Behavioral Ecology and Sociobiology

Cue-conflict experiments between magnetic and visual cues in dunlin Calidris alpina and curlew sandpiper Calidris ferruginea --Manuscript Draft--

| Manuscript Number: | BEAS-D-16-00614R2 | |
|--|---|---|
| Full Title: | Cue-conflict experiments between magnetic and visual cues in dunlin Calidris alpina and curlew sandpiper Calidris ferruginea | |
| Article Type: | Original Article | |
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| Funding Information: | Ministero dell'Istruzione, dell'Università e della Ricerca (PRIN 20083ML4XC) | Dr Dimitri Giunchi |
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| Response to Reviewers: | Dear Dr Bakker, Thank you for your positive decision on the r "Cue-conflict experiments between magnetic and curlew sandpiper Calidris ferruginea". I checked all comments and made the reque p values for the t-tests in the Results section use the exact p value for the Moore test (line from the table in the paper by Moore (1980; | c and visual cues in dunlin Calidris alpina ested corrections. I inserted the requested n (lines 194 and 211); however I can not es 195 and 204) since the values derive |

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Supplementary Material

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| 1 | Cue-conflict experiments between magnetic and visual cues in dunlin Calidris alpina and curlew sandpiper |
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10 Abstract

11 Despite our extensive knowledge on various aspects of their lives, there has been limited investigation into the 12 hierarchical relationships among different compass systems in shorebirds. The aim of this study was to investigate the relationship between magnetic and celestial compasses in two species of shorebirds, the curlew sandpiper (Calidris 13 ferruginea; pre-breeding migration) and the dunlin (Calidris alpina; post-breeding migration) using cue-conflict 14 15 experiments. Birds were captured in a Mediterranean stopover site, after which their magnetic orientation was determined under simulated overcast conditions at sunset using modified Emlen funnels fitted with infrared video 16 17 cameras. Birds that demonstrated a well-defined directional preference were then exposed over two sunsets to conflicting directional information between the local geomagnetic field and the \pm 90° shifted band of maximum 18 19 polarisation. These individuals were tested again for magnetic orientation at sunset in the same conditions as previous 20 test, to determine whether their directional choices had changed after the cue-conflict. Our results showed that 21 individuals from both species did not recalibrate their magnetic compass from visual cues after the cue-conflict, even 22 though at least dunlins did not appear to completely disregard the information derived from celestial cues. This study is 23 one of the few experimental studies on the migratory orientation of Charadriiformes and on the hierarchical 24 relationships between the different compasses used by these birds during their extensive migratory movements.

25

26 Keywords: orientation, cue-conflict, polarized light, shorebirds, magnetic compass

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28 Significance Statement

29 Migrating birds are able to use different compass mechanisms based on geomagnetic or celestial cues and it seems

30 reasonable to hypothesise that birds calibrate their various compasses to maintain the correct direction especially when

the directional information does not agree. The hierarchy among different compasses has been studied largely on night migrating passerines, but it is still poorly understood. We investigated the hierarchy among geomagnetic and celestial cues (band of maximum polarization) in two species of *Charadriiformes* by means of cue-conflict experiments. Our result showed that the geomagnetic cues have a dominant role in the orientation mechanisms of the studied species, even though the information derived from celestial cues did not appear to be completely disregarded.

36

37 Introduction

38 Migrating birds are able to use different compass mechanisms based on geomagnetic (Wiltschko and Wiltschko 1996) 39 or celestial cues, including the sun (Kramer 1953; Moore 1987), stars (Emlen 1975) and skylight polarization pattern 40 (Able and Able 1995; Cochran et al. 2004; Muheim et al. 2006b). However, the redundancy of various cues may be a 41 problem for migrating birds, especially when the directional information does not agree (Wiltschko and Wiltschko 42 1999; Muheim et al. 2006a). Therefore, it seems reasonable to hypothesise that birds calibrate their various compasses 43 to maintain the correct direction (Muheim et al. 2006a; Sjöberg and Muheim 2016), however, the hierarchy among 44 different compasses remains poorly understood (Liu and Chernetsov 2012). While there is general agreement that 45 celestial cues dominate over magnetic cues during the pre-migratory period (Bingman 1983; Able and Able 1990; Prinz 46 and Wiltschko 1992; Wiltschko et al. 1998; Wiltschko and Wiltschko 1999), the relationship between these cues during 47 migration remains unclear (Muheim et al. 2006a; Wiltschko and Wiltschko 2009). It has recently been proposed that at 48 sunset, and possibly also at sunrise, the Band of Maximum Polarization of skylight (BMP) may be the reference system 49 used by birds to recalibrate their magnetic compass during the migratory period (Muheim et al. 2006a, b). This use of 50 BMP would give birds a reference system that is independent from both topography (Liu and Chernetsov 2012) and 51 magnetic anomalies. Unlike the sun compass (Schmidt-Koenig 1990), the use of BMP does not require any time 52 compensation as at sunrise and sunset it intersects vertically the horizon roughly along the North-South axis, 53 independent of latitude (Phillips and Waldvogel 1988; Muheim et al. 2006a). Some cue-conflict (CC) experiments seem 54 to have confirmed this hypothesis (Phillips and Moore 1992; Cochran et al. 2004; Muheim et al. 2006b, 2007), 55 however, the role of polarisation pattern is still a topic of debate (Wiltschko et al. 2008a, b; Muheim et al. 2008). Further studies on both caged and free-flying birds have cast doubts on the use of BMP for recalibrating magnetic 56 57 references (Chernetsov et al. 2011; Schmaljohann et al. 2013; Åkesson et al. 2015), or have suggested that birds may 58 not always reconcile conflicting information from geomagnetic and celestial (star) cues, therefore making a form a 59 compromise between the directions indicated by the two reference systems (Giunchi et al. 2015). Sjöberg and Muheim 60 (2016) recently attempted to integrate the different outcomes by suggesting that the discrepancies observed between

cue-conflict experiments may be due to a combination of both the availability of polarised light information near the
 horizon and stars. While intriguing, this explanation is highly speculative and a number of experimental observations
 remain unexplained (see Sjöberg and Muheim 2016 for further details).

