-63. doi.org/10.1038/nature01226 PMid:12511953
Bühlmann, C., Cheng, K., \& Wehner, R. (2011). Vector based and landmark-guided navigation in desert ants inhabiting landmark-free and landmark-rich environments Journal of Experimental Biology, 214, 2845-2853.
doi.org/10.1242/jeb. $054601 \quad$ PMid: 21832127 way rat. Bethesda, MD: U.S. Department of Health, Eduway rat. Bethesda,
Cartwright, B. A., \& Collett, T. S. (1982). How honey bees use landmarks to guide their return to a food source. $N a$ ture, 295, 560-564. doi.org/10.1038/295560a0
Cartwright, B. A., \& Collett, T. S. (1983). Landmark learn ing in bees. Journal of Comparative Physiology A, 151 521-543. doi.org/10.1007/BF00605469
Cheng, K. (2008). Geometry and navigation. In M. E. Jef feries \& W.-K. Yeap (Eds.), Robotics and cognitive approaches to spatial mapping (pp. 145-161). Berlin, Heidelberg, New York: Springer
doi.org/10.1007/978-3-540-75388-9 9
Cheng, K. (in press). Arthropod navigation: Ants, bees crabs, spiders finding their way. In E. A. Wasserman \& T. R. Zentall (Eds.), Handbook of Comparative Cognition (in press). Oxford: Oxford University Press.
Cheng, K., Collett, T. S., \& Wehner, R. (1986). Honeybees learn the colours of landmarks. Journal of Comparative Physiology A, 159, 69-73. doi.org/10.1007/BF00612497 Cheng, K., Narendra, A., Sommer, S., \& Wehner, R. (2009) Traveling in clutter: Navigation in the Central Australian desert ant Melophorus bagoti. Behavioural Processes, 80 261-268. doi.org/10.1016/j.beproc.2008.10.015 PMid:19049857
Cheng, K., Narendra, A., \& Wehner, R. (2006). Behaviora
ecology of odometric memories in desert ants: acquisition, retention, and integration. Behavioral Ecology, 17, 227 235. $\quad$ doi.org/10.1093/beheco/arj017

Cheng, K., \& Wehner, R. (2002). Navigating desert ants (Cataglyphis fortis) learn to alter their search patterns on their homebound journey. Physiological Entomology, 27, 285-290. doi.org/10.1046/i.1365-3032.2002.00298.x
Christian, K. A., \& Morton, S. R. (1992). Extreme thermo philia in a Central Australian ant, Melophorus bagoti. Physiological Zoology, 65, 885-905.
Colborn, M., Ahmad-Annuar, A., Fauria, K., \& Collett, T. S (1999). Contextual modulation of visuomotor associations in bumble-bees (Bombus terrestris). Proceedings of the Royal Society of London, Series B-Biological Sciences, 266, 2413-2418. doi.org/10.1098/rspb.1999.0940 PMCid:1690472
Collett, M. (2010). How desert ants use a visual landmark for guidance along a habitual route. Proceedings of the National Academy of Sciences USA, 107, 11638-11643. doi.org/10.1073/pnas. 1001401107
PMid:20534539 PMCid:2895085
Collett, M., Collett, T. S., Bisch, S., \& Wehner, R. (1998) Local and global vectors in desert ant navigation. Nature 394, 269-272. doi.org/10.1038/28378
Collett, T. S., \& Collett, M. (2002). Memory use in insect visual navigation. Nature Reviews Neuroscience, 3, 542552. doi.org/10.1038/nrn872 PMid:12094210

