A magnet attached to the forehead disrupts magnetic compass orientation in a migratory songbird

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**Abstract**

For studies on magnetic compass orientation and navigation performance in small bird species, controlled experiments with orientation cages inside an electromagnetic coil system are the most prominent methodological paradigm. These are, however, not applicable when studying larger bird species and/or orientation behaviour during free flight. For this, researchers have followed a very different approach. By attaching small magnets to birds, they intended to deprive them of access to meaningful magnetic information. Unfortunately, results from studies using this approach appear rather inconsistent. As these are based on experiments with birds under free flight conditions, which usually do not allow exclusion of other potential orientation cues, an assessment of the overall efficacy of this approach is difficult to conduct.

Here, we directly test the efficacy of small magnets for temporarily disrupting magnetic compass orientation in small migratory songbirds using orientation cages under controlled experimental conditions. We found that birds which have access to the Earth’s magnetic field as their sole orientation cue show a general orientation towards their seasonally appropriate migratory direction. When carrying magnets on their forehead under these conditions, the same birds become disoriented. However, under changed conditions that allow birds access to other (i.e. celestial) orientation cues, any disruptive effect of the magnets they carry appears obscured.

Our results provide clear evidence for the efficacy of the magnet approach for temporarily disrupting magnetic compass orientation in birds, but also reveal its limitations for application in experiments under free flight conditions.

**Introduction**

Birds are amongst the most mobile animals, with many species ranging over thousands of kilometres and between continents during their year-round movements. Such a mobile lifestyle requires the ability to orient and navigate reliably over long distances using positional (i.e. map) and directional (i.e. compass) information derived from environmental cues (e.g. Kramer 1953, 1957). There is evidence for the use of at least three different compass systems for choosing and maintaining specific directions in birds (Chernetsov 2017). Two of the three systems are based on celestial cues: the time-dependent sun compass, which requires the birds’ internal clock as reference (e.g. Kramer 1949; Schmidt-Koenig 1958) and the time-independent star compass (e.g. Emlen 1967a, b; Mouritsen and Larsen 2001; Pakhomov et al. 2017). The magnetic compass which uses Earth’s magnetic field represents the third system (e.g. Merkel & Wiltschko 1965; Wiltschko 1968).

Since its first description in the 1960s, magnetic compass orientation in birds has received particular attention in many subsequent studies. It was shown that birds use the inclination (dip angle) of Earth’s magnetic field rather than its polarity for orientation (Wiltschko & Wiltschko 1972). Further, there is experimental evidence suggesting that in birds, magnetic compass orientation depends on the wavelengths and the intensity of the available light (Wiltschko et al. 1993; Wiltschko and Wiltschko 1995b; Muheim et al. 2002). The sensory basis for magnetic compass orientation, however, is still not fully resolved and subject of intensive research (e.g. Mouritsen 2015; Nordmann et al. 2017; Mouritsen 2018).

A large number of experiments on avian magnetic senses, magnetic compass orientation and navigation have used migratory songbirds as a model for birds in general. If kept in captivity, migratory songbirds typically express increased locomotor activity (wing whirring and/or hopping) during periods normally used for their migratory flights – a behaviour referred to as migratory restlessness (or ‘Zugunruhe’; e.g. Berthold et al. 2000). As migratory restlessness is commonly concentrated towards the birds’ preferred migratory flight direction (Kramer 1949), it can be used as a proxy for their orientation behaviour that is studied by means of small orientation cages (e.g. Emlen funnels; Emlen & Emlen 1966) under controlled experimental conditions. When placed inside an electromagnetic coil system (e.g. Helmholtz coil system), such orientation cages allow an assessment of the effect of specifically altered magnetic fields on birds’ orientation behaviour (e.g. Wiltschko 1968). To date, the combined use of electromagnetic coil systems with orientation cages represents the most prominent methodological paradigm for studies on magnetic compass orientation and navigation performance in small bird species, especially songbirds.

However, when studying magnetic compass orientation and navigation performance in larger bird species and in the context of their behaviours during free flight, any precise alteration of the magnetic field birds experience is challenging, if not impossible. Instead, researchers have tried to assess birds’ use of Earth’s magnetic field for orientation and navigation by depriving them of access to meaningful magnetic information. The most common approach for this is releasing birds with small magnets attached to their bodies (e.g. to the head or back), which was first suggested by physicist C. Maurain in 1926 (see Wiltschko and Wiltschko 1995a for a review). Permanent magnets such as the widely used rare-earth magnets (neodymium and samarium-cobalt magnets) produce strong magnetic fields which exceed Earth’s magnetic field in total magnetic intensity (total field strength) within a radius of at least several centimetres. When attached close to birds’ putative magnetic receptors, such magnets should strongly interfere with Earth’s magnetic field around them, leading to altered resultant magnetic fields in which the resultant vectors are forced to remain within a certain sector relative to the magnets’ alignment when the birds turn or move (e.g. Mouritsen et al. 2003). Such resultant magnetic fields are generally assumed to be uninterpretable and, thus, useless for orientation and navigation purposes (e.g. Mouritsen et al. 2003). Others, however, have questioned the efficacy of this approach (e.g. Wang et al. 2006; Nimpf, Nordmann et al. 2017).

Results from previous studies using magnets for disrupting magnetic compass orientation and navigation in birds appear rather inconsistent. Whereas some studies report a disruptive effect, suggesting the use of magnetic information, others find no such effect (see Wiltschko and Wiltschko 1995a for a review), or effects that varied between repeated experiments and between years (Ranvaud et al. 1991; Moore 1988). Studies in which the birds apparently had access to other orientation cues (e.g. the sun) during the experiment tend to find no effect of the magnets on orientation and overall navigation performance (e.g. Wiltschko and Wiltschko 1995a and references therein; Mouritsen et al. 2003; Bonadonna et al. 2005; Gagliardo et al. 2013; Pollonara et al. 2015; Padget et al. 2017; but see Southern 1972). This is generally inconclusive, as the lack of control over other orientation cues makes it difficult to rule out that Earth’s magnetic field is actually used to obtain positional (i.e. map) and/or directional (i.e. compass) information under natural conditions. Beyond that, the efficacy of the specific treatment cannot be assessed without any preceding tests under controlled experimental conditions that allow no reversion to other orientation cues. Most of the earlier studies used magnets on pigeons (*Columba livia* f. *domestica*) during homing experiments (see Wiltschko and Wiltschko 1995a for review). More recent studies that combined magnets with satellite telemetry to study birds’ behaviour on a large spatial scale mainly focused on the navigation performance of seabirds like tubenoses (Procellariiformes) during foraging and homing flights towards their breeding colonies (e.g. Mouritsen et al. 2003; Bonadonna et al. 2005; Gagliardo et al. 2013; Pollonara et al. 2015; Padget et al. 2017). To date, however, there appears to be only one previous study using magnets on a songbird species (the barn swallow (*Hirundo rustica*) during homing experiments; Bochenski et al. 1960), despite songbirds being the most extensively studied taxon with regard to magnetic compass orientation and navigation. Further, to the best of our knowledge, there is not a single study applying the magnet approach in the context of compass orientation or navigation during seasonal migration.

