

1 **Moonlight Polarisation Pattern Guides Nocturnal Bull Ants Home**

2
3 Cody A Freas*, Ajay Narendra; Trevor Murray; Ken Cheng

4
5 School of Natural Sciences, Macquarie University, Sydney, NSW Australia

6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25 * Corresponding Author

26
27
28 Address for correspondence:
29 Cody Freas
30 School of Natural Sciences
31 Macquarie University
32 Sydney, NSW 2113 Australia
33 Email: cody.freas@mq.edu.au

34
35
36
37
38
39
40
41
42
43
44

45 **Abstract**

46 Celestial cues are used by many animals to orient and navigate their environment.
47 The sun or moon can provide directional information via their position; however, they
48 can often be obstructed by clouds, canopy or the horizon. Despite being hidden,
49 these bodies can still provide compass information through the polarised light pattern
50 they produce/reflect. Sunlight produces polarised light patterns across the overhead
51 sky as it enters the atmosphere, and polarised light is a well-known compass cue for
52 navigating animals. Moonlight produces a similar pattern, albeit a million times
53 dimmer than sunlight. Polarised moonlight's use has, until now, only been
54 demonstrated for straight-line orientation in nocturnal dung beetles. Here we show
55 the first evidence that polarised moonlight forms part of the celestial compass of
56 navigating nocturnal ants. Nocturnal bull ants leave their nest at twilight and rely
57 heavily on the overhead solar polarisation pattern to navigate. Yet many foragers
58 return home overnight when the sun cannot guide them. We demonstrate that bull
59 ants use polarised moonlight to navigate home during the night, by altering the
60 ambient polarisation pattern above homing ants, who alter their headings in
61 response. Furthermore, these ants can detect this cue throughout the lunar month,
62 even under crescent moons, when polarised light levels are at their lowest. Finally,
63 we present evidence that the polarised moonlight pattern is being incorporated
64 throughout the night into these ants' path integration system for homing, as polarised
65 sunlight is incorporated throughout the day.

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85 **Keywords:** ant navigation, lunar cues, route maintenance, cue detection, polarised
86 light, vector

87

88

89 **Introduction**

90 Many navigating animals attend to the position of the sun or moon to guide their
91 movement (Jander 1957; Koltz and Reid 1993; Perez et al. 1997; Dacke et al. 2014;
92 Warrant and Dacke 2016; Freas and Cheng 2022). Yet these celestial bodies are not
93 always directly visible, often obscured by clouds, the canopy or passing below the
94 horizon, resulting in gaps for navigation relying solely upon direct visual detection.
95 Animals hence rely on the pattern of polarised skylight which is accessible when the
96 celestial bodies are occluded to some extent (Horváth et al., 2014). Polarised light
97 comprises light waves which occur along a single plane and are produced as a by-
98 product of light passing through the upper atmosphere (Horváth & Varjú 2004;
99 Horváth et al., 2014). E-vectors are arranged in concentric circles around the
100 sun/moon with the maximum degree of polarisation located 90° from the source.
101 When the sun/moon is near the horizon, the pattern of polarised skylight is
102 particularly simple with uniform direction of polarisation approximately parallel to the
103 north-south axes (Dacke et al., 1999, 2003; Reid et al. 2011; Zeil et al., 2014). The
104 pattern's stability makes the sky's polarisation a useful directional cue for orientation
105 (Wehner and Müller 2006; Reid et al. 2011; Leebhardt and Ronacher 2013; Warrant
106 and Dacke 2016; Freas et al. 2017a, 2019), which insects detect through specialised
107 photoreceptors located in the dorsal rim area of their eyes (Labhart and Meyer 1999;
108 Homberg and Paech 2002; el Jundi et al. 2015). Like solar polarisation, though a
109 million times dimmer, the moon reflects sunlight, producing a polarised moonlight
110 pattern emanating from the moon's position in the night sky (Gál et al. 2001). Given
111 that the moon creates a much dimmer version of the polarisation pattern formed
112 around the sun, night-navigating insects with eyes highly specialised for low light
113 detection may be able to rely on this pattern to orient and navigate to goals.

114 Currently, only nocturnal dung beetles (*Scarabaeus satyrus* and *S.*
115 *zambesianus*) are known to attend to moonlight polarisation patterns during their
116 movement (Dacke et al. 2003, 2004, 2011; Foster et al. 2019). Yet interestingly,
117 these beetles do not use moonlight to navigate, instead relying on this pattern to
118 keep moving straight in order to roll their dung balls expeditiously away from a
119 central dung pile. As such, this cue has only been documented for heading
120 maintenance over short periods. While it has been theorised that this ability to detect
121 the much dimmer polarisation pattern produced by the moon may be present across
122 nocturnal insects more broadly, including nocturnal bees and crickets (Herzmann
123 and Labhart 1989; Greiner et al. 2007; Rost and Honegger 1987), there is currently
124 no concrete behavioural evidence for its use in goal-directed navigation.