Collett, T. S., Dillmann, E., Giger, A., \& Wehner, R. (1992). Visual landmarks and route following in desert ants. Journal of Comparative Physiology A, 170, 435-442 doi.org/10.1007/BF00191460
Collett, T. S., Fauria, K., \& Dale, K. (2003). Contextual cues and insect navigation. In K. J. Jeffery (Ed.), The neurobiology of spatial behaviour (pp. 67-82). Oxford: Oxford University Press
Collett, T. S., Fauria, K., Dale, K., \& Baron, J. (1997). Places and patterns: a study of context learning in honeybees Journal of Comparative Physiology A, 181, 343-353. doi.org/10.1007/s003590050120
Collett, T. S., Graham, P., \& Harris, R. A. (2007). Novel landmark-guided routes in ants. Journal of Experimental Biology, 210, 2025-2032. doi.org/10.1242/jeb.000315 PMid: 17562876
Collett, T. S., \& Kelber, A. (1988). The retrieval of visuospatial memories by honeybees. Journal of Comparative Physiology A, 163, 145-150. doi.org/10.1007/BF00612004
Cruse, H., \& Wehner, R. (2011). No need for a cognitive map: Decentralized memory for insect navigation. PLoS Computational Biology, 7, e1002009.
oi.org/10.1371/journal pcbi. 100200
PMid:21445233 PMCid:3060166
Dittmar, L. (2011). Static and dynamic snapshots for
goal localization in insects. Communicative \& Inte grative Biology, 4, 14-20.
Dittmar, L., Stürzl, W., Baird, E., Boeddekker, N., \& Egelhaaf, M. (2010). Goal seeking in honeybees matching of optic flow snapshots? Journal of Experimental Biology, 213, 2913-2923.
doi.org/10.1242/jeb.043737
PMid:20709919 Durier, V., Graham, P., \& Collett, T. S. (2003). Snap shot memories and landmark guidance in wood ants. Current Biology, 13, 1614-1618.
doi.org/10.1016/j.cub.2003.08.024 PMid:13678592
Dyer, F. C., Gill, M., \& Sharbowski, J. (2002). Motiva tion and vector navigation in honey bees. Naturwissenschaften, 89, 262-264.
doi.org/10.1007/s00114-002-0311-5 PMid:12146791 Dyer, F. C., \& Gould, J. L. (1983). Honey bee naviga tion. American Scientist, 71, 587-597.
Etienne, A. S., \& Jeffery, K. J. (2004). Path Integration in mammals. Hippocampus, 14, 180-192.
doi.org/10.1002/hipo. 10173 PMid: 15098724
Fukushi, T. (2001). Homing in wood ants, Formica ja ponica: use of the skyline panorama. Journal of Experimental Biology, 206, 535-541.
Fukushi, T., \& Wehner, R. (2004). Navigation in wood ants Formica japonica: context dependent use of landmarks. Journal of Experimental Biology, 207, 2431-3439. doi.org/10.1242/jeb. 01159
PMid:15326219
Fyhn, M., Molden, S., Witter, M. P., Moser, E. I., \& Moser, M.-B. (2004). Spatial representation in the entorhinal cortex. Science, 305, 1258-1264
doi.org/10.1126/science. 1099901 PMid:15333832 Gallistel, C. R. (1990). The organization of learning. Cambridge, MA: MIT Press.
Giurfa, M. (2007). Behavioral and neural analysis of associative learning in the honeybee: a taste from the magic well. Journal of Comparative Physiology A, 193, 801-824. doi.org/10.1007/s00359-007-0235-9 PMid:17639413
Goodale, M. A., \& Milner, A. D. (1992). Separate visual pathways for perception and action. TRENDS in Neurosciences, 15, 20-25.
doi.org/10.1016/0166-2236(92)90344-8
Graham, P., \& Cheng, K. (2009a). Ants use the panoramic skyline as a visual cue during navigation. Current Biology, 19, R935-R937.
doi.org/10.1016/j.cub.2009.08.015 PMid:19889365
Graham, P., \& Cheng, K. (2009b). Which portion of the
natural panorama is used for view based navigation in the Australian desert ant? Journal of Comparative Physiology A, 195, 681-689.
doi.org/10.1007/s00359-009-0443-6 PMid:19404647
Graham, P., Durier, V., \& Collett, T. S. (2004). The binding and recall of snapshot memories in wood ants (Formica rufa L.). Journal of Experimental Biology, 207, 393-398. doi.org/10.1242/jeb.00771 PMid:14691086
Graham, P., Fauria, K., \& Collett, T. S. (2003). The influence of beacon-aiming on the routes of wood ants. Journal of Experimental Biology, 206, 535-541. doi.org/10.1242/jeb. 00115 PMid: 12502774
Graham, P., Philippides, A., \& Baddeley, B. (2010). Animal cognition: Multi-modal interactions in ant learning. Current Biology, 20, R639-R640. doi.org/10.1016/j.cub.2010.06.018 PMid:20692612
Griffin, D. R. (1952). Bird navigation. Biological Reviews, 27, 359-400.
doi.org/10.1111/j.1469-185X.1952.tb01509.x
Hafting, T., Fyhn, M., Molden, S., Moser, M.-B., \& Moser, E. I. (2005). Microstructure of a spatial map in the entorhinal cortex. Nature, 436, 801-806. doi.org/10.1038/nature03721 PMid:15965463
Hills, T. T. (2006). Animal foraging and the evolution of goal-directed cognition. Cognitive Science, 30, 3-41. doi.org/10.1207/s15516709cog0000 50
PMid:21702807
Hoffmann, G. (1983a). The random elements in the systematic search behavior of the desert isopod Hemilepistus reaumuri. Behavioral Ecology and Sociobiology, 13, 81-92. doi.org/10.1007/BF00293798
Hoffmann, G. (1983b). The search behavior of the desert isopod Hemilepistus reaumuri as compared with a systematic search. Behavioral Ecology and Sociobiology, 13, 93-106. doi $\mathrm{org} / 101007$ /BF00293799
Hoffmann, G. (1985a). The influence of landmarks on the systematic search behavior of the desert isopod Hemilepistus reaumuri I. Role of the landmark made by the animal. Behavioral Ecology \& Sociobiology 17, 325-334. doi.org/10.1007/BF00293210
Hoffmann, G. (1985b). The influence of landmarks on the systematic search behavior of the desert isopod Hemilepistus reaumuri II. Problems with similar landmarks and their solution. Behavioral Ecology \& Sociobiology, 17, 335-348
doi.org/10.1007/BF00293211
Horridge, A. (2009). What does an insect see? Journal
of Experimental Biology, 212, 2721-2729 doi.org/10.1242/jeb.030916 PMid:19684204 Hunt, J. H. (2007). The evolution of social wasps. New York: Oxford University Press
doi.org/10.1093/acprof:oso/9780195307979.001.0001 Jacobs, L. F., \& Schenk, F. (2003). Unpacking the cognitive map: The parallel map theory of hippocampal function. Psychological Review, 110, 285-315.