Here, we investigate the effect of small magnets attached to the foreheads of first-year migratory songbirds, which are known to use Earth’s magnetic field for setting their migratory direction. Birds’ orientation behaviour was repeatedly tested during their first autumn migration season, using orientation cages set up in the field. We expected that birds which have access to the Earth’s magnetic field as their sole orientation cue would be generally oriented towards the seasonally appropriate natural migratory direction. When carrying magnets on their forehead under these circumstances, we expected the same birds to become disoriented due to the disruptive effect on their magnetic compass. When circumstances change and birds are allowed access to other orientation cues (i.e. celestial cues) during the tests, however, we expected any disruptive effect of magnets to be obscured, as they would probably fall back on another compass system (i.e. the star compass) for successful orientation. Our study, for the first time, tests the efficacy of magnets for disrupting magnetic compass orientation in songbirds under controlled experimental conditions.

**Methods**

*Study species and site*

We selected the Eurasian reed warbler (*Acrocephalus scirpaceus*, reed warbler hereafter) as our study species, because it represents a well-established model for studying magnetic compass orientation and navigation in songbirds and birds in general. Reed warblers are common long-distance migrants breeding in reed-lined habitats across a large part of Europe to western Asia and overwintering in sub-Saharan Africa (del Hoyo et al. 2006). Fieldwork for this study took place at Lake Neusiedl (Neusiedler See (Austria) or Fertő (Hungary)), a shallow steppe lake situated at the north-western edge of the Pannonian Basin straddling the Austrian-Hungarian border. Experiments were performed at the Biological Station Lake Neusiedl in Illmitz, Burgenland, south-eastern Austria (47° 46' 08.9"N, 16° 45' 57.2"E).

*Assessment of the natural migratory direction*

We obtained bird ring recovery data of reed warblers ringed at Lake Neusiedl during previous years, both from the Austrian Ornithological Centre (AOC) and the Hungarian Bird Ringing Centre (MME). These data were used to assess the natural migratory direction of reed warblers from the study population during autumn migration. We filtered the data for autumn recoveries (September – November) of birds that were ringed during the breeding and early migratory period (late May – August) of the same year and that were found at a distance of > 250 km from the ringing site. The rationale for the latter was to avoid any directional bias due to a disproportionate number of recoveries from nearby ringing sites.

*Orientation experiments*

Ethical statement:

All applicable international, national and/or institutional guidelines for the care and use of animals were followed. The experiments were conducted in accordance with the national animal welfare legislation of Austria and with permission of the state Burgenland (Abteilung 4 - Ländliche Entwicklung, Agrarwesen und Naturschutz; permit: A4/NN.AB-10216-7-2019). Additionally, the experiments received local ethical approval by the animal welfare ethics review body (AWERB) of Bangor University, where the corresponding authors (F.P. and R.A.H.) were employed during the period of data collection.

Capture and husbandry:

For our two orientation experiments (Experiment 1 in 2019 and Experiment 2 in 2020), we captured reed warblers in the reed beds near the Biological Station during early September, i.e. during the species-specific autumn migration season. The birds were captured with mist nets as part of the Biological Station’s bird monitoring and ringing project. Following capture and standard ringing procedures, we transferred a total of 35 first-year birds (17 birds in 2019 and 18 birds in 2020) to outdoor aviaries near the capture site. Each bird was randomly assigned to one of two roughly equal-sized housing groups (group A: 9 birds in 2019 and 9 in 2020; group B: 8 birds in 2019 and 9 in 2020). During the study periods (12th September – 4th October 2019 and 10th – 22nd September 2020), each housing group was kept together in one aviary equipped with perches, reeds, a water basin and food trays. Water and food (a mixture of live meal worms (*Tenebrio molitor*), dried insects and grated carrots) were provided *ad libitum*. The aviaries were made of non-magnetic materials (wood, polyester nets and insect mesh) and weakly magnetic materials (stainless steel screws) to minimize distortion of the Earth’s magnetic field. Further, the aviaries gave the birds an unobstructed view of the surrounding habitat and access to various orientation cues (e.g., the sun and sun-related cues, the stars and local odours). At the end of the study period, i.e. before the end of the reed warbler’s autumn migration season, all birds were released close to the capture site.

Experimental designs and procedures:

Experiment 1: All 17 birds used for this experiment were repeatedly tested in Emlen funnels (specific type of orientation cage; see the section “*Orientation tests*” for details) while being subjected to three different experimental treatments, i.e. control/no attachment (CON), magnet attachment (MAG) and sham attachment (SHAM), during their autumn migration season in 2019. During the orientation tests, independent of the respective experimental treatment, birds were denied access to orientation cues other than the Earth’s magnetic field (see the section “*Orientation tests*” for details). Each bird underwent four orientation tests during each experimental treatment, for a total of 12 orientation tests on 12 different nights within the test period (15 test nights in total for Experiment 1). All birds were tested in four CON tests first. Subsequently, the treatment order differed between the two housing groups (group A and group B; see above). Birds assigned to group A were subjected to four tests during the MAG treatment, followed by four tests during the SHAM treatment. Birds assigned to group B were subjected to four tests during the SHAM treatment, followed by four tests during the MAG treatment (Fig. 1A). The rationale was to have a time balanced design of the tests to exclude any temporal bias in the data (e.g. an ‘‘endogenously controlled change of migratory direction’’ or ‘‘Zugknick’’; Gwinner and Wiltschko 1978; Liechti et al. 2012).

During CON tests, birds were tested with no attachments whatsoever in order to obtain their seasonally appropriate (control) magnetic compass orientation under exclusion of other orientation cues for subsequent comparisons. The lack of such a control orientation would render any following treatment difficult to interpret. For MAG and SHAM tests, we either fitted a small magnet (MAG) or a small non-magnetic sham attachment (SHAM) to the birds’ forehead (Fig. 2). Both magnets and sham attachments were glued tightly to the short forehead plumage of the birds (Fig. 2) using super glue (LOCTITE® Super Glue Gel). We used small (diameter: 3 mm, height: 2 mm, weight: 0.11 g) disc-shaped neodymium magnets (Supermagnete, S-03-02-N, EAN: 7640155436960, material: NdFeB, residual magnetism BR: 1.37 – 1.42 T, coercive field strength iHC: ≥ 955 kA/m, energy product (BH) max: 358 – 382 kJ/m³). The magnets were attached with their North Pole facing down (towards the bird’s skull)/South Pole facing up (away from the bird’s skull; Fig. 2). As non-magnetic sham attachments, we used disk shaped pieces cut from a brass rod, which resembled the magnets in their dimensions and weight (Fig. 2). To habituate the birds to magnets and sham attachments, they were fitted during the night before the first MAG and SHAM test, respectively. If birds had lost their magnet or sham attachment during the day, these were replaced before the next orientation test (replacement required in 30% of the individual orientation tests).