125 The large-eyed *Myrmecia* ants have several species that restrict majority of
126 their navigation to evening twilight (outbound) and morning twilight (inbound)
127 respectively (Narendra et al., 2017). We know that two well-studied nocturnal ant
128 species, *Myrmecia pyriformis* and *Myrmecia midas*, use the overhead solar polarised
129 light pattern, which is still visible during the twilight period to derive compass
130 information (Reid et al. 2011; Freas et al. 2017ab; Freas and Cheng 2018). Because
131 the information required for visual navigation degrades beyond twilight, it has been
132 suggested that animals tend to be less active at night. However, a small proportion of

133 *M. pyriformis* foragers leave the nest (10.7% of daily foraging force) or return home
134 during night (13.3% of daily foraging force) (Reid et al., 2013). In *M. midas*, this
135 nocturnal activity is even more pronounced with a majority of the foraging force
136 returning during night (62.8% of daily foraging force) along with a minority of foragers
137 leaving the nest during night (26.2% of daily foraging force) (Freas et al. 2017a).

138 Nocturnal bull ants navigate using a combination of learned visual cues and
139 homing vectors obtained by integrating pedometric and celestial compass
140 information. This true nocturnal navigation is likely aided by the increase in the light
141 intensity provided by the moon's presence. In *M. pyriformis*, on 'full-moon' nights
142 there was a significantly greater proportion of foragers leaving the nest at night
143 compared to a 'new-moon' night (Reid et al., 2013). This additional light at night may
144 enhance terrestrial visual features foragers have learned and therefore assist in
145 visual guidance. In addition, the moon and the lunar polarisation pattern also likely
146 provides compass information, allowing foragers acquire or follow a homing vector.
147 Several arthropods, including ants and bees directly track the moon's position to
148 obtain compass information (Jander 1957; Koltz and Reid 1993; Dacke et al. 2004;
149 Ugolini et al. 2003). But given the moon may be occluded by cloud or overhanging
150 canopy we aim to identify here whether the lunar polarised skylight can also be used
151 by ants for homing.

152 In the current study, we provide the first evidence of polarised moonlight
153 detection and guidance in ants for goal-directed navigation. We studied foraging
154 ants' ability to orient during lunar twilight, by placing and rotating a linear polarising
155 filter over them as they returned to the nest (Figure 1A). We also explored whether
156 these navigators weigh their attendance to polarised moonlight across the lunar
157 calendar, since during quarter-moon and crescent-moon nights, smaller portions of
158 the moon's surface reflect light (as well as moonless nights). Finally, we
159 characterised changes in weighting that these ant navigators give polarised
160 moonlight, as a function of the moon's consistency in the night sky (waxing vs.
161 waning) and the length of their accumulated path integrator, which should increase
162 the weight given to celestial compass portion of the path integrator when in conflict
163 with terrestrial visual cues (Burkhalter 1972; Narendra 2007; Wystrach et al. 2015;
164 Freas and Cheng 2019). During path integration, ant navigators incorporate
165 movement changes, coupling pedometer and celestial compass estimates to
166 generate a spatial estimate their position relative to their nest (Wehner and
167 Srinivasan 2003).