64 Until recently, almost all available data concerning the orientation mechanisms of wild birds have been derived from experiments on a few species of nocturnal passerine migrants. Furthermore, while there is a significant amount of 65 66 information regarding the migratory ecology and routes of shorebirds (see Colwell 2010), only few studies have 67 investigated the orientation mechanisms of these species. To our knowledge, only four papers have been published on 68 this topic. Sauer (1963) and Sandberg and Gudmundsson (1996) initially investigated shorebird orientation in modified 69 Emlen funnels (Emlen and Emlen 1966), demonstrating that both Pacific golden plovers (Pluvialis fulva) and dunlins 70 (*Calidris alpina*) showed directional preferences consistent with the predicted migratory direction when tested under a 71 clear sky. Following these initial observations, the use of magnetic cues were demonstrated by studies that showed that 72 sanderlings (Calidris alba; Gudmundsson and Sandberg 2000) and sharp-tailed sandpipers (Calidris acuminata; 73 Grönroos et al. 2010) responded as expected to a $\pm 90^{\circ}$ deflection in the local magnetic field. To date, the hierarchy 74 between different compass mechanisms has not yet been investigated in shorebirds. 75 The aim of the present study was to investigate the relationship between magnetic and visual compasses (i.e. 76 the pattern of skylight polarisation) in two species of shorebirds using cue-conflict experiments.

77

78 Materials and methods

79 Study location and experimental birds

80 Birds were caught with mist nets during the pre-breeding (curlew sandpiper *Calidris ferruginea*; April–May) and post-81 breeding (dunlin Calidris alpina; August-September) migrations from 2012 to 2015 in the Lame di San Rossore 82 wetland complex (Massaciuccoli Regional Park, Migliarino, San Rossore, Pisa, Lucca, Italy; 43°41'N, 10°17'E). After 83 the standard ringing procedures (Busse and Meissner 2015) the birds were aged and sexed according to Prater et al. 84 (1977). Due to the migratory strategies of these species (del Hoyo et al. 1996), our experimental sample of curlew 85 sandpiper was composed entirely of adults, while we tested both adults and juvenile dunlins. Birds were weighed using 86 an electronic scale (± 0.1 g) and then held in captivity in the Arnino Field Station (43°39'N 10°18'E; capture site 87 direction = 327°, capture site distance = 3.5 km) for a maximum of 10 days in cylindrical plastic cages (diameter = 100 88 cm, height = 30 cm). The birds were fed with mealworm (*Tenebrio molitor*) larvae and were provided water *ad libitum*. 89 According to the social habits of the considered species (del Hoyo et al. 1996), individual were kept in groups of 3 to 5 90 birds per cage. The cages were placed in rooms with artificial lighting that was synchronised with the natural

- 91 photoperiod so that birds could not observe any celestial cues. During captivity, the mean weight of the curlew
- 92 sandpipers and dunlins increased by 9.7 g ($t_{12} = 6.7$, p < 0.01, paired t-test; capture weight: 49.7 ± 3.6 g SD) and 13.8 g
- 93 (t₉ = 10.4, p < 0.01, paired t-test; capture weight: 39.5 ± 2.7 g), respectively.
- 94 Experimental procedures
- 95 Our experimental protocol was followed as described below:
- 96 1) After a minimum of two days in captivity the magnetic orientation of birds was tested at sunset under
 97 simulated overcast conditions (hereafter referred to as the pre-CC test).
- 98 2) Birds that demonstrated a well-defined directional preference in the pre-CC test were exposed (CC
- 99 exposure) at sunset for two consecutive days to conflicting directional information between the local geomagnetic field
- and the \pm 90° shifted BMP (for further details, see Giunchi et al. 2015). After each CC exposure, birds were returned to
- 101 their indoor cages and prevented from seeing any visual cue.
- 3) Treated birds were tested again the day following the second CC exposure for their magnetic orientation at
 sunset in the same conditions as the pre-CC test. The purpose of this test (post-CC test) was to evaluate the consistency
 of directional choices exhibited before and after the cue-conflict.
- 105

