Ant antennae: are they sites for magnetoreception?

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Migration of the Pachycondyla marginata ant is significantly oriented at 13° with respect to the geomagnetic north–south axis. On the basis of previous magnetic measurements of individual parts of the body (antennae, head, thorax and abdomen), the antennae were suggested to host a magnetoreceptor. In order to identify Fe³⁺/Fe²⁺ sites in antennae tissue, we used light microscopy on Prussian/Turnbull’s blue-stained tissue. Further analysis using transmission electron microscopy imaging and diffraction, combined with elemental analysis, revealed the presence of ultra-fine-grained crystals (20–100 nm) of magnetite/maghaemite (Fe₃O₄/γ-Fe₂O₃), haematite (α-Fe₂O₃), goethite (α-FeOOH) besides (alumo)silicates and Fe/Ti/O compounds in different parts of the antennae, that is, in the joints between the third segment/pedicel, pedicel/scape/thead, respectively. The presence of (alumo)silicates and Fe/Ti/O compounds suggests that most, if not all, of the minerals in the tissue are incorporated soil particles rather than biomineralized by the ants. However, as the particles were observed within the tissue, they do not represent contamination. The amount of magnetic material associated with Johnston’s organ and other joints appears to be sufficient to produce a magnetic-field-modulated mechanosensory output, which may therefore underlie the magnetic sense of the migratory ant.

Keywords: ant antennae; iron oxides; Johnston’s organ; magnetoreception

1. INTRODUCTION

Geomagnetic sensitivity and magnetic orientation behaviour have been intensively studied in a range of animals across all major groups (Wiltschko & Wiltschko 1995; Gould 2008; Lohmann et al. 2008), but the mechanisms by which animals can perceive the geomagnetic field and transduce it into a nerve signal remain to be identified. Current research into magnetoreception is driven mainly by two hypotheses that rely on two fundamentally different but not mutually exclusive physical principles. The radical-pair hypothesis assumes a special class of molecules that undergo magnetically anisotropic chemical reactions as the basis for the magnetic compass sense (Ritz et al. 2000, 2004; Maeda et al. 2008). Migratory birds, for example, appear to have the radical-pair mechanism for the inclination compass and additionally a magnetite-based system for the navigational map (Wiltschko et al. 2008). The magnetite hypothesis postulates the involvement of ferrimagnetic material (e.g. biogenic magnetite) coupled to mechanosensitive structures so as to transmit the information on the geomagnetic field in the form of a torque or force into the nervous system (Kirschvink & Gould 1981; Sherbakov & Winklhofer 1999; Davila et al. 2003, 2005; Ferreira et al. 2005). The first step towards confirming the magnetite hypothesis is to demonstrate the presence of magnetic nanoparticles in or around innervated structures in animal tissues (Diebel et al. 2000; Fleissner et al. 2003).

The discovery of magnetotactic bacteria (Blakemore 1975) defined a milestone for the magnetite hypothesis. Intracellular chains of biomineralized single-domain (SD) magnetite crystals arranged parallel to the motility axis of these micro-organisms make the bacteria orient themselves passively along the geomagnetic field lines, taking them to nutrient-rich and oxygen-poor environments.

Magnetic particles were found in specific tissues or in extracts of body parts of animals (O’Leary et al. 1981; Vilches-Troya et al. 1984; Mann et al. 1988; Walker et al. 1997; Diebel et al. 2000; Hanzlik et al. 2000), but only in the rainbow trout Oncorhynchus mykiss (Walker et al. 1997) and the homing pigeon (Hanzlik et al. 2000; Fleissner et al. 2003) the magnetite found

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seems to have a direct neural connection. Behavioural studies with social insects such as termites, bees and ants have shown orientational and navigational responses to the geomagnetic field (Wiltschko & Wiltschko 1995; Vacha 1997), and a magnetoreceptor system based on superparamagnetic (SPM) magnetite particles in the abdomen was proposed for honeybees (Gould et al. 1978; Hsu et al. 2007).