## doi.org/10.1037/0033-295X.110.2.285

PMid:12747525
Jeffery, K. J. (2010). Theoretical accounts of spatia learning: A neurobiological view (commentary on Pearce, 2009). Quarterly Journal of Experimental Psychology, 63, 1683-1699.
doi.org/10.1080/17470210903540771
PMid:20204918 PMCid:3160474
Jeffery, K. J., \& Burgess, N. (2006). A metric for the cognitive map: found at last? TRENDS in Cognitive Sciences, 10, 1-3. doi.org/10.1016/j.tics.2005.11.002 PMid:16309947
Knaden, M., \& Wehner, R. (2005). Nest mark orientation in desert ants Cataglyphis: what does it do to the path integrator? Animal Behaviour, 70, 1349-1354. doi.org/10.1016/j.anbehav.2005.03.030
Kohler, M., \& Wehner, R. (2005). Idiosyncratic route memories in desert ants, Melophorus bagoti: How do they interact with path integration vectors? Neurobiology of Learning and Memory, 83, 1-12.

## doi.org/10.1016/i.nlm.2004.05.011

## PMid:15607683

Koltermann, R. (1971). 24-Std-Periodik in der Langzeiterinnerung an Duft- und Farbsignale bei der Ho nigbiene [Circadian memory rhythm after scent and colour training with honey-bees]. Zeitschrift fuir Vergleichende Physiologie, 75, 49-68.

## doi.org/10.1007/BF00335137

Legge, E. L. G., Spetch, M. L., \& Cheng, K. (2010) Not using the obvious: desert ants, Melophorus bago$t i$, learn local vectors but not beacons in an arena. An imal Cognition, 13, 849-860.
doi.org/10.1007/s10071-010-0333-x
PMid:20567867
Lent, D., Graham, P., \& Collett, T. S. (2010). Image matching during ant navigation occurs through saccade-like body turns controlled by learnt visual features. Proceedings of the National Academy of Sciences USA, 107, 16348-16353 doi.org/10.1073/pnas. 1006021107

## PMid:20805481 PMCid:294132

Lohmann, K. J., Lohmann, C. M. F., Ehrhart, L. M., Bagley, D. A., \& Swing, T. (2004). Geomagnetic map used in sea-turtle navigation. Nature, 428, 909-910. doi.org/10.1038/428909a