106 Experimental apparatus

107 The orientation tests were performed using modified Emlen funnels which were made out of non-magnetic transparent 108 material (LEXAN; Sabic Innovative Plastic, Pittsfield, MA), the size of which was consistent with that described by 109 Sandberg and Gudmundsson (1996) (top external diameter = 52 cm, base diameter = 18 cm and height = 22 cm). Each 110 funnel was divided into eight sectors of 45° using strips of tape, and then placed in plastic containers (70 cm high). An 111 infrared camera was positioned at the bottom of the plastic containers (see Online Resources, Fig. S1) which was 112 powered by a 12 V battery and connected to a digital video recorder. Each funnel was covered by plastic mesh netting 113 $(2 \times 2 \text{ mm})$. We measured the magnetic field inside the funnels using a fluxgate magnetometer (Applied Physics 114 Systems, Mountain View, CA) to ensure that the magnetic field perceived by the experimental birds was not affected by 115 the experimental apparatus. Before the beginning of each test, each funnel was oriented randomly with respect to the 116 magnetic North. A cubic wooden box $(90 \times 90 \times 50 \text{ cm})$ with four square windows $(48 \times 48 \text{ cm})$ was placed on the top 117 of each funnel in the centre of each side. The top of the funnel and the windows were covered by milky-white 118 Plexiglass lids (3 mm thickness). This structure prevented birds from perceiving the light polarisation (Sandberg et al. 119 1988; Åkesson et al. 2015) and other visual cues while also avoiding excessive darkness inside the funnels (Fig. S1).

To manipulate the skylight polarisation pattern we replaced the Plexiglass windows on the same cubic wooden boxes with panels of two outer layers of pseudo-depolarising filters composed of two sheets of polyester (180 μm thick; Metalloy Italiana, Vicenza, Italy) which were aligned at an angle of 45° relative to each other, in addition to one inner layer composed of a polarising filter (0.8 mm thick, cellulose triacetate; Intercast Europe, Parma, Italy), as described by Gaggini et al. (2010). The polarising filters from the two opposite windows were aligned so that the e-vector was vertical, while the e-vector was horizontal for the other two windows (Gaggini et al. 2010). The top of each box was covered by a milky-white Plexiglass lid (3 mm thick).

During CC exposure, the birds were kept in a small wooden cage $(40 \times 20 \text{ cm})$ with netting on each side, which was placed in the centre of the larger wooden box. The small size of this holding cage compared to the box was used to reduce any parallax effect on the alignment of the artificial polarisation axis (Muheim et al. 2006b). When the vertical filters were aligned with the solar azimuth, the caged birds perceived the BMP shifted by \pm 90° relative to natural conditions (Gaggini et al. 2010). The entire structure was placed on wooden stands (70 cm high) to give the birds a broad view of the horizon through the polarising filters.

We performed both orientation tests and exposures to cue-conflict only in calm evenings with a wind speed of < 10 m/s and no rain. We did not perform exposures to cue-conflict when cloud cover was higher than 3/8. The orientation tests lasted for 40 minutes and were performed between 20 minutes before and 70 minutes after the civil twilight (sun elevation $< -6^\circ$). Exposures lasted for 40 minutes and occurred between 20 minutes before and 20 minutes after civil twilight. The orientation of each individual was tested over a maximum of three consecutive evenings, and birds that remained inactive (see below) during all these trials were released close to the capture site. The first test in which the individual showed to be active was considered for the analyses.

140

141 Data analyses

142 The video recordings were analysed to determine the directional preferences of each individual. The first 10 minutes of 143 the recordings were discarded in order to allow the birds to acclimatise to the funnel. The remaining 30 minutes were 144 subdivided into 600 frames (1 frame every 3 seconds) which was then combined into a single multipage Tiff file. A 145 circle was drawn on each image 6 cm from the bottom of the funnel for use as the reference line. Only well-defined 146 movements (WDMs) were considered for the analysis, i.e. only when birds crossed the reference line with at least a 147 third of the front of its body. The WDMs within a given sector were assigned to the mid-angle of the sector. Only active 148 birds (WDMs \geq 10) were included in the analyses. Individual directional preferences were calculated as the circular 149 mean of the WDM distribution (Batschelet 1981). Each of the multipage Tiff files was independently examined by three observers blind to the experimental condition. As the circular means estimated by the three observers did not differ by more than 30°, all active birds were included in the analysis. The orientation of each bird was calculated by averaging the three directional choices estimated by the three observers. Birds were excluded from the analyses if the result of the Rayleigh test (Batschelet 1981) applied to the circular distribution of their individual headings was not significant (p >0.05). As previously reported (see e.g. Muheim and Åkesson 2002), the Rayleigh test was only used as a guideline as the movement of birds inside the funnel could not be considered to be independent. Actually all active birds of both species turned out to be oriented according to this criterion.

157 The group statistics were calculated from individual mean directions so that each data point represented one bird (Batschelet 1981). When the mean vector length resulting from doubling the angles was larger than the unimodal 158 159 vector length, we used a mean axis of orientation as the basis of the analysis. Randomness was tested with the Rayleigh 160 test and the 95% confidence interval for the mean group directions were calculated using bootstrap methods with 5000 161 resampled datasets (Adams and Anthony 1996) only for significantly oriented samples. The effect of CC exposure was 162 determined by analysing the distribution of the angular differences of individual headings between the pre-CC and post-163 CC tests (post-pre) and using the non-parametric paired-sample test described by Moore (1980). As either unimodal 164 and axially bimodal circular distributions did not seems to describe satisfactorily the post-pre sample distribution in 165 dunlins, we used a model based clustering approach (Hornik and Grün 2014 and references therein) by means of the R 166 package Directional 2.4 (Tsagris et al 2016) to evaluate whether a mixture of two von Mises-Fisher distributions fitted 167 the data better than the standard unimodal model. We firstly calculated the Bayesian information criterion (BIC; 168 Schwarz 1978) of the models with one or two components using the bic.mixvmf function, assuming a mixture of two 169 von Mises-Fisher distributions. The model with the lowest BIC value was chosen to best describe the data. If a bimodal 170 model had the lowest BIC value then a mixture of two von Mises-Fisher distributions were fitted to the data using the 171 mix.vmf function, which gave the predicted group assignment for each observation. As a check of this approach, we 172 applied this method to all samples (Table S1). All statistical analyses were performed with the software R 3.3.2 (R Core 173 Team 2016); standard circular statistic analyses were performed using the R package circular 0.4–7 (Agostinelli and 174 Lund 2013).