Most of the ant species are known primarily for being ‘chemical’ animals. Pheromones, the main mediator of communication in these animals, are used by a forager to transfer information to the other workers about food sources or to warn the ants about an imminent attack. Landmarks and polarized light are important for homing. The influence of the geomagnetic field in ant species was reported for the species Formica pratensis (Çamlitepe et al. 2005), Solenopsis invicta (Anderson & Vander Meer 1993), Formica rufa (Çamlitepe & Stradling 1995) and Atta colombica (Rivers & Strygle 2008).

Migration of the ant Pachycondyla marginata is significantly oriented at a 13° angle with respect to the geomagnetic north–south axis (Acosta-Avalos et al. 2001). Magnetic measurements on individual parts of the body (head, antenna, abdomen and thorax) showed that the strongest magnetic signal comes from the antennae (Wajnberg et al. 2004). In addition, behavioural experiments carried out with the Myrmica ruginodis ant showed that the antennae respond to magnetic fields with the pedicel being the most influenced part (Vowles 1954). Based on these results and considering the still-emerging knowledge on the magnetic material in animals and the preliminary magnetic data on the P. marginata ant (Acosta-Avalos et al. 1999; Wajnberg et al. 2000, 2004), we used light microscopy (LM) and transmission electron microscopy (TEM) in this study to identify a possible magnetic sensor in the antennae, a promising sensory organ for magnetoreception.

2. MATERIAL AND METHODS

Pachycondyla marginata workers were collected in the act of foraging in the region of Campinas, São Paulo, Brazil, in September 2006 (Sep06) and May 2007 (May07). The animals were transported alive in boxes containing soil from the nest and its neighbourhood (Sep06) and free of soil (May07). Some of the ants died and were separated only for extraction procedures. For electron microscopic measurements, alive ants were kept in the freezer until no movement was observed. The ants were divided into head with antenna, thorax and abdomens, put separately in ethanol 70 per cent containing soil from the nest and its neighbourhood (May07). The animals were transported alive in boxes for magnetic extractions, antennae and heads of 47 (Sep06) and 50 (May07) individuals were separated from each other, and the procedure in Acosta-Avalos et al. (1999) was followed. Ant body parts were macerated in the presence of 5 per cent sodium hypochlorite (NaOCl) solution (Reagen, Brazil), and the cuticle was removed. Approximately equal amounts of each sample were placed in 1.5 ml Eppendorf tubes, more NaOCl solution was added to half fill the tubes and they were left overnight at 4°C. The tubes were centrifuged at 15 700g for 8 min to pellet the insoluble material. The supernatant consisting of dissolved organic matter was discarded, and a further 0.75 ml of NaOCl solution was added to each pellet. After resuspending the pellets both manually and using ultrasonic vibration for 15 min, the material was subjected to magnetic concentration by placing a strong (Sm-Co) magnet on the lateral wall (not the bottom) of the tubes for 10 min. The resultant red/orange pellet was partially removed with the supernatant, and successive magnetic concentrations were performed to reduce the pellet to a minimum. The same centrifugation, sonication and magnetic concentration procedures were applied to two soil samples. The magnetic concentration of soil resulted in a black concentrated material on the wall of the Eppendorf tube, but this process did not remove all magnetic content of the pellet. The magnetic concentrate and the resuspended pellet, called soil solution, were kept.

After several NaOCl centrifugation steps, chloroform was used to remove fat, with washing procedures similar to the one described earlier. This process was repeated until most of the organic matter and fat had been digested and the orange precipitate did not change its colour. The pellet was washed in distilled water and precipitated by centrifugation. Just before the measurement, the material was sonicated and again magnetically concentrated, and the resulting material of each part on the wall (not observed visually) was dropped onto a TEM grid covered with a carbon film. In the antennae sample, even the pellet was not observed visually. Therefore, the solution was stirred for several minutes with a magnetic finger and the adhering drop was air-dried on a TEM grid. All body parts from Sep06 samples were measured, while only antennae from May07 samples were analysed. Bright field images and selected area electron diffraction (SAED) patterns were taken at 100 kV with a transmission electron microscope Jeol JEM100 CX.