PMid:15118716
Macquart, D., Garnier, L., Combe, M., \& Beugnon, G. (2006). Ant navigation en route to the goal: signature routes facilitate way-finding of Gigantiops destructor. Journal of Comparative Physiology A, 192, 221234. doi.org/10.1007/s00359-005-0064-7 PMid:16240146
McNaughton, B. L., Battaglia, F. P., Jensen, O., Moser, E. I., \& Moser, M.-B. (2006). Path integration and the neural basis of the 'cognitive map'. Nature Reviews Neuroscience, 7, 663-678. doi.org/10.1038/nrn1932 PMid:16858394
Menzel, R., Greggers, U., Smith, A., Berger, S., Brandt, R., Brunke, S., et al. (2005). Honeybees navigate according to a map-like spatial memory. Proceedings of the National Academy of Sciences USA, 102, 3040 3045. doi.org/10.1073/pnas. 0408550102

PMid:15710880 PMCid:549458
Merkle, T., Knaden, M., \& Wehner, R. (2006). Uncertainty about nest position influences systematic search strategies in desert ants. Journal of Experimental Biology, 209, 3545-3549
doi.org/10.1242/jeb.02395 PMid:16943494
Merkle, T., \& Wehner, R. (2009). How flexible is the systematic search behaviour of desert ants? Animal Behaviour, 77, 1051-1056.
doi.org/10.1016/j.anbehav.2009.01.006
Merkle, T., \& Wehner, R. (2010). Desert ants use foraging distance to adapt the nest search to the uncertainty of the path integrator. Behavioral Ecology, 21, 349-355. doi.org/10.1093/beheco/arp197
Milner, A. D., \& Goodale, M. A. (1995). The visual brain in action. New York, NY: Oxford University Press.
Mishkin, M., Ungerleider, L. G., \& Macko, K. A (1983). Object vision and spatial vision: Two cortical pathways. TRENDS in Neurosciences, 6, 414-417. doi.org/10.1016/0166-2236(83)90190-X
Mittelstaedt, M.-L., \& Mittelstaedt, H. (1980). Homing by path integration in a mammal. Naturwissenschaften, 67,566. doi.org/10.1007/BF00450672 Möller, R. (2001). Do insects use templates or parameters for landmark navigation? Journal of Theoretical Biology, 210, 33-45. doi.org/10.1006/jtbi.2001.2295

PMid:11343429
Möller, R. (2002). Insects could exploit UV-green contrast for landmark navigation. Journal of Theoretical Biology, 214, 619-631. doi.org/10.1006/jtbi.2001.2484 PMid:11851371
Moreau, C. S., Bell, C. D., Vila, R., Archibald, S. B., \& Pierce, N. E. (2006). Phylogeny of the ants: Diversification in the age of angiosperms. Science, 312, 101-104.
doi.org/10.1126/science. 1124891