175

176 Expectations

- 177 We expected that in the pre-CC test the birds would orient themselves according to the distribution of
- 178 recovery/recapture of birds ringed in Italy during the spring (curlew sandpiper) or autumn (dunlin) migration (Spina and

179 Volponi 2008). Depending on the hierarchy determined by the information provided by the celestial cue (BMP) and the

- 180 geomagnetic field, each individuals could respond to the cue-conflict in two possible ways:
- 181 If the magnetic cue was dominant over BMP, we expected that there would be no differences in the orientation 182 of the birds between the pre-CC and post-CC tests (Fig. 1).
- 183 If BMP was used to calibrate the magnetic compass, we expected to observe a significant $\pm 90^{\circ}$ shift in
- 184 directional preferences in the post-CC test compared to the pre-CC test (Fig. 1; Cochran et al. 2004; Muheim et al.
- 185 2006b).
- 186
- 187 **Results**
- 188
- 189 Curlew sandpiper

190 30 adult curlew sandpipers tested in the spring of 2013 and 2014, 15 were active in the pre-CC test. Two birds were 191 inactive during the post-CC test for three consecutive trials, therefore, they were excluded from the analyses. The 192 directional preferences of the remaining 13 individuals were bimodally distributed during the pre-CC test along the 193 WSW-ENE axis (Fig. 2). We did not find any difference in body weight between WSW [225°-300°] and ENE [350°-194 110°] oriented birds (54.5 \pm 4.3 g vs 57.4 \pm 3.2 g; t_{10.8} = 1.37, p = 0.2, Welch's t-test). The distributions before (pre-CC 195 test) and after (post-CC test; see Fig. 2) the CC exposure were not significantly different ($r_m = 0.12$, p > 0.05, Moore 196 test). The angular differences between the post-CC test and the pre-CC test (post-pre) were unimodally distributed and 197 the mean direction ($\alpha = 13^{\circ}$) was not significantly different from 0° (95% CI = [331°-35°]; Fig. 2).

198

199 Dunlin

200 We tested 19 dunlins during the post-breeding migration between 2012 and 2015. Eleven birds were active in the pre-201 CC test, but only 10 (five juveniles and five adults) were found to be active in the post-CC test and hence included in 202 the analyses. In the pre-CC test (Fig. 2) birds headed in a SSW direction, and no significant differences were recorded 203 between the heading distributions before (pre-CC test) and after (post-CC test; see Fig. 2) the CC exposure ($r_m = 0.70, p$ 204 > 0.05, Moore test). The angular differences between post-CC test and pre-CC test (post-pre, Fig 2) were unimodally 205 distributed with a mean direction of - 58° (95% CI: [261°-342°]). The model based clustering approach analysis 206 indicated that a model with two components (a mixture of two von Mises-Fisher distribution) described the post-pre 207 distribution better than a unimodal model (BIC = 33.07 vs 33.47, respectively; see Table S1). This approach identified 208 two well-defined clusters: birds belonging to one cluster shifted their directional preferences 109° counter-clockwise, while individuals belonging to the second cluster did not change their directional preferences after the cue conflict (Fig. 2; Table S1). The mean weight of birds belonging to the two cluster, recorded just before the post-CC test, was not significantly different (cluster $1 = 52.5 \pm 7.2$ g vs cluster $2 = 54.1 \pm 0.7$ g; $t_{4.08} = 0.52$, p = 0.63, Welch's t-test)

212

213 Discussion

This work is one of the few studies aimed at investigating the migratory orientation of *Charadriiformes* and the hierarchical relationship between the visual and magnetic compass systems. Our results show that both curlew sandpipers and dunlins are able to orient themselves under simulated overcast conditions, as previously demonstrated for sanderlings (Gudmundsson and Sandberg 2000) and sharp-tailed sandpipers (Grönroos et al. 2010).

218 The percentage of inactive individuals was noticeable (ca. 50% in both species), but it is consistent with the 219 results of a number of orientation studies on waders, which reported that these birds are more likely to not show 220 migratory restlessness when tested under overcast conditions (see e.g. Sandberg and Gudmundsson 1996; 221 Gudmundsson and Sandberg 2000, but see Grönroos et al. 2010). Even though free-flying waders can depart from 222 stopover/staging sites and orient correctly under solid overcast, it should be noted that (1) Alerstam et al. (1990) 223 reported that in most occasions knot and turnstone flocks departed under clear sky conditions and (2) all the 277 224 departures of wader flocks observed by Piersma et al. (1990) in the Dutch Wadden Sea occurred when the sun was 225 clearly visible. Furthermore, it should be noted that the migratory behaviour of shorebirds is often quite different from 226 that of passerines, being characterized by long flights followed by prolonged stay at staging site (Piersma 1987); this 227 behaviour might affect the motivation to migrate of birds captured while refuelling thus increasing the inter-individual 228 variability in the expression of migratory restlessness. For this reasons we suggest that the fraction of inactive birds is 229 not an artifact due to our experimental apparatus, but a consequences of having tested birds without available visual 230 cues.