2.2. Turnbull’s/Prussian blue reaction and LM

To locate regions containing iron particles focusing on magnetite, Prussian blue (PB for FeIII) and Turnbull’s blue (TB for FeII) reagents were applied for 10 min to serial histological sections (10 μm thickness, longitudinal (L) or transversal (T) to the antennae) of half heads with antennae or only antennae (total: 8). In the presence of HCl, FeIII and FeII react with potassium ferrocyanide and ferricyanide, respectively, to yield the dark blue ferric ferrocyanide:

\[
PB : \quad K_4[Fe^{III}(CN)_{6}] + Fe^{III}Cl_3 \\
\leftrightarrow KFe^{III}Fe^{II}(CN)_6 + 3KCl,
\]

\[
TB : \quad K_3[Fe^{III}(CN)_{6}] + Fe^{II}Cl_2 \\
\leftrightarrow KFe^{II}Fe^{III}(CN)_6 + 2KCl.
\]
The samples were immediately inspected and, when possible, the presence of Fe regions in consecutive slices of the same sample or in the same region of another sample was verified to discard the iron of contaminant soil particles. Magnetite is the iron oxide most frequently found in animals and, owing to its magnetic properties, it is also desirable from the point of view of magnetoreception. Stoichiometric magnetite consists of one-third ferrous and two-thirds ferric iron, but since ferric iron also occurs in iron storage proteins, we consider the ferrous iron reaction (TB) to be more appropriate for the detection of magnetite. The PB was used at the same time as a control.

2.3. Thin sectioning

Ant antennae are elbow-shaped and divided into segments: the scape, the longest one connected to the head, followed by the pedicel and then the other segments that compose the flagellum (figure 1). Before cutting, the flagellum was aligned with the scape providing only one cutting direction, transverse or longitudinal through the whole antenna. A longitudinal cut was performed in the frontal part of the head to facilitate diffusion of the fixatives. Heads with antennae of samples Sep06 and May07 were cut in two or three positions, respectively, as shown in figure 1.

Sep06 sample parts were fixed in cacodylate-buffered 4 per cent paraformaldehyde and 2.5 per cent glutaraldehyde (Karnovsky’s fixative) solution for 48 h, while May07 samples were fixed in cacodylate-buffered 2.5 per cent glutaraldehyde solution overnight. Both samples were post-fixed in cacodylate-buffered 1 per cent osmium tetroxide solution for 40 min and 1 h, respectively. The May07 samples were embedded in 2.5 per cent low melting point agarose. Samples were then dehydrated with ethanol, plastic-embedded in SPURR and ultrathin sectioned to 100–200 nm. Selected area electron diffraction patterns and bright field images were taken from one Sep06 sample (third segment to head) and two May07 samples (third segment to scape) at 100 kV with a Jeol 100 CX or at 120 kV with a Jeol 2000 FX equipped with an EDAX Genesis system for X-ray energy dispersive spectroscopy (EDX).

3. RESULTS

The collected soil is red/orange and the magnetic extraction procedure resulted in a black concentrated material on the lateral wall of the tube, close to the magnet position, which appeared as agglomerated particles on TEM (data not shown). Figure 2a shows a TEM micrograph of the sample called soil solution in the magnetic extraction section. Energy dispersive spectroscopy analysis of the particles in figure 2a revealed the presence of Fe/O, Si/Al/O, Fe/Ti/O and Ti/O particles. Clusters of 14 μm and particles from <200 nm size up to 6 μm were observed.

An orange precipitate appeared at the bottom of the tube in the antenna, head, thorax and abdomen magnetic extractions (Sep06), with the abdomen having the highest amount and the antenna the lowest. The amount of orange precipitate was strongly reduced in May07 samples, and was not observed in the antenna extracts. TEM micrographs of magnetically concentrated particles of head, abdomen and thorax were obtained after the washing procedure, and a thorax image is shown, as an example, in figure 2b. Clusters of 14 μm and particles from <100 nm size up to 7 μm were observed. Energy dispersive spectroscopy measurements showed that these particles are composed of Fe/O, Si/Al/O, Ti/O and Fe/Ti/O. Powder diffraction patterns of these particles and of those in the soil solution were taken, and their calculated d-spacings are compared with magnetite in table 1.