Experiment 2: The 18 birds used for this experiment were repeatedly tested in Emlen funnels (see the section “*Orientation tests*” for details) while being subjected to two different experimental treatments, i.e. magnet attachment (MAG) and magnet attachment with celestial cues available (MAG + CEL) during their autumn migration season in 2020. During the orientation tests, the birds’ access to orientation cues other than the Earth’s magnetic field depended on the respective experimental treatment (see the section “Orientation tests” for details). Each bird underwent four orientation tests during each experimental treatment, for a total of 8 orientation tests on 8 different nights within the test period (10 test nights in total for Experiment 2). In contrast to Experiment 1, birds from both housing groups (group A and group B) were subjected to the same treatment in the same order during the whole test period. We started with MAG tests during the first test nights, which were then alternated with MAG + CEL tests during subsequent nights, depending on the local weather conditions and the availability of stars/absence of clouds during the test period. Orientation tests were conducted until each bird was tested four times under each of the two experimental treatments (Fig. 1B).

For both MAG and MAG + CEL tests, we attached a small magnet to the birds’ forehead as in Experiment 1. Dimensions and properties of the magnets were identical to the ones previously used (see above). Contrary to Experiment 1, however, the magnets were attached with their North Pole facing up (away from the bird’s skull)/South Pole facing down (towards the bird’s skull; Fig. 2). The rationale was to gather additional information on whether a different orientation (i.e. a reversed polarity) of the attached magnet would result in an altered treatment effect. The attachment took place during the night before the first orientation test and magnets were replaced before the next orientation test if they got lost during the day (replacement required in 19% of the individual orientation tests).

Placement of the magnets:

The sensory basis for magnetic compass orientation is subject of active research and intense scientific debate (e.g. Mouritsen 2015; Nordmann et al. 2017; Mouritsen 2018). Several different models and hypotheses have been put forward to date, proposing very distinct avian magnetic senses that could be used to acquire directional (i.e. compass) information from Earth’s magnetic field for orientation purposes (e.g. Mouritsen 2015; Mouritsen 2018). The radical pair model, a leading concept for the magnetic compass of birds, proposes reversible light-dependent chemical reactions inside the retina of the birds’ eyes as the basis for the avian magnetic sense providing directional information, with the yield of these reactions depending on the alignment of a specific type of molecule (cryptochromes) to the magnetic field (e.g. Ritz et al. 2000; Hore & Mouritsen 2016; Xu et al. 2021). The magnetite model, in contrast, proposes an avian magnetic sense based on biogenic magnetite (Fe3O4) structures located within nerve endings of the trigeminal nerve’s ophthalmic branch V1 in the birds’ upper beak (e.g. Kirschvink and Gould 1981; Falkenberg et al. 2010; Heyers et al. 2010). Apart from these two models, an avian magnetic sense located within the semi-circular canals of the inner ear has been hypothesized, which might be based on electromagnetic induction (Nimpf et al. 2019).

We decided to fit magnets and sham attachments of our MAG, SHAM and MAG + CEL treatments to the foreheads of the birds. This way, magnets were located in close proximity (< 2 cm) to any target tissues holding putative magnetoreceptors, regardless of whether these are actually found in the retina of the birds’ eyes, nerve endings in their upper beak or semi-circular canals of their inner ears. While conducting some example measurements at Bangor University, United Kingdom, we found that the small disc-shaped neodymium magnets we used (see the section “Experimental procedures” for details) increase the total intensity of the local magnetic field by ~83,000 nT at a distance of 2 cm above their South Pole. It is worth mentioning, however, that the magnetic field induced by such neodymium magnets changes strongly and anisotropically as a function of both distance and direction from its centre. The total intensity of Earth’s magnetic field shows a global range of approx. 25,000-65,000 nT (<https://www.ngdc.noaa.gov/geomag/>; accessed 1st June 2021), and amounts to approx. 48,800 nT (estimated for 17th September 2020; <https://www.ngdc.noaa.gov/geomag/calculators/magcalc.shtml>, accessed 1st June 2021) at our study site in Austria. This means that, during our MAG and MAG + CEL treatments, birds’ putative magnetoreceptors located in close proximity (< 2 cm) to the attached magnets were exposed to a magnetic field with the total intensity increased by probably more than 120% of the natural global maximum. Further, putative magnetoreceptors that are situated in more than a single location would most probably be exposed to very different conditions due to the steep and anisotropic magnetic gradients induced by the attached magnet.

Orientation tests:

Orientation tests lasted for 30 min each and were conducted in two successive sessions within the same night (one session for each of the housing groups, i.e., group A and group B), with the first session starting about 90 min after sunset (approximately at the end of the evening twilight period when the sun and sun-related cues are unavailable for orientation purposes). The assignment to the first and second session was alternated between the housing groups to allow a balanced experimental design and avoid any temporal bias in the data. We used modified Emlen funnels – the classical approach for testing migratory orientation in songbirds since its establishment by Emlen & Emlen (1966). The Emlen funnels were made of aluminium (top diameter: 350 mm, bottom diameter: 100 mm, slope 45°). Before the orientation tests, we placed up to nine Emlen funnels on a levelled wooden table (tabletop: ca. 1 m x 1 m) set up under the open sky on a meadow at a distance of approximately 150 m to the Biological Station. We provided no artificial light during the orientation tests. Instead, we surrounded the table with vertical wooden panels (up to ca. 40 cm above the tabletop) to screen off any artificial light sources at the horizon (see Kishkinev, Packmor et al. 2021 fig. S2C).

During all orientation tests of Experiment 1 (CON, MAG and SHAM) and during MAG orientation tests of Experiment 2, the top of each Emlen funnel was covered with a translucent acrylic glass lid that served as light diffuser and prevented the birds from seeing surroundings and gathering information from orientation cues other than the local geomagnetic field. During MAG + CEL orientation tests of Experiment 2, however, the top of each Emlen funnel was covered with insect mesh that gave the birds an unobstructed view of the starry, clear (< 50% cloud cover), moonless night sky and, thus, access to celestial orientation cues (i.e. stars) during the tests.

The directionality of the birds’ activity, i.e. their orientation, was recorded as their scratch marks left on a print film coated with a dried mixture of whitewash and glue. When an Emlen funnel is lined with such a print film, its two ends slightly overlap. During orientation tests, the alignment of the different funnels was alternated, with the overlapping point facing in different cardinal directions (i.e. north and south). The funnel alignment was unknown to the researchers who assessed the birds’ mean directions based on the distribution of the scratch marks from each orientation test. Instead, mean directions were estimated assuming an alignment to the North and later corrected according to the actual alignment from the record. This procedure was meant to avoid any observer bias with regard to directional assessment as well as to avoid providing unintentional visual cues to the birds. Two researchers (F.P. and F.B., F.P. and B.K. or F.P. and C.M.) independently assessed each bird’s mean direction from the distribution of the scratch marks. At least one of the researchers was unaware of the respective experimental treatment during the directional assessment, except for MAG and SHAM tests of Experiment 1 during which both researchers were unaware of the respective experimental treatment as these were conducted during the same period. The resultant direction from the two researchers’ recordings was taken into further analysis. If both researchers considered the scratch marks to be randomly distributed or their assessed directions deviated by more than 30°, a test was considered to be not oriented and, thus, discarded (24% (Experiment 1) and 25% (Experiment 2) of the orientation tests). To ensure comparability with previous studies, only tests with at least 35 scratch marks (a common activity threshold, Wiltschko et al. 1998) and a clear unidirectional orientation were taken into analysis. Tests with less than 35 scratch marks were considered to reveal a lack of migratory activity and, thus, discarded (12% (Experiment 1) and 8% (Experiment 2) of the orientation tests). Birds’ individual directions were used to calculate individual mean directions for each of the three experimental treatments. From individual mean directions, group mean directions were calculated for the different experimental treatments of Experiment 1 and 2.