231

232 Orientation preferences in the pre-CC test

Dunlins showed a clear directional preference toward a SSW direction, which is in agreement with the distribution of the ringing recoveries reported by Spina and Volponi (2008) and to the expected direction for the post-breeding migration (Cramp and Simmons 1983). The bimodal distribution of the directional preferences of curlew sandpipers is more puzzling. According to previous studies (Wilson et al. 1980; Delany et al. 2009), curlew sandpipers crossing the Mediterranean appear to follow an eastern route during spring migration, moving from Western Africa to their breeding grounds via Tunisia, Crimea and the Caspian Sea. The number of Italian recoveries of abroad ringed birds during the 239 spring migration is quite low; however, most of the recoveries were located along a SW-NE axis between the African 240 Atlantic coast and the Crimea peninsula (Scebba and Moschetti 2006; Spina and Volponi 2008). Therefore, the axis of 241 orientation of tested curlew sandpipers is consistent with the migration axis of migrating Curlew sandpipers in spring, 242 as supported by Italian ringing data. The fact that the majority of birds preferred a westward direction is however 243 problematic. The orientation of experimental birds may have been affected by a positive phototaxis towards the setting sun (see e.g. Åkesson and Sandberg 1994; Muheim et al. 2006a), as the solar azimuth in the mid-period of the 244 245 experiments was approximately 295°. This explanation seems however unlikely because the experiments were 246 performed under simulated overcast conditions obtained by opaque milky plexiglass lids, which diffuse the low ambient 247 light present at the time of the experiments. A phototactic response should characterize mostly birds tested in the first 248 period of the experiments (20 minutes before to 20 minutes after the civil twilight), but this was not confirmed by the 249 data reported in Fig. 2 (pre- and post-CC tests). Furthermore, this kind of response was absent in dunlins, that were all 250 tested in the first period of the experiments (sun azimuth in the mid-period of the experiments = $\sim 287^{\circ}$). The directional 251 preferences of the sub-group westward oriented can be interpreted as reverse migration (Åkesson et al. 1996). While reverse migration is more likely when the access to visual cues is prevented (Åkesson et al. 2001; Nilsson and Sjöberg 252 253 2016), such as in this study, this behaviour is usually shown by lean birds facing an ecological barrier (Sandberg and 254 Moore 1996; Åkesson et al. 1996; Sandberg et al. 2002; Deutschlander and Muheim 2009; Schmaljohann et al. 2011). The individuals tested were found to be in good condition (see Materials and Methods) and did not appear to be facing 255 256 an ecological barrier, therefore, this explanation seems unlikely. It can be speculated that at least a part of the tested 257 individuals oriented toward the sea-coast (located about 1.7 km west of the test site), possibly relying on infra-sounds, 258 as previously suggested for homing pigeons (Hagstrum 2000, 2013). In addition to providing an important food source (Colwell 2010), the coastline is also an important landmark for migrating birds (Åkesson 1993), and therefore, the 259 260 directional preferences of the curlew sandpipers may have been affected by the closeness of the coast. However, it 261 should be noted that this effects was absent in dunlins; moreover, as reported by Holland (2014) the effect of infrasound 262 on bird migration has never been experimentally demonstrated.

263

264 Response to the cue-conflict

We did not record any significant difference between pre- and post-CC distributions in both species, which suggest that neither dunlins nor curlew sandpipers significantly changed their directional preferences after CC exposure, contrary to that expected if birds had recalibrated their magnetic compass based on the artificial skylight polarisation pattern (Cochran et al. 2004; Muheim et al. 2006a, b, 2007, 2009). Our results are hence in broad agreement with recent studies 269 indicating the primary role of the magnetic cues in the migratory orientation of passerines (Gaggini et al. 2010; 270 Chernetsov et al. 2011; Schmaljohann et al. 2013; Åkesson et al. 2015). Actually, the majority of songbird species 271 tested did not appear to recalibrate their magnetic compass based on the skylight polarisation pattern (Åkesson et al. 272 2015). The results from the current study add two species of non-passerine long-distance migrants to this pattern, and 273 taken together, these results suggest that recalibration is less widespread than that hypothesized by Muheim et al. 274 (2006a). However, the outcomes from some studies may have been affected by issues in methodology (see also Giunchi 275 et al. 2015). According to Wiltschko et al. (2008a) the experimental protocol also used in our experiments may lead to 276 artefacts due to the totally artificial nature of the band of polarization. It should to be noted that the experiments 277 performed by Muheim et al. (2009), where birds were exposed either to shifted magnetic fields or to shifted artificial 278 polarization pattern, produced consistent results, thus suggesting that the potential artefacts due to polarizing filters *per* 279 se are at least negligible. Sjöberg and Muheim (2016) recently modified the model proposed by Muheim et al. (2006a) 280 to suggest that the discrepancies observed in the various experiments may be due to a combination of the availability of 281 polarised light information near the horizon and of the stars during the conflict. According to this model, birds 282 recalibrate their magnetic compass using polarised light cues, provided they have access to the BMP and the view of 283 surroundings. In our study birds had a full view of the BMP and distant landmarks (a pinewood strip running parallel to 284 the coast, in particular). Therefore, according to Sjöberg and Muheim (2016), we would still expect to observe 285 recalibration of the magnetic compass even though the birds did not have access to stars information. This recalibration 286 however was not observed, as no significant difference between pre- and post-CC tests was recorded for both species. 287 Birds might not had recalibrated their magnetic compass because of a "switching off" of the calibration after prolonged 288 exposure to visual landmarks in the same area (Sjöberg and Muheim 2016). Our tested birds were displaced from the 289 capture site and held indoor with no access to landmarks before and during the experiments; they could see the 290 surroundings only during the exposure to the cue-conflict. It thus seems quite unlikely that they did not pay attention to 291 calibration cues due to the familiarity to local landmarks. It can be suggested that birds did no recalibrate their magnetic 292 compass because they could rely on infra-sounds to sense the sea-coast which can be used as a prominent directional 293 cue. While deserving further investigation, this explanation has to be regarded as highly speculative, as no experiment 294 has demonstrated yet the effect of infra-sounds on the orientation of migratory birds (Holland 2014). The reaction of 295 birds to the cue-conflict might depend also on the reliability of the compass information that they experienced in 296 different geographic areas (Chernetsov et al. 2011; Åkesson et al. 2015). It should be noted, however, that the analysis 297 reported by Sjöberg and Muheim (2016) did not find any relationship between the temporal and spatial variability of