The d-spacings of the soil solution are common to many Al/Si/O compounds and Al, Si, O compositions with other elements such as Mn, Na and Ca (American Mineralogist Crystal Structure Database), while the body extract d-spacings indicate the presence of Al/Si/O and iron-containing particles.

Compared with the other body part extracts, a large amount of 50 nm Fe/O particles (figure 3a) and a significant decrease in the amount of Si/Al/O particles were observed in the antenna magnetic extracts of the May07 sample. The 50 nm Fe/O particles displayed electron diffraction patterns, as shown in figure 3b. The calculated d-spacings of 2.52 Å together with the measured angle of 60° between two directions (i.e. the angle between two crystalline planes) were used to identify the crystal as haematite.
As soil contaminants can be present in the magnetic extracts, the biogenic origin of single crystals should be proven by their detection within well-preserved embedded tissue. We therefore observed PB/TB reactions in semi-sections by LM to roughly identify sites of iron compounds. Fe$^{3+}$ and Fe$^{2+}$ sites were only observed by LM in the joint of the scape with the head (data not shown).

Ultrathin sectioning and electron microscopy were used to obtain the precise position within the tissue. Transmission electron microscopy bright field images of ultrathin sections showed iron concentrations in several transversal ultrathin sections through the three main antennal joints: pedicel/third segment, scape/pedicel and head/ scape, not observed in the semi-sections by LM, probably because the regions were too small to be observed by LM.

Micrographs of the transversal ultrathin sections through the third segment/pedicel joint (figure 4a, inset) are shown in figure 4a,b. In figure 4a, an empty knob shows a long sensory process, while the overall EDX spectrum of the particles in figure 4b indicated the presence of Fe, Si, Al and O. The region with particles in figure 4b is magnified in figure 4c. Several diffraction patterns of the iron-containing particles were taken and they indicated the presence of goethite and/or haematite and other iron-containing particles that could not be identified. As an example, the diffraction pattern of the particle indicated by the arrow in figure 4c is shown (figure 4d). The particles were found inside the epicuticular invagination (Masson & Gabouriaux 1973) or cuticular knobs (Tsujiuchi et al. 2007), into which the long sensory process (arrow in figure 4a) (Hallberg 1981) from the scolopidia comes.

Iron oxides, aluminosilicates, silicates and Ti/O particles were also found in transversal ultrathin sections through the pedicel (P)/scape (S) joint (figure 5a, inset). Particles surrounding a cell-like structure were

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Table 1. Comparison of the experimental lattice distances ($d$), in angstroms (Å), of ant body extracts and soil solution to the known patterns of three iron oxides. (The percentages in parentheses are the relative intensities measured from X-ray powder diffraction and are directly related to the probability that each diffraction point appears in any diffraction pattern. X-ray powder diffraction data were obtained from the Joint Committee for Powder Diffraction Standards (JCPDS) database. The corresponding card numbers in this database are 19-629 for magnetite, 13-534 for haematite and 29-713 for goethite.)

<table>
<thead>
<tr>
<th></th>
<th>magnetite, $d$ (Å)</th>
<th>haematite, $d$ (Å)</th>
<th>goethite, $d$ (Å)</th>
<th>soil (solution), $d$ (Å)</th>
<th>thorax (Sep06), $d$ (Å)</th>
<th>head (Sep06), $d$ (Å)</th>
<th>antennae (Sep06), $d$ (Å)</th>
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<tr>
<td>2.53 (100)</td>
<td>2.51</td>
<td>2.58/2.53</td>
<td>2.42</td>
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<td>2.53</td>
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<tr>
<td>2.42 (8)</td>
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<td>2.30</td>
<td>2.32</td>
<td>2.42</td>
<td>2.32</td>
<td></td>
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<tr>
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<td>1.97</td>
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<td>2.07</td>
<td>2.09/2.01/1.92</td>
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<td>1.80/1.77/1.72</td>
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<td>1.62 (30)</td>
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<td>1.64</td>
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Figure 2. Transmission electron microscopy micrograph of (a) soil solution and (b) magnetic extracts from thorax of the *P. marginata* ant.
Figure 3. (a) Transmission electron microscopy micrograph of 50 nm Fe/O particles in antenna magnetic extracts of the May07 sample, identified as haematite by (b) the diffraction pattern of one of the crystals.