*Statistics*

Statistical analyses were conducted using the software R version 4.0.4(R Core Team 2021).

We analysed bird ring recovery data to estimate the natural migratory direction of reed warblers from our study population during autumn migration. We used the package ‘geosphere’ (Hijmans et al. 2019) to calculate bearings between the ringing sites (at Lake Neusiedl) and the respective recovery sites (Fig. 3). Then, we tested whether these bearings significantly differed from a uniform distribution using the Rayleigh test of uniformity and calculated their circular mean using the package ‘circular’ (Lund & Agostinelli 2007). The circular mean was adopted as the study population’s approximate natural migratory direction.

To test whether the orientation data obtained during the different experimental treatments of Experiment 1 and 2 significantly differed from a uniform distribution, the Rayleigh test of uniformity was used. Additionally, we assessed the likelihood of the ten models for orientation behaviour described by Schnute and Groot (1992) for the orientation data obtained during each experimental treatment of Experiment 1 and 2 using the model selection procedure implemented in the package ‘CircMLE’ (Fitak & Johnsen 2017). These comprise uniform (M1), unimodal (M2A), symmetric modified unimodal (M2B), modified unimodal (M2C), homogenous symmetric bimodal (M3A), symmetric bimodal (M3B), homogenous axial bimodal (M4A), axial bimodal (M4B), homogenous bimodal (M5A), and bimodal (M5B) orientation models. We compared the models by means of the corrected Akaike information criterion (AICc; Hurvich & Tsai 1989) and the corresponding AICc model weights.

In order to compare orientation data between the experimental treatments during which birds were significantly oriented (according to the Rayleigh test of uniformity), we used the non-parametric Mardia-Watson-Wheeler test implemented in the package ‘circular’ (Lund & Agostinelli 2007). For comparisons of orientation data between the experimental treatments during which birds were not significantly oriented with those during which they were significantly oriented, we followed a bootstrap approach applied by Chernetsov et al. (2017). This approach uses the mean resultant vectors (r-values; a measure of directedness) obtained during the different experimental treatments and compares whether the r-value that derives from a not significantly oriented sample falls within some confidence intervals for another r-value that derives from a significantly oriented sample. In a first step, a random sample of *n* orientation directions is drawn with replacement from the original (significantly oriented) sample of *n* orientation angles obtained during the respective experimental treatment (e.g. *n* = 17 for the control tests) and the corresponding r-value is calculated. This procedure is repeated 100,000 times, each time with a new randomization. In a second step, the resulting 100,000 r-values are ranked in ascending order, with values at the ranks 2500 and 97500, 500 and 99500, and 50 and 99,950 defining the 95%, 99%, and 99.9% confidence limits for the observed r-value of the significantly oriented sample, respectively. If the observed r-value of the not significantly oriented sample is outside these confidence intervals, the significantly oriented sample is more directed with a confidence of > 95%, > 99%, and > 99.9%, respectively.

**Results**

*Natural migratory direction*

Autumn bird ring recoveries of reed warblers ringed at Lake Neusiedl suggested a natural migratory direction towards the SE to SSE (mean direction: α = 152° [all directions are indicated relative to the magnetic North]; Rayleigh test: r = 0.80, *P* < 0.001, *n* = 19; 95% confidence interval of the group mean direction [95% CI mean] = 140° ̶ 170°; Fig. 3).

*Orientation experiments*

Experiment 1: We repeatedly tested a total of 17 first-year reed warblers for their magnetic compass orientation while they were subjected to three different experimental treatments (CON, MAG, SHAM) during the autumn migration season. The circular distributions obtained were best described by unimodal orientation models (Table 1) and birds showed a mean orientation towards the SE to SSE during both CON tests (mean direction (CON): α = 147°; Rayleigh test: r = 0.47, *P* = 0.02, *n* = 17; 95% CI mean = 105° ̶ 187°; Fig. 4A) and SHAM tests (mean direction (SHAM): α = 167°; Rayleigh test: r = 0.65, *P* = 0.002, *n* = 14; 95% CI mean = 136° ̶ 195°; Fig. 4A). The two circular distributions were not found to be statistically distinguishable (Mardia-Watson-Wheeler test: W = 0.56, *P* = 0.76). During MAG tests the birds were randomly oriented (Rayleigh test: r = 0.17, *P* = 0.70, *n* = 12; Fig. 4A) and their circular distribution was best described by the uniform orientation model (Table 1). This apparent lack of a mean orientation was distinguishable from the same birds’ orientation during both CON tests (> 95% confidence: the bootstrapped 95% confidence interval for the r-value from CON tests was 0.21 < r < 0.76, which does not overlap with the r-value of 0.17 obtained during MAG tests) and SHAM tests (> 99.9% confidence: the bootstrapped 99.9% confidence interval for the r-value from SHAM tests was 0.30 < r < 0.93, which does not overlap with the r-value of 0.17 obtained during MAG tests).

Experiment 2: We repeatedly tested a total of 18 first-year reed warblers for their magnetic and celestial compass orientation while they were subjected to two different experimental treatments (MAG, MAG + CEL) during the autumn migration season. During MAG tests the birds were not oriented in a unimodal way (Rayleigh test: r = 0.23, *P* = 0.48, *n* = 15; Fig. 4B) and their circular distribution was best described by either the homogenous symmetric bimodal orientation model or the uniform orientation model (Table 2). The circular distribution obtained during MAG + CEL tests were best described by a unimodal orientation model (Table 2) and birds showed a mean orientation towards the SE to SSE (mean direction (MAG + CEL): α = 146°; Rayleigh test: r = 0.51, *P* = 0.01, *n* = 16; 95% CI mean = 104° ̶ 184°; Fig. 4B). The lack of a unimodal mean orientation during MAG tests was distinguishable from the same birds’ orientation during MAG + CEL tests (> 95% confidence: the bootstrapped 95% confidence interval for the r-value from MAG + CEL was 0.30 < r < 0.76, which does not overlap with the r-value of 0.23 obtained during MAG tests).