168

169

170

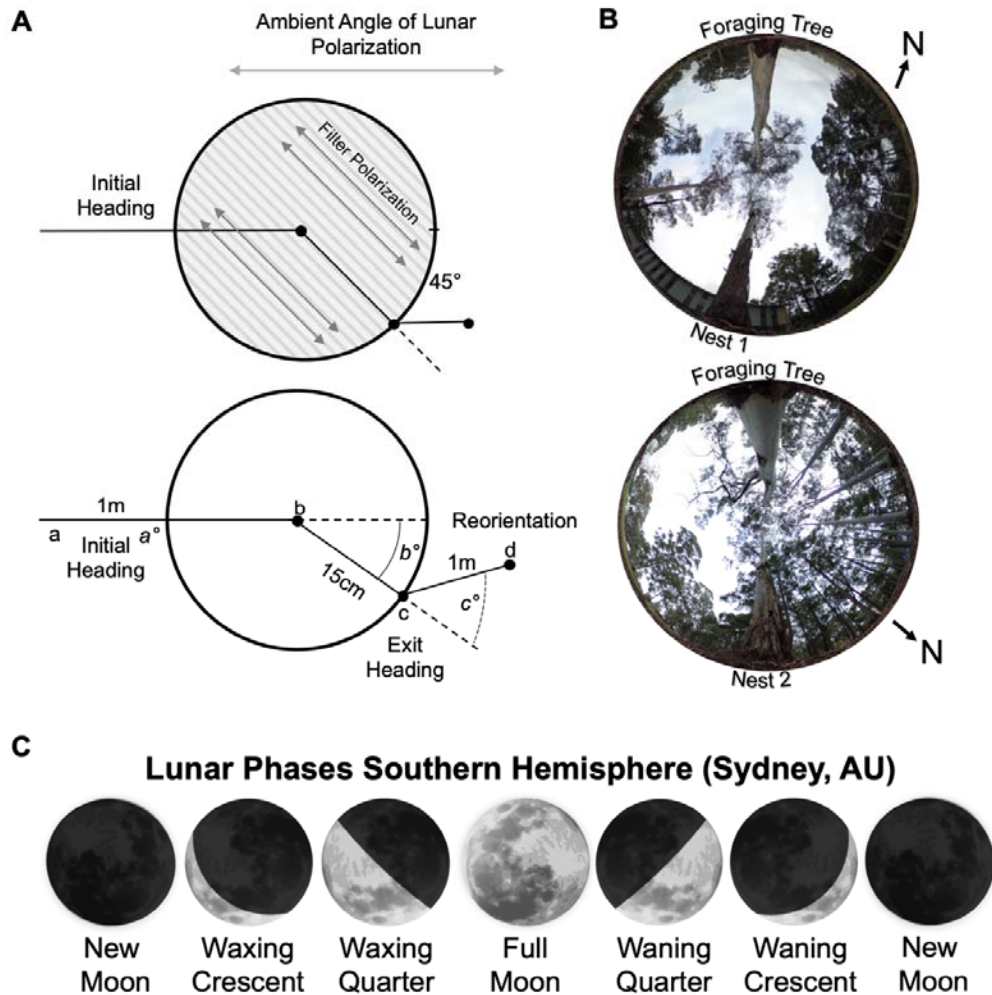
171

172

173

174

175



176
177
178
179
180
181
182
183
184
185
186
187
188
189
190

Figure 1. Diagram of the polarisation filter and changes to the ambient lunar polarisation pattern. (A) During the inbound journey, a linear polarisation filter was placed over the forager, rotating the overhead e-vector by $\pm 45^\circ$. Panel depicts the positional measurements recorded during testing. Initial orientation routes were measured from the foraging tree release point (a) to when the polarisation filter was placed over each forager (b). Exit orientations were measured from the filter centre (b) to the forager's exit location at the filter edge (c). Route directions under the filter (b°) were calculated from the forager's initial route direction zeroed. Reorientations were measured from the forager's exit location from the polarisation filter (c) to the forager's path 1m after exit (d). Reorientation directions (c°) were calculated from the under-filter route direction zeroed. (B) Images of the sky and canopy cover at both nests. Photos were taken at the on-route midpoint between the foraging and nest trees. (C) Lunar phases denote the sunlit part of the moon's surface and where this area is increasing (Waxing) or decreasing (Waning). The lunar phase cycle repeats every 29.5 days. moon images are public domain art accessed through wiki commons (<https://commons.wikimedia.org/>).

191
192
193
194
195
196
197

Methods

Study site

Experiments were conducted from April through October 2023 on two *Myrmecia midas* nests on the Macquarie University campus in Sydney, Australia ($33^\circ 46' 11''\text{S}$, $151^\circ 06' 40''\text{E}$). *M. midas* nests are typically located within stands of *Eucalyptus* trees with the nest entrance located near ($<30\text{cm}$) a tree. *M. midas* is nocturnal, with foraging onset occurring $\sim 20\text{min}$ after sunset when foragers leave the nest to travel

198 to and up one of several surrounding foraging trees overnight (Freas et al. 2018).
199 Inbound navigation is more variable with foragers returning to the nest entrance
200 overnight and into morning twilight (Freas et al. 2017a). For this study, we chose two
201 nests under open canopies to ensure foragers had unobstructed visual access to the
202 overhead sky (Figure 1B). The understory of these areas is naturally barren of
203 vegetation, and we cleared the foraging column of debris to aid visual tracking.

204

205 *Apparatus: Polarisation filter*

206 For each condition, we altered the ambient pattern of polarised moonlight by
207 rotating a linear polarisation filter (30cm diameter; same apparatus used in Freas et
208 al. 2019) above each ant along their inbound journey. This rotation alters the
209 dominant ambient e-vector direction of the sky above the navigator. This filter was
210 held by a circular 1cm thick wooden ring and lifted 10cm off the ground by four
211 equally spaced thin metal legs (Figure 1). All testing was conducted overnight before
212 morning solar twilight, during the lunar twilight for each tested lunar phase. For each
213 night we obtained the moon's position as it reached the horizon based on the
214 Astronomical Almanac (<http://asa.usno.navy.mil>) and set the ambient lunar e-vector
215 perpendicular from the moon's position at moonset. We relied on a digital compass
216 mobile application (Apple™) to locate the ambient e-vector and rotated the linear
217 polariser by $\pm 45^\circ$ from this direction for each ant.