## PMid:16601190

Morris, R. G. M. (1981). Spatial localization does not require the presence of local cues. Learning and Motivation, 12, 239-260.
doi.org/10.1016/0023-9690(81)90020-5
Morris, R. G. M., Garrud, P., Rawlins, J. N. P., \& O'Keefe, J. (1982). Place navigation impaired in rats with hippocampal lesions. Nature, 297, 681-683. doi.org/10.1038/297681a0 PMid:7088155
Müller, M., \& Wehner, R. (1994). The hidden spiral: systematic search and path integration in desert ants, Cataglyphis fortis. Journal of Comparative Physiology A, 175, 525-530
Müller, M., \& Wehner, R. (2010). Path integration provides a scaffold for landmark learning in desert ants. Current Biology, 20, 1368-1371.
doi.org/10.1016/j.cub.2010.06.035 PMid:20619653 Muser, B., Sommer, S., Wolf, H., \& Wehner, R. (2005) Foraging ecology of the thermophilic Australian desert ant, Melophorus bagoti. Australian Journal of Zoology, 53, 301-311. doi.org/10.1071/ZO05023
Narendra, A. (2007a). Homing strategies of the Australian desert ant Melophorus bagoti I. Proportional path integration takes the ant half-way home. Journal of Experimental Biology, 210, 1798-1803.
doi.org/10.1242/jeb.02768 PMid:17488943
Narendra, A. (2007b). Homing strategies of the Australian desert ant Melophorus bagoti II. Interaction of the path integrator with visual cue information. Journal of Experimental Biology, 210, 1804-1812. doi.org/10.1242/jeb. 02769 PMid:17488944 doi.org/10.1242/jeb. 02791
Narendra, A., Cheng, K., Sulikowski, D., \& Wehner R. (2008). Search strategies of ants in landmark-rich habitats. Journal of Comparative Physiology A, 194 929-938. doi.org/10.1007/s00359-008-0365-8 PMid:18781312
Narendra, A., Cheng, K., \& Wehner, R. (2007). Acquiring, retaining and integrating memories of the out-
bound distance in the Australian desert ant Melophorus bagoti. Journal of Experimental Biology, 210, 570-577 doi.org/10.1242/jeb. 02678
PMid:17267642
Narendra, A., Si, A., Sulikowski, D., \& Cheng, K. (2007). Learning, retention and coding of nest-associated visual cues by the Australian desert ant, Melophorus bagoti. Behavioral Ecology and Sociobiology, 61, 1543-1553. doi.org/10.1007/s00265-007-0386-2 Nehmzow, U. (2008). Emergent cognitive mappings in mobile robots through self-organisation. In M. E. Jefferies \& W.-K. Yeap (Eds.), Robotics and cognitive approaches to spatial mapping (pp. 83-104). Berlin, Heidelberg, New York: Springer. doi.org/10.1007/978-3-540-75388-9 6
Nicholson, D. J., Judd, P. D., Cartwright, B. A., \& Collett, T. S. (1999). Learning walks and landmark guidance in wood ants (Formica rufa). Journal of Experimental Biology, 202, 1831-1838. PMid: 10359685
O'Keefe, J. (1976). Place units in the hippocampus of the freely moving rat. Experimental Neurology, 51 78-109. doi.org/10.1016/0014-4886(76)90055-8
O'Keefe, J., \& Dostrovsky, J. (1971). The hippocampus as a spatial map. Preliminary evidence from unit activity in the freely-moving rat. Brain Research, 34, 171-175. doi.org/10.1016/0006-8993(71)90358-1
O'Keefe, J., \& Nadel, L. (1978). The hippocampus as a cognitive map. Oxford: Clarendon Press.
Pahl, M., Zhu, H., Pix, W., Tautz, J., \& Zhang, S. (2007). Circadian timed episodic-like memory - a bee knows what to do when, and also where. Journal of Experimental Biology, 210, 3669-3567.
doi.org/10.1242/jeb.005488 PMid:17921157
Philippides, A., Baddeley, B., Cheng, K., \& Graham, P. (2011). How might ants use panoramic views for route navigation? Journal of Experimental Biology, 214, 445-451. PMid:21228203
Prabhu, C., \& Cheng, K. (2008). One day is all it takes: Circadian modulation of the retrieval of colour memories in honeybees. Behavioral Ecology and Sociobiology, 63, 11-22. doi.org/10.1007/s00265-008-0631-3 Putman, N. F., Endres, C. S., Lohmann, C. M. F., \& Lohmann, K. J. (2011). Longitude perception and bicoordinate magnetic maps in sea turtles. Current Biology, 21, 463-466. doi.org/10.1016/j.cub.2011.01.057 PMid:21353561
Ronacher, B. (2008). Path integration as the basic navi-
gation mechanism of the desert ant Cataglyphis fortis (Forel, 1902) (Hymenoptera: Formicidae). Myrmecological News, 11, 53-62.
Rosengren, R. (1971). Route fidelity, visual memory and recruitment behaviour in foraging wood ants of the genus Formica (Hymenoptera, Formicidae). Acta Zoologica Fennica, 133, 1-105.
Rossel, S., \& Wehner, R. (1986). Polarization vision in bees. Nature, 323, 128-131. doi.org/10.1038/323128a0 Russell, J. C., McMorland, A. J. C., \& MacKay, J. W B. (2010). Exploratory behaviour of colonizing rats in novel environments. Animal Behaviour, 79, 159 164. doi.org/10.1016/j.anbehav.2009.10.020