**Discussion**

In this study, we show the disruptive effect of small magnets on magnetic compass orientation in a migratory songbird, the reed warbler, under controlled experimental conditions. During CON and SHAM tests, i.e. when Earth’s magnetic field was available as the sole orientation cue, reed warblers were oriented towards the SE to SSE (Fig. 4A), matching the seasonally appropriate, natural, migratory direction of the study population during autumn migration (Fig. 3). During MAG tests, however, when birds had small magnets attached to their foreheads intended to deprive them of gathering meaningful magnetic information, birds appeared disoriented, with the obtained orientation data being best described either by a uniform distribution (Fig. 4A; Table 1) or by symmetric bimodal and uniform distributions (Fig. 4B; Table 2). When provided with an unobstructed view of the starry night sky during MAG + CEL tests, in contrast, the birds were able to orient in the seasonally appropriate, natural, migratory direction towards the SE to SSE, despite having small magnets attached to their foreheads (Fig. 4B). This suggests that the birds can fall back on other orientation cues if they find Earth’ magnetic field to be unusable for determining direction.

Magnetic compass orientation has repeatedly been reported in various songbird species and is now well established for this taxon (e.g., Mouritsen 2015; Chernetsov 2017). Furthermore, there is evidence for songbirds using the Earth’s magnetic field not only as a source of directional (i.e. compass), but also positional (i.e. map) information during the navigation process (e.g., Kishkinev et al. 2013, 2015; Chernetsov et al. 2017; Pakhomov et al. 2018; Kishkinev, Packmor et al. 2021). With regard to other bird taxa that include larger species, however, evidence for the use of Earth’s magnetic field for orientation or navigation appears less coherent. Although, some experimental studies show magnetic compass orientation in homing pigeons (*Columba livia domestica*) (e.g. Keeton 1969, 1971; Walcott & Green 1974; Ioalè 1984, 2000; Visalberghi and Alleva 1979; Gagliardo et al. 2009; Mora and Walker 2012), others criticised these studies and/or failed to replicate the prior results (e.g. Lamotte 1974; Moore 1988; Ranvaud et al. 1991; reviewed in Wiltschko and Wiltschko 1995a, Wallraff 2005). For gulls (Laridae) there is evidence for magnetic compass orientation from displacement experiments with fledglings (e.g., Southern 1973), but the significance of magnetic information for their navigational performance has been questioned based on data from an experimental study tracking long-distance flight behaviour of gulls by means of satellite telemetry (Wikelski et al. 2015). The latter holds true for tubenoses (Procellariiformes) as well, which to the best of our knowledge have not been shown to use Earth’s magnetic field to obtain directional (i.e. compass) or positional (i.e. map) information for orientation and navigation during their offshore flights in any experimental study yet (e.g. Massa et al. 1991; Mouritsen et al. 2003; Benhamou et al. 2003; Bonadonna et al. 2003, 2005; Gagliardo et al. 2013; Pollonara et al. 2015; Padget et al. 2017; Syposz et al. 2020). A recent correlational study, however, suggests that the inclination of Earth’s magnetic field plays a role for finding and selecting breeding colonies during the recruitment phase in a tubenose species (Wynn et al. 2020).

The vast majority of studies that aimed to investigate the use and overall significance of magnetic information for orientation and navigation in “non-passerine” birds and in the context of free flight behaviour used rare-earth magnets in a way comparable to our current study (e.g. Wiltschko and Wiltschko 1995a; Mouritsen et al. 2003). Hence, we expected that our results would be comparable with those of the previous studies and could help to put them into context. Generally, it can be stated that studies which found a disruptive effect of the magnets on birds’ orientation behaviour were usually able to largely exclude orientation cues other than the Earth’s magnetic field during the experiments (i.e. complete overcast conditions). Studies in which the birds apparently had access to other (e.g. celestial) orientation cues (i.e. clear skies), however, show a tendency towards finding no effect of the magnets on the orientation behaviour and overall navigation performance (e.g. Keeton 1969, 1971; Ioalè 1984, 2000; Mouritsen et al. 2003; Bonadonna et al. 2005; Gagliardo et al. 2013; Pollonara et al. 2015; Padget et al. 2017; but see Southern 1972; reviewed in Wiltschko and Wiltschko 1995a; Wallraff 2005). This is in general agreement with the results of the current study. The magnets attached to the birds’ foreheads were found to disrupt their magnetic compass orientation, but the disruptive effect was not detectable when the birds had access to celestial orientation cues, which apparently gave them the opportunity to fall back on another compass system for successful orientation. If orientation cues are accessible, birds may integrate all their different compass systems (magnetic and celestial compasses; Wiltschko and Wiltschko 2001), but for successful orientation they seem to require merely one at a time (e.g. Mouritsen 2015). Impaired access to celestial orientation cues may occur under overcast conditions, and the access to Earth’s magnetic field for orientation and navigation may get severely impacted by both natural (e.g. solar storms; Bianco et al. 2019) and anthropogenic (e.g. electromagnetic noise; Engels et al. 2014) causes that birds must cope with. Hence, independent of any underlying compass system hierarchy and calibration (e.g. Muheim et al. 2006; Liu and Chernetsov 2012; Pakhomov & Chernetsov 2020) birds need their compass integration mechanisms to allow reversion to the other systems, if one becomes temporarily inaccessible or unreliable. This is supported by both the current results and previous studies (e.g. Wiltschko et al. 1987; Mouritsen 1998).

Our study species, the reed warbler, represents a well-established model for migratory songbirds and has repeatedly been shown to use Earth’s magnetic field for orientation and navigation in a migratory context (e.g. Holland 2010; Kishkinev et al. 2015; Chernetsov et al. 2017; Kishkinev, Packmor et al. 2021). Thus, by studying reed warblers in orientation cages under controlled experimental conditions, we are able to provide clear evidence for the general efficacy of the magnet approach for temporarily disrupting magnetic compass orientation in birds. Further, the efficacy of the magnet approach appears independent of the relative alignment of the magnets in our orientation experiments (i.e. North Pole facing down/South pole facing up in Experiment 1 vs. North Pole facing up/South pole facing down in Experiment 2). Studies that applied the magnet approach to investigate the significance of Earth’s magnetic field for birds’ orientation and overall navigation performance under free flight conditions have previously been criticised for the potential inefficacy of the experimental treatment (e.g. Wang et al. 2006), which is comprehensible given that many of them did not find a disruptive effect. With regard to our current results, however, we argue that this general criticism is no longer tenable. Instead, we would like to emphasise that the detectability of a disruptive effect of the magnets appears highly *context dependent*. This is crucial for the interpretation of results from studies following the magnet approach under free flight conditions, which usually do not allow the restriction of birds’ access to other cues for orientation and navigation.

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**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

Conceptualization: F.P., D.K., L.C.Z., T.G., R.A.H.; Methodology: F.P., D.K., L.C.Z., T.G., R.A.H.; Formal analysis: F.P.; Investigation: F.P., F.B., B.K., C.M.; Resources: T.Z., R.A.H.; Data curation: F.P.; Writing – original draft: F.P., R.A.H.; Writing - review & editing: F.P., D.K., L.C.Z., T.G., R.A.H.; Visualization: F.P.; Supervision: R.A.H.; Project administration: F.P., R.A.H.; Funding acquisition: R.A.H.