218 Across all conditions, we recorded four positions: the release point, their
219 position when the filter was placed overhead, the filter exit point, and reorientation
220 after ~1m, taking care to slowly follow the ant and mark their positions so as to not
221 disturb their travel. These positions determine each forager's initial orientation, exit
222 orientation and reorientation directions (Figure 1). After testing, each forager was
223 marked with acrylic paint (Tamiya™) to prevent retesting. Testing was conducted at
224 distinct lunar phases which predicably occur throughout the lunar month cycle (29.5
225 days).

226

227 *Full moon*

228 For full moon testing we chose nights during which the waxing moon's lunar phase
229 was near full but with clear separation between solar and lunar twilights. Testing on
230 true full moon nights is difficult as solar and lunar twilights fully overlap and the solar
231 polarised light pattern would overpower the lunar counterpart. Testing during the
232 nights preceding the full moon (waxing phase) ensured the moon's presence in the
233 night sky overnight and testing occurred on nights in which the lunar twilight (1am –
234 4am) was clearly separated from the start of morning solar twilight (5:22am) with
235 illumination above 80% of the lunar surface.

236 Outbound *M. midas* foragers from two nests were followed as they left the
237 nest during evening twilight and collected as they climbed onto their foraging tree,
238 (Nest 1: 6.0m; Nest 2: 3.1m from the nest entrance). Each forager was provided a
239 small amount of honey and held within a clear plastic phial on the ground 5m from
240 the foraging tree with an unobstructed view of the sky (Figure 1). Foragers were held
241 in these phials overnight until the moon's position was within $\pm 10^\circ$ of the horizon

242 (large stands of trees and buildings near the western horizon occluded the moon's
243 position during all testing; Figure 1).

244

245 *Waxing Lunar phase*

246 We observed clear shifts in forager headings at both nests under *Full Moon*
247 conditions, yet nights with over 80% lunar illumination only account for nine nights
248 per lunar cycle. To assess if polarised moonlight can be used throughout the lunar
249 month, we tested foragers in three further conditions representing distinct lunar
250 phases: a *Waxing Quarter Moon*, a *Waxing Crescent Moon*, and a *No Moon* control.
251 For the *Quarter Moon* and *Crescent Moon* conditions, we tested ants identically to
252 full moon conditions; for overnight testing, however, the moon has a different
253 temporal period when it is positioned near the horizon (12am and 9pm, respectively).

254 The procedure was slightly modified for *No Moon* testing as we did not test
255 these foragers on the new moon night (the new moon is only present during the
256 day). We hypothesised that foragers with no available ambient polarisation and
257 suddenly presented with an e-vector might fall back on a memory of the morning
258 solar e-vector as many foragers return during morning twilight and this direction
259 remains consistent across nights. In order to test on a night when there were distinct
260 directional differences between the lunar and morning solar e-vectors (Lunar e-
261 vector: 329°; Morning solar e-vector: 7°; Evening Solar e-vector: 353°), we chose to
262 test on a quarter moon night when the moon was well below the horizon. Testing
263 commenced at 9:00pm, when the moon was 30° below the horizon and we rotated
264 the filter around to the future lunar e-vector direction (moonrise at 12:39am). If
265 foragers were relying on a solar e-vector memory, we expected to see unequal shifts
266 between $\pm 45^\circ$ (smaller shifts in the $+45^\circ$ condition and larger shifts in the -45°
267 condition).

268

269 *Waning Lunar Phases*

270 While there is a consistent presence of polarised light during the waxing phase,
271 during waning lunar phases there is a gap which may impede this pattern's use as a
272 compass cue. Testing during the waxing lunar phase, as the moon's illuminated
273 surface increases, corresponds with periods in which the moon rises prior to sunset
274 and sets overnight. This creates a consistent presence of polarised light (solar then
275 lunar) that foragers could use to continuously update their celestial compass. In
276 contrast, the waning lunar phase corresponds with the moon rising overnight, leading
277 to a gap in this cue as solar twilight ends. We hypothesised that the absence of the
278 moon's presence as a cue overnight might decrease its weighting or degrade its
279 integration into the celestial compass. We added two conditions to test this
280 hypothesis: a *Waning Full Moon* and *Waning Quarter Moon* conditions. Foragers
281 were tested identical to previous conditions; only, they were tested during moonrise
282 (10pm and 1am) overnight rather than moonset.