Figure 4. Transmission electron microscopy micrographs of transversal ultrathin sections of the third segment/pedicel joint cut through the antenna. (a) A cuticular knob (diameter, 4–5 μm). The white arrow points to a long sensory process (diameter, 1 μm). Inset: SEM of P. marginata antenna, showing the third segment (3rd) and the pedicel (P). (b) Another cuticular knob in which particles were found. (c) Magnified knob region in which haematite and goethite (arrow) crystals together with silicates/alumosilicates were identified. (d) Diffraction pattern of the goethite particle (arrow in c).
observed continuously over approximately 35 µm. The three-dimensional image of this structure could be associated with an ellipsoid with an a semi-axis that varies from 7.5 to 12.5 µm and b semi-axis from 3 to 4 µm (figure 5a). Energy dispersive spectroscopy and SAED data were obtained and allowed us to identify haematite, goethite and magnetite/maghaemite. The arrow in the inset of figure 5b indicates a magnetite/maghaemite crystal.

Iron-containing areas were observed in the scape/head joint region after PB reaction (data not shown), and this result was confirmed through the diffraction patterns and EDX analysis obtained. The EDX data of the particles in this area indicated the presence of silicates/alumosilicates, Fe/O and Fe/Ti/O particles. Micrographs of ultrathin sections of these regions show a large amount of small (5–20 nm) Fe/O particles surrounding large particles (approx. 500 nm to 2 µm) of Ti/Fe/O or Fe/O (data not shown).

4. DISCUSSION

Magnetic measurements (Acosta-Avalos et al. 1999; Wajnberg et al. 2000, 2004) have suggested the presence of biogenic magnetite in the body parts (especially the antenna) of the P. marginata ant. In this work, TEM was used for the first time to localize and characterize iron-containing regions within antennal tissue. In magnetic extracts of head, antennae, thorax and abdomen of P. marginata, pure Fe/O particles were found together with Al/Si/O particles, with the latter forming the major part. But even if magnetite is present in the extracts, an unambiguous attribution to a biogenic or non-biogenic origin is not possible.

Several methods and criteria have been developed to distinguish between biotic/biogenic magnetite crystals and abiotic/non-biogenic ones, based on a wide dataset of cultured, mutant and wild-type bacteria (Guyodo et al. 2006; Kopp & Kirschvink 2008). Magnetite crystals (magnetosomes) synthesized intracellularly in magnetic bacteria possess unique features in terms of size, morphology and mineralogy. An identification method for magnetofossils in soils based on seven properties obtained by rock magnetic, TEM, scanning electron microscopy (SEM) and ferromagnetic resonance measurements (Kopp et al. 2006) was recently proposed. However, all these criteria are based on the characteristics of bacterial magnetosomes. For other organisms, it is important to keep in mind that their crystal features are not necessarily those of bacterial magnetosomes as they are supposed to be involved in the complex magnetoreception mechanism, instead of the passive magnetotactic response. Indeed, cluster arrangements of SPM instead of SD magnetite crystals were found in the upper beak skin of homing pigeons (Hanzlik et al. 2000). Therefore, the task of finding specific characteristics that can be unambiguously ascribed to a biological origin still remains (Fortin & Langley 2005).

Extraneous iron minerals (from the soil) can adhere to the cuticle of all body parts and/or be ingested in the thorax and abdomen. The extraction method cannot guarantee that the extraneous material is completely washed off after cleaning. Therefore, to make sure that the particles are from within the tissue and to clarify their possible physiological context, it is necessary to localize them within the tissue. The search for PB/TB reaction products under the LM is a fast screening method for regions containing Fe inside the tissue. However, as only Fe sites of the order of 1 µm can be detected under the LM, small-scale reaction products may be missing even if present. For example, we detected Fe/O crystals under the TEM, but no blue

![Figure 5](http://rsif.royalsocietypublishing.org/)
stains were visible in the corresponding regions when observed under the LM. Moreover, in poorly fixated/infiltrated samples, a displacement of the blue dots through the reagents from their original positions may occur.