Santschi, F. (1911). Sur le mécanisme de l'orientation chez les fourmis [On mechanisms of orientation in ants]. Revue Suisse de Zoologie, 19, 303-338.
Santschi, F. (1913). Comment s'orientent les fourmis [How ants orient]. Revue Suisse de Zoologie, 21, 347-426.
Schultheiss, P., \& Cheng, K. (2011). Finding the nest: inbound searching behaviour in the Australian desert ant Melophorus bagoti. Animal Behaviour, 81, 10311038 doi.org/10.1016/i anbehav201102008
Schultheiss, P., Schwarz, S., \& Wystrach, A. (2010) Nest relocation and colony founding in the Australian desert ant, Melophorus bagoti Lubbock (Hymenoptera: Formicidae). Psyche, 2010,
doi: 10.1155/2010/435838.
doi.org/10.1155/2010/435838
Schwarz, S., \& Cheng, K. (2010). Visual associative learning in two desert ant species. Behavioral Ecology and Sociobiology, 64, 2033-2041.
doi.org/10.1007/s00265-010-1016-y
Schwarz, S., Narendra, A., \& Zeil, J. (2011). The properties of the visual system in the Australian desert ant Melophorus bagoti. Arthropod Structure \& Development, 40, 128-134. doi.org/10.1016/j.asd.2010.10.003 Shettleworth, S. J. (2010). Cognition, evolution, and behavior, second edition. New York: Oxford University Press.
Shettleworth, S. J., \& Sutton, J. E. (2005). Multiple systems of spatial learning: Dead reckoning and beacon homing in rats. Journal of Experimental Psychology: Animal Behavior Processes, 31, 125-141.
doi.org/10.1037/0097-7403.31.2.125 PMid:15839771 Sheynikhovich, D., Chavarriaga, R., Strösslin, T., Arleo, A., \& Gerstner, W. (2009). Is there a geometric module for spatial orientation? Insights from a rodent
navigational model. Psychological Review, 116, 540 566. doi.org/10.1037/a0016170 PMid:19618986 Sommer, S., von Beeren, C., \& Wehner, R. (2008). Multiroute memories in desert ants. Proceedings of the National Academy of Sciences USA, 105, 317-322. doi.org/10.1073/pnas. 0710157104
PMid:18160534 PMCid:2224209
Srinivasan, M. V., Zhang, S., Altwein, M., \& Tautz, J. (2000). Honeybee navigation: nature and calibration of the "odometer". Science, 287, 757-920.
Srinivasan, M. V., Zhang, S. W., \& Bidwell, N. J. (1997). Visually mediated odometry in honeybees. Journal of Experimental Biology, 200, 2513-2522. PMid:9320443
Stürzl, W., \& Zeil, J. (2007). Depth, contrast and viewbased homing in outdoor scenes. Biological Cybernetics, 96, 519-531.
doi.org/10.1007/s00422-007-0147-3 PMid:17443340 Tolman, E. C. (1948). Cognitive maps in rats and men. Psychological Review, 55, 189-208.
doi.org/10.1037/h0061626 PMid:18870876
Towne, W. F. (2008). Honeybees can learn the relationship between the solar ephemeris and a newly-experienced landscape. Journal of Experimental Biology, 211,3737-3743. doi.org/10.1242/jeb.003640 PMid:19011214
Towne, W. F., \& Moscrip, H. (2008). The connection between landscapes and the solar ephemeris in honeybees. Journal of Experimental Biology, 211, 37292736. doi.org/10.1242/jeb. 003640 PMid: 19011214 doi.org/10.1242/jeb.022970 PMid:19011213
von Frisch, K. (1967). The dance language and orientation of bees. Cambridge, MA: Belknap.
von Frisch, K., \& Lindauer, M. (1954). Himmel und Erde in Konkurrenz bei der Orientierung der Bienen [Sky and Earth in competition in the orientation of bees]. Naturwissenschaften, 41, 245-253.
doi.org/10.1007/BF00634944
Wahl, O. (1932). Neue Untersuchungen über das Zeitgedächtnis der Bienen [New research on time memory in bees]. Zeitschrift für Vergleichende Physiologie, 16, 529-589.
Wehner, R. (1994). The polarization-vision project: championing organismic biology. Fortschritte der Zoologie, 39, 103-143.