283

284 *Vector testing*

285 After noticing differences in heading shift magnitude between nests which correlated
286 with PI vector lengths, we tested the hypothesis that, similar to solar polarisation
287 (Freas et al. 2017b), the navigator's accumulated vector length impacts orientation to
288 rotated lunar polarised light. While foragers at Nest 1 (6.0m vector) reoriented fully,
289 those at Nest 2 (3.1m vector) reoriented only halfway on near-full-moon nights. To
290 test this hypothesis we collected foragers of Nest 1 on near (waxing) full moon nights
291 and tested them at a site close (2m) to their nest tree with either a 2m remaining
292 vector, or with a larger (~6m) full corridor remaining vector (*Halfway Collect &*
293 *Release* or *Halfway Release* conditions). In the *Halfway Collect & Release* condition,
294 outbound foragers were collected at the halfway point, held overnight and released
295 at this site to be tested ($\pm 45^\circ$ rotations) at ~2m from the nest. In the *Halfway Release*
296 condition, we followed outbound foragers to their foraging tree, collected and held
297 them until lunar twilight (2-4am) identically to previous full moon conditions. These
298 foragers were released along the route at the halfway point from the nest/foraging
299 tree and we tested each forager with their vector state near full but their true location
300 close (~2m) to the nest tree. In both conditions, we again recorded the initial
301 orientation, filter exit orientation and post-filter reorientation of each forager.

302

303 *Statistical analysis*

304 Data were analysed with circular statistics with the statistics package Oriana
305 Version 4 (Kovach Computing Services). Each ant had a slightly different inbound
306 heading due to their stereotypical route along the foraging corridor and we corrected
307 this variance by designating the initial headings (pre-filter) as 0° for calculating shift
308 magnitudes under the filter. To assess shift magnitude between -45° and $+45^\circ$
309 foragers within conditions, we calculated the mirror of shift in each -45° condition,
310 allowing shift magnitude comparisons within each condition. As -45° and $+45^\circ$ shifts
311 were not significantly different in any condition (Watson-Williams F-tests), they were
312 combined for between condition comparisons. Within-individual comparisons (Initial
313 Orientation vs. Filter Exit and Filter Exit vs. Reorientation) were analysed using
314 Moore's Paired Tests. Across-condition shift magnitudes were analysed using
315 Watson-Williams F-tests. In the lunar phase comparisons where full, quarter and
316 crescent shift magnitudes, as well as Waning and Waxing phases were compared,
317 Holm-Bonferroni corrections were applied to the p -value to account for multiple
318 comparisons.

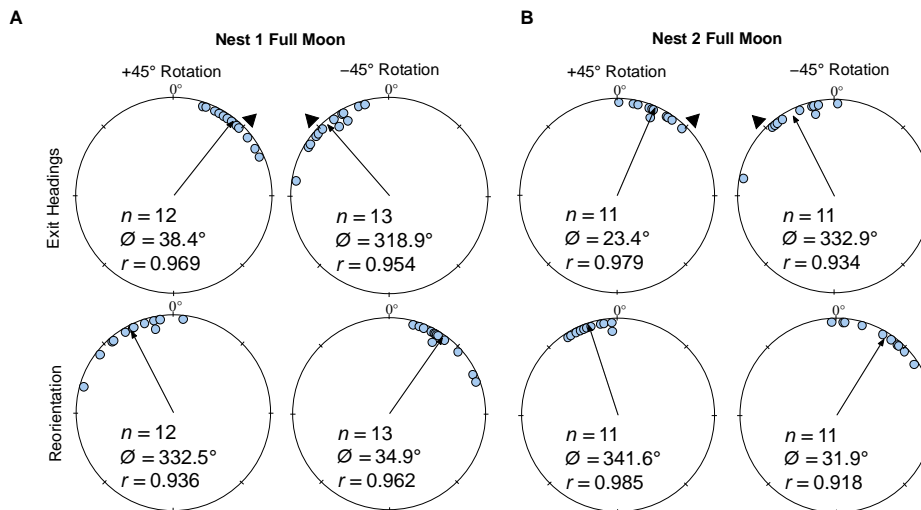
319

320 **Results**

321 *Full moon testing*

322 At both nests, when the polariser was rotated clockwise ($+45^\circ$), exit orientations
323 were shifted to the right of initial headings (mean \pm s.e.m. Nest 1: $38.4 \pm 4.8^\circ$; Nest 2:
324 $23.4 \pm 4.2^\circ$; Figure 2AB), and these changes were significant (Moore's Paired Test,
325 Nest 1: $R=1.639$, $p<0.001$; Nest 2: $R=1.592$, $p<0.001$). Forager headings also
326 changed predictably when the filter was rotated counter-clockwise (-45°) with exit
327 orientations to the left of initial headings (Nest 1: $-41.1 \pm 5.5^\circ$; Nest 2: $-27.1 \pm 7.4^\circ$;
328 Figure 2AB), and these changes were significant (Moore's Paired Test, Nest 1:

329 $R=1.794$, $p<0.001$; Nest 2: $R=1.310$, $p<0.01$). After exiting the $+45^\circ$ or -45° rotated
 330 filter, foragers reoriented significantly to the left (Moore's Paired Test, Nest 1:
 331 $R=1.598$, $p < 0.001$; Nest 2: $R=1.383$, $p<0.005$) or right respectively (Moore's Paired
 332 Test, Nest 1: $R=1.604$, $p<0.001$; Nest 2: $R=1.328$, $p<0.005$). Shift magnitudes did not
 333 differ between $+45^\circ$ and -45° conditions (Watson-Williams F-test, Nest 1:
 334 $F_{(1,23)}=0.155$, $p=0.697$; Nest 2: $F_{(1,20)}=0.234$, $p=0.634$). Combined shift magnitudes
 335 were significantly larger at Nest 1 compared to Nest 2 (Watson-Williams F-test,
 336 $F_{(1,45)}=8.672$, $p=0.005$), exhibiting shift magnitudes near the full e-vector change
 337 ($39.7\pm 3.2^\circ$) while foragers at Nest 2 only exhibited shift magnitudes at about half the
 338 45° rotation ($25.2\pm 3.7^\circ$).
 339



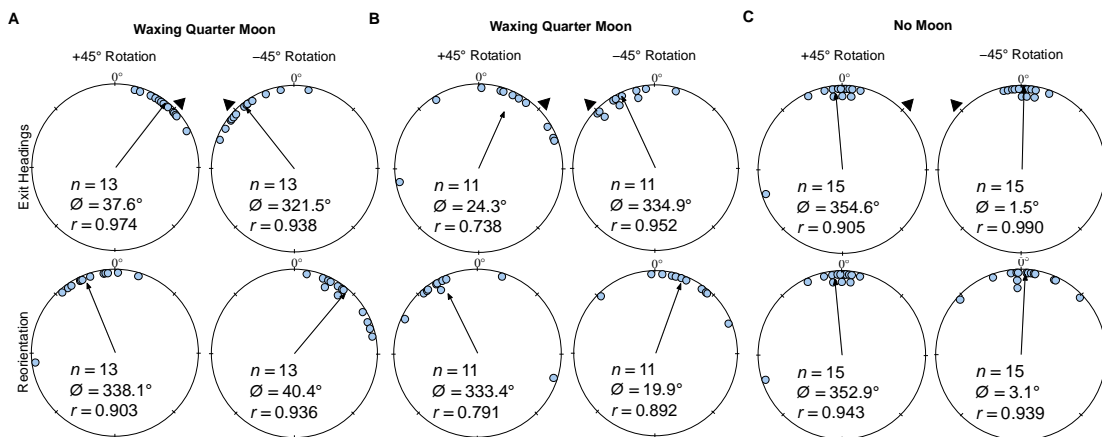
340
 341 **Figure 2.** Circular distributions of headings during the original full moon conditions. In both conditions,
 342 testing occurs in the nights preceding the full moon (illumination > 80%) with the moon waxing.
 343 Circular plot shifts show the exit orientations of individual foragers from their initial headings while the
 344 reorientation represents the change in headings 1m after exiting the filter. Triangles denote the $\pm 45^\circ$
 345 e-vector rotation. The arrow denotes the length and direction of the mean vector. (A) Nest 1 foragers,
 346 5m from the nest (6.0m foraging route). (B) Nest 2 foragers, 2m from the nest (3.1m foraging route).
 347 n , number of individuals; $\bar{\theta}$, mean vector; r , length of the mean vector.
 348
 349

350 Waxing Lunar phases

351 Under a *Waxing Quarter Moon* (lunar illumination ~50%), when the e-vector
 352 was rotated clockwise ($+45^\circ$), exit orientations were again significantly shifted
 353 (Moore's Paired Test, $R=1.787$, $p<0.001$) to the right of their initial heading
 354 (mean \pm s.e.m.; $37.6\pm 4.1^\circ$; Figure 3A). Forager headings were similarly
 355 significantly altered (Moore's Paired Test, $R=1.734$, $p<0.001$) when the overhead e-
 356 vector was rotated counter-clockwise (-45°) with exit orientations to the left of initial
 357 headings (mean \pm s.e.m. $-38.5\pm 6.4^\circ$; Figure 3A). After exiting the filter in both
 358 conditions, foragers reoriented significantly back to the ambient lunar e-vector either
 359 to the left ($+45^\circ$: Moore's Paired Test, Nest 1: $R=1.616$, $p<0.001$) or right (-45° :
 360 Moore's Paired Test, Nest 1: $R=1.664$, $p<0.001$) of their filter exit heading direction.
 361