Pure Fe/O (or Fe oxides) particles were detected in different parts of the *P. marginata* antennae: third segment/pedicel, pedicel/ scape and scape/head joint. These three main joints are rich in sensorial structures common to the class Insecta and open up a new aspect in iron oxide function. In the cuticular surface of scape and pedicel, there are two types of mechanoreceptors: sensilla campaniform and chaeticum (J. Ferreira de Oliveira & M. Hanzlík 2008, unpublished data). The first responds to deformations in the cuticle (Ehmer & Gronenberg 1997a,b), while the second has taste and mechanoreceptive functions (Zacharak 1980). Moreover, on two joints, pedicel/ scape and scape/head, some groups of hair plates and also sensilla campaniform were observed. The hair plate is composed of Böhm’s bristles, which is a proprioceptor perceiving large changes in the antennal position (Sane et al. 2007) and also sensilla campaniform were observed.

Localized in the antennal pedicel, Johnston’s organ functions as a mechanosensory organ, perceiving alterations of the flagellum in relation to the pedicel. It thus serves in gravireception, hearing and flight control (Sandeman 1976). This organ is composed of sensorial units called scolopidia, which are distributed symmetrically inside the pedicel. Different functions are attributed to this organ, and if the iron oxide crystals present in the cuticular knobs are somehow linked to this receptor, a magnetic function might be expected. The Johnston’s organ in flies (Diptera) has been shown to be responsible for gravireception (Kamikouchi et al. 2009), but it is generally assumed that it lacks statoliths (Étsock 2009). Our data show that the Johnston’s organ in Hymenoptera has dense mineral particles incorporated, which may well serve as statoliths in gravireception. The observed inhomogeneous distribution among the knobs is probably deceptive and caused by the unknown particle distribution in the whole three-dimensional structure. Recent papers on fruit fly antennae (Kamikouchi et al. 2009; Yorozu et al. 2009) have shown that different clusters of neurons in this organ preferentially respond to gravity/wind, whereas others preferentially respond to sound. If the functions in the Johnston’s organ are compartmentalized, the presence of particles in all knobs perhaps is not necessary. Vowles (1954) reported that the Johnston’s organ is able to perceive gravity and magnetic fields that work as a stimulus for orientation in *M. ruginodis* and *M. laevinodis* ants.

In the pedicel–scape joint, a cell-like structure surrounded by different kinds of particles including iron-containing ones was found (figure 5a). The whole structure is assumed to be a proprioceptor because of its special location. The unknown function of this structure motivated us to test it as a magnetosensor of the geomagnetic field. The presence of magnetic particles turns the surrounding material (the outward surface) magnetizable and it represents a ferromembrane (Winklhofer 1998). Importantly, the elongated surface loaded with the magnetic particles implies that its magnetic behaviour is anisotropic, that is, it can be magnetized more easily along the long axis of the elongated structure than along the short axis (figure 5a). As a consequence of this magnetic anisotropy, a torque will arise if the ferromembrane is magnetized at an oblique angle with respect to the long axis. The torque tries to rotate the structure into alignment with the external magnetic field. There is no need of an actual mechanical rotation, however, in order for the proprioceptor to detect the torque. The magnetic torque acting on the proprioceptor will bias the mechanical torques (for example, owing to movement of the filaments), and by comparing the output of a magnetically coated proprioceptor with a non-magnetic adjacent one, the magnetic torque can be isolated. This is conceptually similar to the magnetite null detector suggested by Edmonds (1992), who considered the torque balance between hypothetical magnetite-loaded hair cell detectors and non-magnetic ones. The derivation of our theoretical model can be found in appendix A. To provide a quantitative assessment of the magnetic field sensitivity of the ferromembrane, we have to make an assumption about the intrinsic low-field magnetic susceptibility $k$, which depends on the size and concentration of the magnetic particles in the shell as well as on the intrinsic magnetic parameters of the particles. Nanocrystalline haematite particles, as can be observed in figure 3a, are often found to be SPM, for which low-field magnetic susceptibilities may be up to 0.01 G/Oe (Bodker et al. 1994; Raming et al. 2002). For a magnetic shell of average thickness of approximately 2.5 $\mu$m ($\lambda = 1.075$; appendix A) and intrinsic low-field susceptibility of $k = 0.03$, the torque is just enough to balance the thermal energy at room temperature (figure 6). This is a worst-case scenario, which does not involve the possibility of using strongly ferromagnetic compounds such as maghaeite or magnetite (figure 5b inset) as well. A small amount of 5 per cent incorporated with either component would significantly amplify the torque.