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**Data availability**

All relevant data will be made available in the Dryad digital repository upon acceptance for publication.

**References**

Benhamou S, Jouventin P, Bonadonna F (2003) Successful homing of magnet-carrying white-chinned petrels released in the open sea. *Animal Behaviour* 65: 729-734.

Berthold P, Fiedler W, Querner U (2000) Die Zugunruhe bei Vögeln — eine Darstellung nach Videoaufnahmen bei Infrarotlichtbeleuchtung. *Journal für Ornithologie* 141: 285-299.

Bianco G, Ilieva M, Åkesson S (2019) Magnetic storms disrupt nocturnal migratory activity in songbirds. *Biology Letters* 15: 20180918.

Bochenski Z, Dylewska M, Gieszczykiewicz J (1960) Homing experiments on birds. Part XI: Experiments with swallows *Hirundo rustica* L. concerning the influence of earth magnetism and partial eclipse of the sun on their orientation. *Zesz Nauk Uniw Jagiellonk* 33: 125-130 (in Polish)

Bonadonna F, Bajzak C, Benhamou S, Igloi K, Jouventin P, Lipp HP, Dell'Omo G (2005) Orientation in the wandering albatross: interfering with magnetic perception does not affect orientation performance. *Proceedings of the Royal Society B* 272: 489-495.

Bonadonna F, Chamaillé-Jammes S, Pinaud D, Weimerskirch H (2003) Magnetic cues: are they important in Black-browed Albatross *Diomedea melanophris* orientation? *Ibis* 145: 152-155.

Chernetsov N (2017) Compass systems. *Journal of Comparative Physiology A* 203: 447-453.

Chernetsov N, Pakhomov A, Kobylkov D, Kishkinev D, Holland RA, Mouritsen H (2017) Migratory Eurasian reed warblers can use magnetic declination to solve the longitude problem. *Current Biology* 27: 2647-2651.e2.

del Hoyo J, Elliott A, Christie DA (2006) The Handbook of the Birds of the World, Vol. 11 Old World Flycatchers to Old World Warblers. Barcelona: Lynx Edicions.

Emlen ST, Emlen JT (1966) A technique for recording rigratory orientation of captive birds. *The Auk* 83: 61-367.

Emlen ST (1967a) Migratory orientation in the Indigo Bunting, *Passerina cyanea*: Part I: evidence for use of celestial cues. *The Auk* 84: 309-342.

Emlen ST (1967b) Migratory orientation in the Indigo Bunting, *Passerina cyanea*: Part II: mechanism of celestial orientation. *The Auk* 84: 463-489.

Falkenberg G, Fleissner G, Schuchardt K, Kuehbacher M, Thalau P, Mouritsen H, Heyers D, Wellenreuther G, Fleissner G (2010) Avian magnetoreception: elaborate iron mineral containing dendrites in the upper beak seem to be a common feature of birds. *PLoS ONE* 5(2): e9231.

Fitak RR, Johnsen S (2017) Bringing the analysis of animal orientation data full circle: model-based approaches with maximum likelihood. *Journal of Experimental Biology* 220: 3878-3882.

Gagliardo A, Bried J, Lambardi P, Luschi P, Wikelski M, Bonadonna F (2013) Oceanic navigation in Cory's shearwaters: evidence for a crucial role of olfactory cues for homing after displacement. *Journal of Experimental Biology* 216: 2798-2805.

Gagliardo A, Ioalè P, Savini M, Wild M (2009) Navigational abilities of adult and experienced homing pigeons deprived of olfactory or trigeminally mediated magnetic information. *Journal of Experimental Biology* 212: 3119-3124.

Gwinner E, Wiltschko W (1978) Endogenously controlled changes in migratory direction of the garden warbler, *Sylvia borin*. Journal of Comparative Physiology 125: 267-273.

Heyers D, Zapka M, Hoffmeister M, Wild JM, Mouritsen H (2010) Magnetic field changes activate the trigeminal brainstem complex in a migratory bird. *Proceedings of the National Academy of Science of the United States of America* 107: 9394-9399.

Hijmans RJ, Williams E, Vennes C (2019) Package: ‘geosphere’ in R. Available <https://cran.r-project.org/web/packages/geosphere/index.html>.

Holland RA (2010) Differential effects of magnetic pulses on the orientation of naturally migrating birds. *Journal of The Royal Society Interface* 7: 1617-1625.

Hore PJ, Mouritsen H (2016) The radical-pair mechanism of magnetoreception. *Annual Review of Biophysics* 45: 299-344.

Hurvich CM, Tsai CL (1989) Regression and time-series model selection in small samples. *Biometrika* 76: 297-307.

Ioalè P (1984) Magnets and pigeon orientation. *Monitore Zoologico Italiano* 18: 347-358.

Ioalè P (2000) Pigeon orientation: effects of the application of magnets under overcast skies. *Naturwissenschaften* 87 : 232-235.

Keeton WT (1969) Orientation by pigeons: is the sun necessary? Science 165: 922-928.

Keeton WT (1971) Magnets interfere with pigeon homing. *Proceedings of the National Academy of Science of the United States of America* 68: 102-106.

Kirschvink JL, Gould JL (1981) Biogenic magnetite as a basis for magnetic field detection in animals. *Biosystems* 13: 181-201.

Kishkinev D, Chernetsov N, Heyers D, Mouritsen H (2013) Migratory reed warblers need intact trigeminal nerves to correct for a 1,000 km eastward displacement. *PLoS ONE* 8(6): e65847

Kishkinev D, Chernetsov N, Pakhomov A, Heyers D, Mouritsen H (2015) Eurasian reed warblers compensate for virtual magnetic displacement. Current Biology 25: R822-R824.

Kishkinev D, Packmor F, Zechmeister T, Winkler H-C, Chernetsov N, Mouritsen H, Holland RA (2021) Navigation by extrapolation of geomagnetic cues in a migratory songbird. Current Biology 31: 1563-1569.e4.

Kramer G (1949) Über Richtungstendenzen bei der nächtlichen Zugunruhe gekäfigter Vögel. Ornithologie als Biologische Wissenschaft (eds. Mayr E & Schüz E) 269-283.

Kramer G (1950) Weitere Analyse der Faktoren, welche die Zugaktivität des gekäfigten Vogels orientieren. *Naturwissenschaften* 37: 377-378.

Kramer G (1953) Wird die Sonnenhöhe bei der Heimfindeorientierung verwertet? *Journal für Ornithologie* 94: 201-219.

Kramer G (1957) Experiments in bird orientation and their interpretation. *Ibis* 99: 196-227.

Lamotte MM (1974) The influence of magnets and habituation to magnets on inexperienced homing pigeons. *Journal of comparative Physiology* 89: 379-389.

Liechti F, Komenda-Zehnder S, Bruderer B (2012) Orientation of passerine trans-Sahara migrants: the directional shift (‘Zugknick’) reconsidered for free-flying birds. *Animal Behaviour* 83: 63-68.

Liu X, Chernetsov N (2012) Avian orientation: multi-cue integration and calibration of compass systems. *Chinese Birds* 3: 1-8.