362 Foragers continued to show evidence of attending to the lunar polarisation
 pattern even under a *Waxing Crescent Moon* (lunar illumination ~20%). Here, when

363 the ambient e-vector was rotated clockwise (+45°), exit orientations were again
 364 significantly shifted (Moore's Paired Test, $R=1.175$, $p<0.025$) to the right of their initial
 365 heading (mean \pm s.e.m.; $29.3\pm 9.9^\circ$; Figure 3B). Forager headings were similarly
 366 significantly altered (Moore's Paired Test, $R=1.39$, $p<0.005$) when the overhead e-
 367 vector was rotated -45° counter-clockwise, with exit orientations to the left of initial
 368 headings (mean \pm s.e.m.; $-27.6\pm 4.7^\circ$; Figure 3B). After exiting the filter in both
 369 conditions, foragers reoriented significantly back to the ambient lunar e-vector either
 370 to the left (+45°: Moore's Paired Test, Nest 1: $R=1.324$, $p<0.005$) or right (-45° :
 371 Moore's Paired Test, Nest 1: $R=1.223$, $p<0.025$) of their filter exit heading direction.
 372 Shift magnitudes were not significantly different between $\pm 45^\circ$ conditions (Watson-
 373 Williams F-test, $F_{(1,20)}=0.03$, $p=0.863$).
 374



375 **Figure 3.** Circular distributions of headings during (A) Waxing Quarter Moon, (B) Waxing Crescent
 376 Moon and (C) No Moon conditions. Circular plot shifts show the exit orientations of individual foragers
 377 from their initial headings while the reorientation represents the change in headings 1m after exiting
 378 the filter. Triangles denote $\pm 45^\circ$ e-vector rotation. The arrow denotes the length/direction of the mean
 379 vector. n , number of individuals; $\bar{\theta}$, mean vector; r , length of the mean vector.
 380

381
 382 When no ambient lunar e-vector was present and the polariser was rotated
 383 either clockwise (+45°) or counter-clockwise (-45°), foragers did not significantly
 384 alter their paths under the filter (+45°: Moore's Paired Test, $R=0.226$, $p>0.50$;
 385 mean \pm s.e.m.: $-5.4\pm 7.4^\circ$; -45° : Moore's Paired Test, $R=0.650$, $p>0.10$;
 386 mean \pm s.e.m.: $0.7\pm 7.6^\circ$; Figure 3C). Foragers also did not significantly reorient
 387 after exiting the filter (+45°: Moore's Paired Test, $R=0.294$, $p>0.50$; mean \pm s.e.m.:
 388 $-7.1\pm 5.6^\circ$; -45° : Moore's Paired Test, $R=0.611$, $p>0.10$; mean \pm s.e.m.: $1.8\pm 5.1^\circ$).
 389 Shift magnitudes were not significantly different between $\pm 45^\circ$ conditions (Watson-
 390 Williams F-test, $F_{(1,24)}=0.016$, $p=0.899$).

391 When comparing shift magnitude across lunar phases, Full Moon foragers
 392 were not significantly different from either the Waxing Quarter Moon or Waxing
 393 Crescent Moon (Watson-Williams F-test, $p>0.05$; Figure 5).

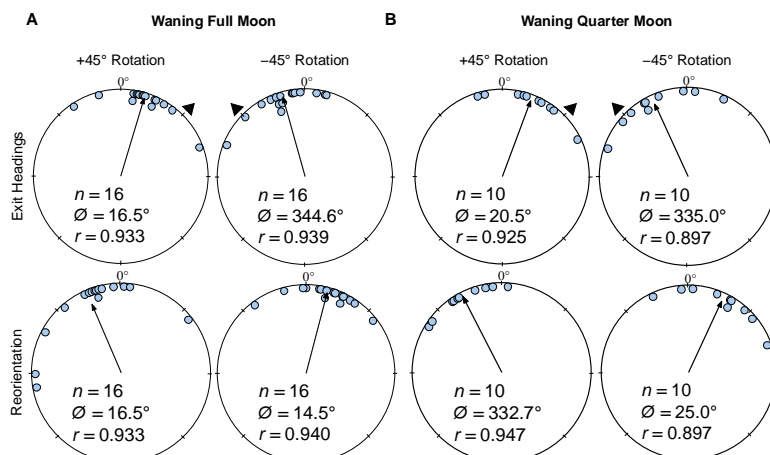
394 Waning Lunar Phase

395 Under a Waning Full Moon, when the ambient e-vector was rotated clockwise (+45°),
 396 exit orientations were shifted significantly to the right of initial headings
 397