In ants, the scape is connected to the head by a ball joint. It can be moved by four extrinsic muscles, two
levator and two depressor muscles, which are located within the head capsule. This gives the scape, and hence the entire antenna, a large freedom of movement. The second antennal segment, the pedicel, is connected to the scape by a hinge joint. It can only be moved in a single plane by a pair of muscles confined within the scape and referred to as intrinsic antennal muscles. All these muscles together control antennal movements, ranging from slow position-maintaining tasks to fast antennal flicks (Ehmer & Gronenberg 1997a,b). Within the head capsule, the Janet’s organ is found. Scolopales and dendrites, which compose this organ, together form an elastic ribbon that is stretched by antennal movement. One end of the scolopale inserts at internal processes at the basis of the scape (Ehmer & Gronenberg 1997a,b). The iron-containing particles found inside the ball joint and surrounding regions suggest that this part can also be involved, together with other structures, in the detection of the Earth’s magnetic field.

The presence of magnetic particles found within the tissue close to mechanosensitive structures in very specific areas along the antenna parts, probably incorporated during the growing process, suggests a possible magnetoreception function.

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APPENDIX A

A.1. Magnetic torque experienced by a ferromembrane in an external magnetic field

To derive an expression for the magnetic torque, we assume that the ferromembrane has homogeneous magnetic susceptibility and that its shape can be approximated by a prolate ellipsoidal shell. The magnetic energy $W$ of a homogeneously magnetized shell confined by two confocal ellipsoids is given by:

$$ W = \frac{1}{2} M H_0 V, \quad (A1) $$

where $M$ is the magnetization, $H_0$ the external field strength (in Oe) and $V$ the volume of the shell, given by $V = \frac{4\pi r^3}{3}(\lambda^2 - 1) - c^3 (1 - e^2)$, where $e^2 = 1 - (a/c)^2$ and $\lambda > 1$ is the ratio of similitude between the inner ellipsoid ($c$, $a$) and the outer ellipsoid ($c_l = \lambda c$, $a_l = \lambda a$). Equation (A1) can be rewritten as

$$ W = \frac{1}{2} (\kappa'_c \cos^2 \theta + \kappa'_l \sin^2 \theta) H_0^2 V, \quad (A2) $$

where $\theta$ is the polar angle of the external magnetic field axis with respect to the long axis of the body and $\kappa'_c$ and $\kappa'_l$ are the apparent susceptibilities along the long and short axes of either ellipsoid, respectively, with

$$ \kappa'_v = \frac{\kappa}{(1 + N_v \kappa)}, \quad (v = a, c), \quad (A3) $$

where $\kappa$ is the intrinsic magnetic susceptibility and $N_v$, the geometric demagnetizing factor along the long axis, is given as

$$ \frac{N_v}{4\pi} = 1 - \frac{e^2}{e^2 - 1} \left( \text{Artanh}(e) - e \right) \quad (A4) $$

(e.g. Osborn 1945; Stoner 1945), and $N_a = N_c = (4\pi - N_v) / 2$. The magnetically coated structure described in the article has a long axis $2c = 35 \mu m$ and short axes $2a \approx 2b \approx 7 \mu m$. For $a/c = 3.5/17.5$ ($e^2 = 0.96$), $N_c = 0.70$ and $N_a = 5.93$. From equation (A2), we obtain the torque as

$$ T = \frac{dW}{d\theta} = \frac{1}{2} (\kappa'_c - \kappa'_l) \sin 2\theta H_0^2 V. \quad (A5) $$

The sin $2\theta$ dependence of $T$ implies an axial field response (figure 7), as can be expected for an SPM system.

REFERENCES


American Mineralogist Crystal Structure Database. See http://rtruff.geo.arizona.edu/AMS/result.php.


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