Lund U, Agostinelli C (2007) Package: ‘circular’ in R. Available at <http://cran.r-project.org/web/packages/circular/index.html>.

Massa B, Benvenuti SP, Ioalè P, Lo Valvo M, Papi F (1991) Homing of Cory’s shearwaters (*Calonectris diomedea*) carrying magnets. *Bollettino di zoologia* 58: 245-247.

Merkel FW, Wiltschko W (1965) Magnetismus und Richtungsfinden zugunruhiger Rotkehlchen (*Erithacus rubecula*). *Vogelwarte* 23: 71-77.

Moore BR (1988) Magnetic fields and orientation in homing pigeons: experiments of the late W. T. Keeton. *Proceedings of the National Academy of Science of the United States of America* 85: 4907-4909.

Mora CV, Walker MM (2012) Consistent effect of an attached magnet on the initial orientation of homing pigeons, *Columba livia*. *Animal Behaviour* 84: 377-383.

Mouritsen H (1998) Redstarts, *Phoenicurus phoenicurus*, can orient in a true-zero magnetic field. *Animal Behaviour* 55: 1311-1324.

Mouritsen H (2015) Magnetoreception in birds and its use for long-distance migration. Sturkie's avian physiology (ed. Scanes C) 113-133.

Mouritsen H (2018) Long-distance navigation and magnetoreception in migratory animals. *Nature* 558: 50-59.

Mouritsen H, Huyvaert KP, Frost BJ, Anderson DJ (2003) Waved albatrosses can navigate with strong magnets attached to their head. *Journal of Experimental Biology* 206: 4155-4166.

Mouritsen H, Larsen ON (2001) Migrating songbirds tested in computer-controlled Emlen funnels use stellar cues for a time-independent compass. Journal of Experimental Biology 204: 3855-3865.

Muheim R, Bäckman J, Åkesson S (2002) Magnetic compass orientation in European robins is dependent on both wavelength and intensity of light. *Journal of Experimental Biology* 205: 3845-3856.

Muheim R, Phillips JB, Åkesson S (2006) Polarized light cues underlie compass calibration in migratory songbirds. *Science* 313: 837-839.

Nimpf S, Nordmann GC, Kagerbauer D, Malkemper EP, Landler L, Papadaki-Anastasopoulou A, Ushakova L, Wenninger-Weinzierl A, Novatchkova M, Vincent P, Lendl T, Colombini1Matthew Mason MJ, Keays DA (2019) A putative mechanism for magnetoreception by electromagnetic induction in the pigeon inner ear. *Current Biology* 29: 4052-4059.e4.

Padget O, Dell’Ariccia G, Gagliardo A, González-Solís J, Guilford T (2017) Anosmia impairs homing orientation but not foraging behaviour in free-ranging shearwaters. *Scientific Reports* 7: 9668.

Pakhomov A, Anashina A, Chernetsov N (2017) Further evidence of a time-independent stellar compass in a night-migrating songbird. *Behavioral Ecology and Sociobiology* 71: 1-6.

Pakhomov A, Anashina A, Heyers D, Kobylkov D, Mouritsen H, Chernetsov N (2018) Magnetic map navigation in a migratory songbird requires trigeminal input. *Scientific Reports* 8: 11975.

Pakhomov A, Chernetsov N (2020) A hierarchy of compass systems in migratory birds. Biological Communications 65: 262-276.

Pollonara E, Luschi P, Guilford T, Wikelski M, Bonadonna F, Gagliardo A (2015) Olfaction and topography, but not magnetic cues, control navigation in a pelagic seabird: displacements with shearwaters in the Mediterranean Sea. *Scientific Reports* 5: 16486.

Ranvaud R, Schmidt-Koenig K, Ganzhorn JU, Kiepenheuer J, Gasparotto OC, Britto LRG (1991) The initial orientation of homing pigeons at the magnetic equator: compass mechanisms and the effect of applied magnets. *Journal of Experimental Biology* 161: 299-314.

R Core Team (2021) R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. Retrieved from http://www.r-project.org/

Ritz T, Adem S, Schulten K (2000) A model for photoreceptor-based magnetoreception in birds. *Biophysical Journal* 78: 707-718.

Schmidt-Koenig K (1958) Der Einfluß experimentell veränderter Zeitschätzung auf das Heimfindevermögen bei Brieftauben. *Die Naturwissenschaften* 45: 47-48.

Schnute JT, Groot K (1992) Statistical analysis of animal orientation data. *Animal Behaviour* 43: 15-33.

Southern WE (1972) Influence of disturbances in the earth's magnetic field on Ring-billed Gull orientation. *The Condor* 74: 102-105.

Syposz M, Padget O, Wynn J, Gillies N, Fayet AL, Guilford T (2021) An assay to investigate factors influencing initial orientation in nocturnally fledging seabirds. Journal of Avian Biology 52: e02613.

Visalberghi E, Alleva E (1979) Magnetic influences on pigeon homing. *The Biological Bulletin* 56: 246-256.

Walcott C, Green RP (1974) Orientation of homing pigeons altered by a change in the direction of an applied magnetic field. *Science* 184: 180-182.

Wallraff HG (2005) Avian navigation: pigeon homing as a paradigm. Heidelberg: Springer Verlag.

Wang K, Mattern E, Ritz T (2006) On the use of magnets to disrupt the physiological compass of birds. Physical Biology 3: 220-231.

Wikelski M, Arriero E, Gagliardo A, Holland RA, Huttunen MJ, Juvaste R, Mueller I, Tertitski G, Thorup K, Wild M, Alanko M, Bairlein F, Cherenkov A, Cameron A, Flatz R, Hannila J, Hüppop O, Kangasniemi M, Kranstauber B, Penttinen M-L, Safi K, Semashko V, Schmid H, Wistbacka R (2015) True navigation in migrating gulls requires intact olfactory nerves. *Scientific Reports* 5: 17061.

Wiltschko W (1968) Über den Einfluß statischer Magnetfelder auf die Zugorientierung der Rotkehlchen (*Erithacus rubecula*). *Zeitschrift für Tierpsychologie* 25: 537-558.

Wiltschko W, Daum P, Fergenbauer-Kimmel A, Wiltschko R (1987) The Development of the Star Compass in Garden Warblers, *Sylvia borin*. *Ethology* 74: 285-292.

Wiltschko W, Munro U, Ford H, Wiltschko R (1993) Red light disrupts magnetic orientation of migratory birds. *Nature* 364: 525-527.

Wiltschko W, Munro U, Ford H, Wiltschko R (1998) Effect of a magnetic pulse on the orientation of silvereyes, *Zosterops l. lateralis*, during spring migration. *Journal of Experimental Biology* 201: 3257-3261.

Wiltschko R, Wiltschko W (1995a) Magnetic orientation in animals. Berlin: Springer Verlag.

Wiltschko W, Wiltschko R (1972) Magnetic compass of European robins. *Science* 176(4030): 62-64.