magnetic field properties (declination, intensity and inclination) and the outcomes from cue-conflict experimentsperformed in North America and Europe.

300 While the heading distributions obtained before and after the CC exposure was not significantly different in 301 dunlins, the 95% CI of the mean direction of the angular differences between pre- and post-CC tests did not include 0°, 302 while they marginally included 90°, thus indicating that dunlins shifted their orientation after the CC exposure at least 303 slightly. Actually, according to the model-based clustering approach, the circular distribution of these angular 304 differences should be properly described as a mixture of two unimodal distributions, which suggests that the way 305 dunlins reacted to the cue-conflict was individually variable, but that the skylight polarisation pattern is not completely 306 disregarded. According to this model, half of the tested dunlins (cluster 1 in Fig. 2) did not change their directional 307 preferences after the cue-conflict, whereas the remaining individuals (cluster 2 in Fig. 2) shifted their directional 308 preferences close to the 90° shift predicted if birds were to recalibrate their magnetic compass based on the skylight 309 polarisation pattern. It can be speculated that this pattern might be due to the inclusion of birds in different phases of 310 migration (see e.g. Wiltschko et al. 2008a), but this seems unlikely as dunlins were tested well after the beginning of 311 their migration (Cramp and Simmons 1983). The heterogeneity observed in the responses to the cue-conflict might be 312 ascribe also to population specific differences (i.e. birds breeding at different latitudes with different reliability of 313 magnetic cues - Muheim et al. 2003; Åkesson and Bianco 2015) or to the previous experience of tested individuals, as 314 the ecological context and the regional availability of orientation cues may affect the use of various compasses 315 (Sandberg and Moore 1996; Chernetsov et al. 2011; Giunchi et al. 2015). Interestingly, the same pattern was not evident 316 in curlew sandpipers, although even in this species the individual variability was not negligible. While we acknowledge 317 that the sample size was small, these inter-specific differences may suggest an effect of different migratory periods (but 318 see Chernetsov et al. 2011), even though possible inter-specific differences in the way the different cues are integrated 319 might be considered. In addition, we cannot excluded an effect of the proximity of the coastline, as this could have also 320 affected the directional preferences of curlew sandpipers during the post-CC test, which may have been responsible for 321 the lack of an effect of the cue-conflict on the majority of tested birds.

In conclusion, the data reported in this study suggest that the variability observed in cue-conflict experiments may be affected by the responses of individuals in prioritising different cues. This individual variability should be taken into account when interpreting the results of cue-conflict studies, particularly with species that tend to live longer than small songbirds, and therefore, have more opportunities to learn from previous migrations.