Wehner, R. (2003). Desert ant navigation: how miniature brains solve complex tasks. Journal of Comparative Physiology A, 189, 579-588.
doi.org/10.1007/s00359-003-0431-1 PMid:12879352 Wehner, R. (2008). The architecture of the desert ant's navigational toolkit (Hymenoptera: Formicidae). Myrmecological News, 12, 85-96.
Wehner, R., Boyer, M., Loertscher, F., Sommer, S., \& Menzi, U. (2006). Ant navigation: One-way routes rather than maps. Current Biology, 16, 75-79.
doi.org/10.1016/j.cub.2005.11.035
PMid:16401425
Wehner, R., Michel, B., \& Antonsen, P. (1996). Visual navigation in insects: coupling of egocentric and geocentric information. Journal of Experimental Biology, 199, 129-140. PMid:9317483
Wehner, R., \& Müller, M. (2006). The significance of direct sunlight and polarized skylight in the ant's celestial system of navigation. Proceedings of the National Academy of Sciences USA, 103, 12575-12579. doi.org/10.1073/pnas. 0604430103
PMid:16888039 PMCid:1567920
Wehner, R., \& Müller, M. (2010). Piloting in desert ants: pinpointing the goal by discrete landmarks. Journal of Experimental Biology, 213, 4174-4179. doi.org/10.1242/jeb.050674 PMid:21112998
Wehner, R., \& Räber, F. (1979). Visual spatial memory in desert ants, genus Cataglyphis (Formicidae, Hymenoptera). Experientia, 35, 1569-1571.
doi.org/10.1007/BF01953197
Wehner, R., \& Srinivasan, M. V. (1981). Searching behaviour of desert ants, genus Cataglyphis (Formicidae, Hymenoptera). Journal of Comparative Physiology A, 142, 315-338. doi.org/10.1007/BF00605445 Wehner, R., \& Srinivasan, M. V. (2003). Path integration in insects. In K. J. Jeffery (Ed.), The neurobiology of spatial behaviour (pp. 9-30). Oxford: Oxford University Press.
Wehner, R., \& Wehner, S. (2011). Parallel evolution of thermophilia: daily and seasonal foraging patterns of heat-adapted desert ants: Cataglyphis and Ocymyrmex species. Physiological Entomology, 36, 271-281. doi.org/10.1111/j.1365-3032.2011.00795.x
Wittlinger, M., Wehner, R., \& Wolf, H. (2006). The ant odometer: Stepping on stilts and stumps. Science, 312, 1965-1967. doi.org/10.1126/science. 1126912 PMid:16809544
Wittlinger, M., Wehner, R., \& Wolf, H. (2007). The desert ant odometer: a stride integrator that accounts for stride length and walking speed. Journal of Experimental Biology, 210, 198-207.
doi.org/10.1242/jeb.02657 PMid:17210957
Wystrach, A., Beugnon, G., \& Cheng, K. (2011). Landmarks or panoramas: what do navigating ants attend to for guidance? Frontiers in Zoology, 8, 21.
doi.org/10.1186/1742-9994-8-21
PMid:21871114 PMCid:3177867
Wystrach, A., Beugnon, G., \& Cheng, K. (2012). Ants might use different view-matching strategies on and off the route. Journal of Experimental Biology, 215, 44-55. doi.org/10.1242/jeb.059584
PMid:22162852
Wystrach, A., Cheng, K., Sosa, S., \& Beugnon, G. (2011). Geometry, features, and panoramic views: Ants in rectangular arenas. Journal of Experimental Psychology: Animal Behavior Processes, 37, 420435. doi.org/10.1037/a0023886

PMid:21604907
Wystrach, A., Schwarz, S., Schultheiss, P., Beugnon, G., \& Cheng, K. (2011). Views, landmarks, and routes: how do desert ants negotiate an obstacle course? Journal of Comparative Physiology A, 197, 167-179. doi.org/10.1007/s00359-010-0597-2 PMid:20972570
Zeil, J., Hofmann, M. I., \& Chahl, J. S. (2003). Catchment areas of panoramic snapshots in outdoor scenes. Journal of the Optical Society of America, 20, 450469. doi.org/10.1364/JOSAA.20.000450

Ziegler, P., \& Wehner, R. (1997). Time-courses of memory decay in vector-based and landmark-based systems of navigation in desert ants, Cataglyphis fortis. Journal of Comparative Physiology A, 181, 1320. doi.org/10.1007/s003590050088