Wiltschko W, Wiltschko R (1995b) Migratory orientation of European robins is affected by the wavelength of light as well as by a magnetic pulse. *Journal of Comparative Physiology A* 177: 363-369.

Wiltschko W, Wiltschko R (2001) Light-dependent magnetoreception in birds: the behaviour of European robins, *Erithacus rubecula*, under monochromatic light of various wavelengths and intensities. *Journal of Experimental Biology* 204: 3295-3302.

Wynn J, Padget O, Mouritsen H, Perrins C, Guilford T (2020) Natal imprinting to the Earth’s magnetic field in a pelagic seabird. *Current Biology* 30: 2869-2873.e2.

Xu J, Jarocha LE, Zollitsch T et al. (2021) Magnetic sensitivity of cryptochrome 4 from a migratory songbird. *Nature* 594: 535-540.

**Figure 1:** Schematic overview of the experimental procedure during the repeated orientation tests of Experiment 1 and 2. (A) Experiment 1: each of the 17 birds included in the experiment underwent a total of 12 orientation tests within a test period of 15 nights, four orientation tests under each experimental treatment (CON, MAG, SHAM). Birds in housing group A (9 birds) and housing group B (8 birds) were subjected to the MAG and SHAM treatment in opposing order and had no access to celestial cues in either test. (B) Experiment 2: each of the 18 birds included in the experiment underwent a total of 8 orientation tests within a test period of 10 nights, four orientation tests under each experimental treatment (MAG, MAG + CEL). Birds in both housing groups were subjected to the MAG treatment during the test period. Access to celestial cues (CEL) during the orientation tests was given or denied in alternating fashion among the test nights, depending on the local weather (cloud cover) during the test period. For both experiments the number of birds tested each night of the test periods is given.

**Figure 2:** Magnets (MAG) and sham attachments (SHAM; brass) used for the experiments and their position on the Eurasian reed warblers’ foreheads. Magnets were attached with their North Pole facing down/South pole facing up (Experiment 1) or North Pole facing up/South pole facing down (Experiment 2). Magnet and sham attachment dimensions: Ø 3 mm, height 2 mm; weight ca. 0.11 g.

**Figure 3:** Natural migratory direction of Eurasian reed warblers from Lake Neusiedl during autumn as derived from recoveries of birds ringed at Lake Neusiedl (red dot) both in Austria and Hungary during summer and early autumn the same year. Black arrows depict great circle lines between the ringing site and the respective recovery sites (>250 km distance from ringing site); red arrow depicts the birds’ mean migratory direction and mean recovery distance based on these recoveries. Bird ring recovery data were provided by the Austrian Ornithological Centre and the Hungarian Bird Ringing Centre. The map is an orthographic projection with Lake Neusiedl as the projection centre.

**Figure 4:** Results from the orientation tests of Experiment 1 and 2. (A) Magnetic compass orientation of first-year Eurasian reed warblers tested under the different experimental treatments (CON, MAG, SHAM) of Experiment 1 during autumn migration. (B) Magnetic and celestial compass orientation of first-year Eurasian reed warblers tested under the different experimental treatments (MAG, MAG + CEL) of Experiment 2 during autumn migration. Circular diagrams depict the birds’ individual mean directions from repeated orientation tests (filled circles). Birds that showed directionality only once during repeated orientation tests (open circles) were not considered in subsequent analyses. Arrows show the group-specific mean directions and the mean resultant vector lengths of significantly oriented treatment groups (according to the Rayleigh test of uniformity); dashed circles indicate the radius the respective mean resultant vector needs for the 5% and 1% significance levels according to the Rayleigh test of uniformity; solid lines flanking the arrows give the 95% confidence intervals for the group-specific mean directions; all directions are depicted relative to the magnetic North (mN).

**Table 1:** Summary of the ΔAICc and the AICc model weights (*w*) from the 10 orientation models described by Schnute and Groot (1992) assessed for orientation data obtained under the different experimental treatments (CON, MAG, SHAM) of Experiment 1 (2019). Values from the respective best models with a ΔAIC < 2 are shown in bold and highlighted by asterisks.

|  |  |  |  |
| --- | --- | --- | --- |
| Model | **CON** | **MAG**  | **SHAM** |
| ΔAICc | *w* | ΔAICc | *w* | ΔAICc | *w* |
| M1 (uniform) | 4.71 | 0.05 | **0.00\*** | **0.71\*** | 8.56 | 0.01 |
| M2A (unimodal) | **1.51\*** | **0.23\*** | 4.59 | 0.07 | **0.00\*** | **0.71\*** |
| M2B (symmetric modified unimodal) | **0.00\*** | **0.49\*** | 4.42 | 0.08 | 3.13 | 0.15 |
| M2C (modified unimodal) | 2.96 | 0.11 | 7.95 | 0.01 | 4.43 | 0.08 |
| M3A (homogenous symmetric bimodal) | 8.34 | 0.01 | 4.20 | 0.87 | 12.35 | 0.00 |
| M3B (symmetric bimodal) | 4.09 | 0.06 | 7.52 | 0.02 | 7.14 | 0.02 |
| M4A (homogenous axial bimodal) | 6.23 | 0.02 | 7.00 | 0.02 | 9.40 | 0.01 |
| M4B (axial bimodal) | 6.81 | 0.02 | 11.04 | 0.00 | 14.81 | 0.00 |
| M5A (homogenous bimodal) | 8.12 | 0.01 | 11.69 | 0.00 | 6.89 | 0.02 |
| M5B (bimodal) | 10.74 | 0.00 | 17.26 | 0.00 | 11.94 | 0.00 |
| *w* = AICc model weights |

**Table 2:** Summary of the ΔAICc and the AICc model weights (*w*) from the 10 orientation models described by Schnute and Groot (1992) assessed for orientation data obtained under the different experimental treatments (MAG, MAG + CEL) of Experiment 2 (2020). Values from the respective best models with a ΔAIC < 2 are shown in bold and highlighted by asterisks.

|  |  |  |
| --- | --- | --- |
| Model | **MAG** | **MAG + CEL** |
| ΔAICc | *w* | ΔAICc | *w* |
| M1 (uniform) | **1.39\*** | **0.22\*** | 3.95 | 0.07 |
| M2A (unimodal) | 4.85 | 0.04 | **0.00\*** | **0.52\*** |
| M2B (symmetric modified unimodal) | 5.30 | 0.03 | 2.21 | 0.17 |
| M2C (modified unimodal) | 7.12 | 0.01 | 4.15 | 0.07 |
| M3A (homogenous symmetric bimodal) | **0.00\*** | **0.44\*** | 8.19 | 0.01 |
| M3B (symmetric bimodal) | 3.17 | 0.09 | 5.38 | 0.04 |
| M4A (homogenous axial bimodal) | 3.12 | 0.09 | 7.03 | 0.02 |
| M4B (axial bimodal) | 6.92 | 0.01 | 8.37 | 0.01 |
| M5A (homogenous bimodal) | 4.03 | 0.06 | 3.57 | 0.09 |
| M5B (bimodal) | 8.66 | 0.00 | 7.73 | 0.01 |
| *w* = AICc model weights |