326

327 Acknowledgments

| 328 | We would like to thank all the people who helped us in the field: F. Chini, S. Formento, M. Ricciardi, S. Pardini, S. |
|-----|---|
| 329 | Volpi and all the ringers working in the ringing station of the "Osservatorio F. Caterini" in San Rossore, and in |
| 330 | particular R. Gambogi and A. Galardini. The comments of three anonymous reviewers greatly improved an |
| 331 | earlier draft of the manuscript. This paper have been submitted to ProofReading-Service.com for editing and |
| 332 | proofreading. |
| 333 | |
| 334 | Ethical statement |
| 335 | Ethical approval: All protocols performed in studies involving animals complies with the ethical standards and Italian |
| 336 | laws on animal welfare. All procedures involving animals were approved by the Italian Istituto Superiore per la |
| 337 | Protezione e la Ricerca Ambientale (ISPRA). |
| 338 | |
| 339 | Funding: This study was partly funded by the Italian Ministero dell'Istruzione, dell'Università e della Ricerca (MIUR - |
| 340 | PRIN 20083ML4XC). |
| 341 | |
| 342 | Conflict of Interest: The authors declare that they have no conflict of interest. |
| 343 | |
| 344 | Informed consent: Informed consent was obtained from all individual participants included in the study. |
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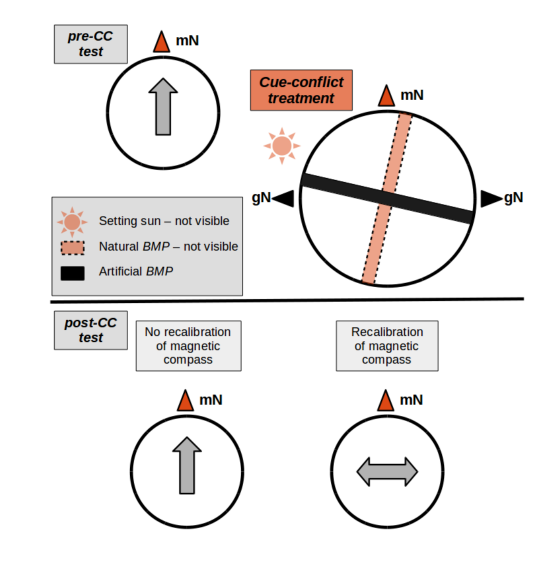
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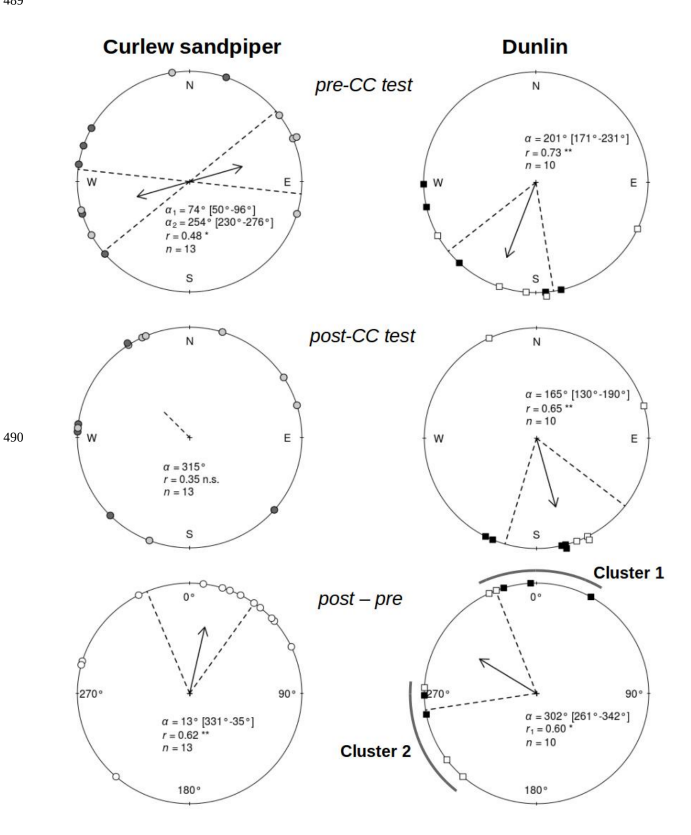
470 Fig. 1 Scheme of the expectations from the cue-conflict experiments with curlew sandpipers, with an expected direction 471 of north. The expectations for dunlins were similar, however, the expected direction was south instead of north. The 472 direction of the magnetic north and geographic north are represented by orange and black triangles, respectively. See 473 the Materials and Methods section for further details

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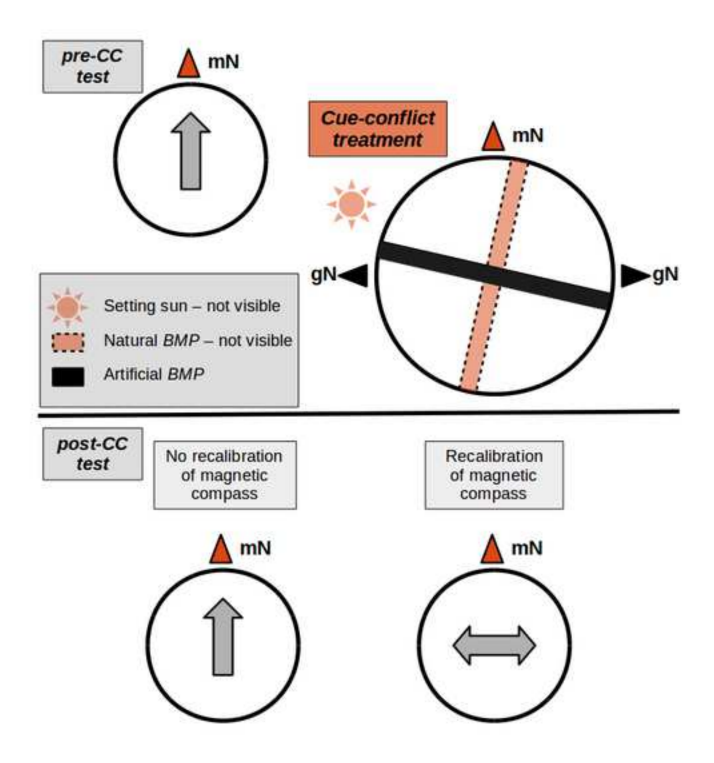
475 Fig. 2 Heading distribution of curlew sandpipers and dunlins prior (pre-CC test) and after (post-CC test) the cue conflict 476 and distribution of the angular differences in the heading of individuals between post-CC test and pre-CC test (post – 477 pre). In curlew sandpipers, different colours were used to identify birds tested in the first (20 minutes before -20478 minutes after the civil twilight; light grey circles) or in the second experimental period (30 minutes - 70 minutes after 479 the civil twilight; dark grey circles). Open and filled circles in dunlins were used to identify adults and juveniles, 480 respectively. Cluster 1 and 2 correspond to the clusters identified by the model based clustering approach (see Materials 481 and Methods and Table S1). The mean vector (α) of each distribution is represented by an arrow, the length (r) was drawn relative to the radius of the circle = 1. The significance is indicated by * p < 0.05, ** p < 0.01, *** p < 0.001, and 482 483 n.s. p > 0.05 as determined by the Rayleigh test. The 95% confidence intervals (broken lines) were reported only for 484 significantly oriented samples

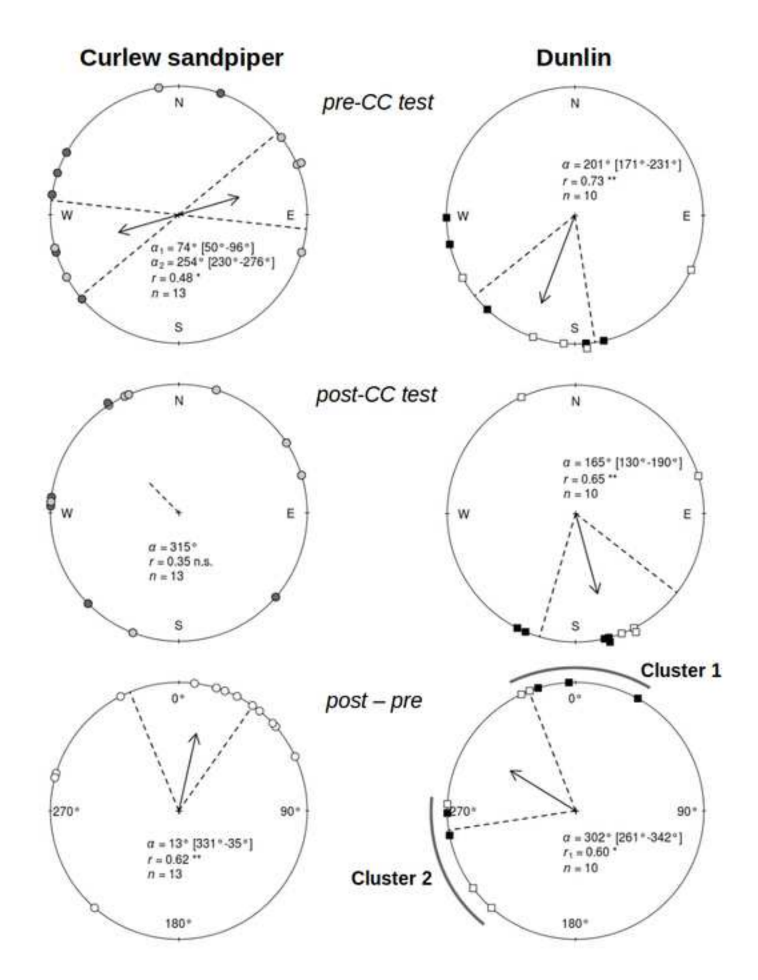




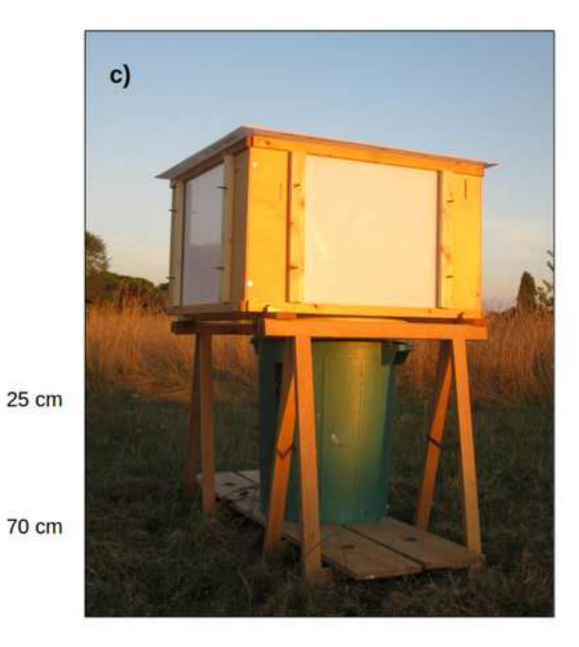


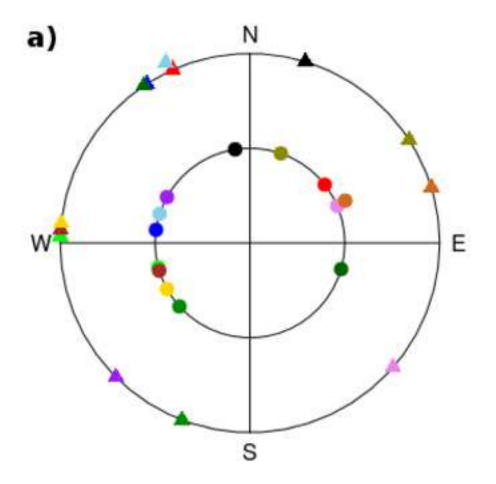


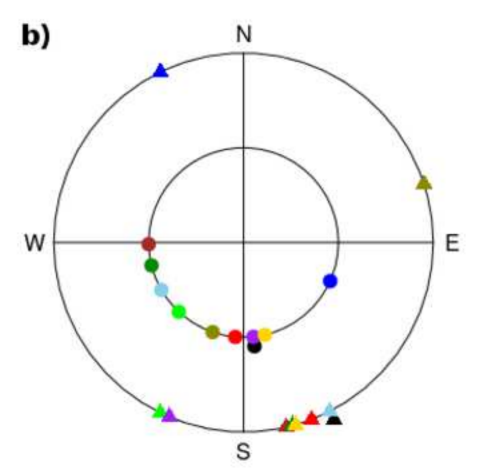




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