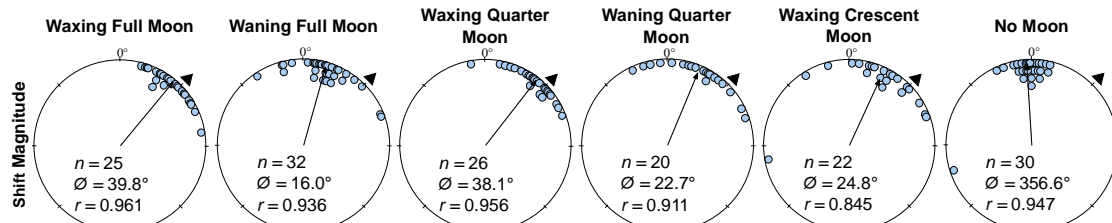
398 (mean \pm s.e.m. = $16.5 \pm 5.3^\circ$; Moore's Paired Test, $R=1.468$, $p<0.005$; Figure 4A).
 399 Headings also shifted significantly to the left of initial headings (mean \pm s.e.m.: $-15.4 \pm 5.1^\circ$; Moore's Paired Test, $R=1.513$, $p<0.001$; Figure 4A) when the overhead e-
 400 vector was rotated -45° counter-clockwise. After exiting the filter in both conditions,
 401 foragers reoriented significantly back to the ambient lunar e-vector either to the left
 402 ($+45^\circ$: Moore's Paired Test, Nest 1: $R=1.548$, $p<0.001$) or right (-45° : Moore's Paired
 403 Test, Nest 1: $R=1.247$, $p<0.025$) of their filter exit heading direction. Shift magnitudes
 404 were not significantly different between the $\pm 45^\circ$ conditions (Watson-Williams F-test,
 405 $F_{(1,30)}=0.022$, $p=0.884$).
 406

407 Under a *Waning Quarter Moon* (lunar illumination $\sim 50\%$), when the ambient e-
 408 vector was rotated clockwise ($+45^\circ$), exit orientations were shifted to the right of
 409 initial headings (mean \pm s.e.m. = $20.5 \pm 8.4^\circ$; Figure 4B), and these changes were
 410 significant (Moore's Paired Test, $R=1.33$, $p<0.005$). Headings also changed
 411 significantly (Moore's Paired Test, $R=1.31$, $p<0.01$) when the overhead e-vector was
 412 rotated counter-clockwise (-45°) with exit orientations to the left of initial headings
 413 (mean \pm s.e.m.: $-22.5 \pm 11.7^\circ$; Figure 4B). After exiting the filter in both conditions,
 414 foragers reoriented significantly back to the ambient lunar e-vector either to the left
 415 ($+45^\circ$: Moore's Paired Test, Nest 1: $R=1.504$, $p<0.001$) or right (-45° : Moore's Paired
 416 Test, Nest 1: $R=1.246$, $p<0.025$) of their filter exit heading direction. Shift magnitudes
 417 were not significantly different between the $\pm 45^\circ$ conditions (Watson-Williams F-test,
 418 $F_{(1,18)}=0.154$, $p=0.700$).
 419

420 When comparing shift magnitudes between Waxing and Waning phases, shift
 421 magnitude was significantly higher in both *Waxing Full Moon* and *Waxing Quarter*
 422 *Moon* foragers when compared to their *Waning* counterparts (39.8° vs. 16.0° and
 423 38.1° vs. 22.7° respectively; Full Moon: Watson-Williams F-test, $F_{(1,55)}=21.62$,
 424 $p<0.001$; Quarter Moon: Watson-Williams F-test, $F_{(1,44)}=5.889$, $p=0.038$; Figure 5).



425 **Figure 4.** Circular distributions of headings during (A) *Waning Full Moon*, (B) *Waning Quarter Moon*
 426 conditions. Circular plot shifts show the exit orientations of individual foragers from their initial
 427 headings while the reorientation represents the change in headings 1m after exiting the filter.
 428 Triangles denote $\pm 45^\circ$ e-vector rotation. The arrow denotes the length/direction of the mean vector. n ,
 429 number of individuals; $\bar{\theta}$, mean vector; r , length of the mean vector.
 430
 431



432

433

Figure 5. Shift magnitudes for lunar phase conditions at Nest 1. Each circular plot shows the $\pm 45^\circ$ combined shifts for each condition. Triangles denote $+45^\circ$ e-vector rotation; data from -45° were mirrored and combined). The arrow denotes the length and direction of the mean vector. n , number of individuals; $\bar{\theta}$, mean vector; r , length of the mean vector.

436

437

438

Vector testing

439

440

441

442

443

444

445

446

447

448

449

450

451

452

453

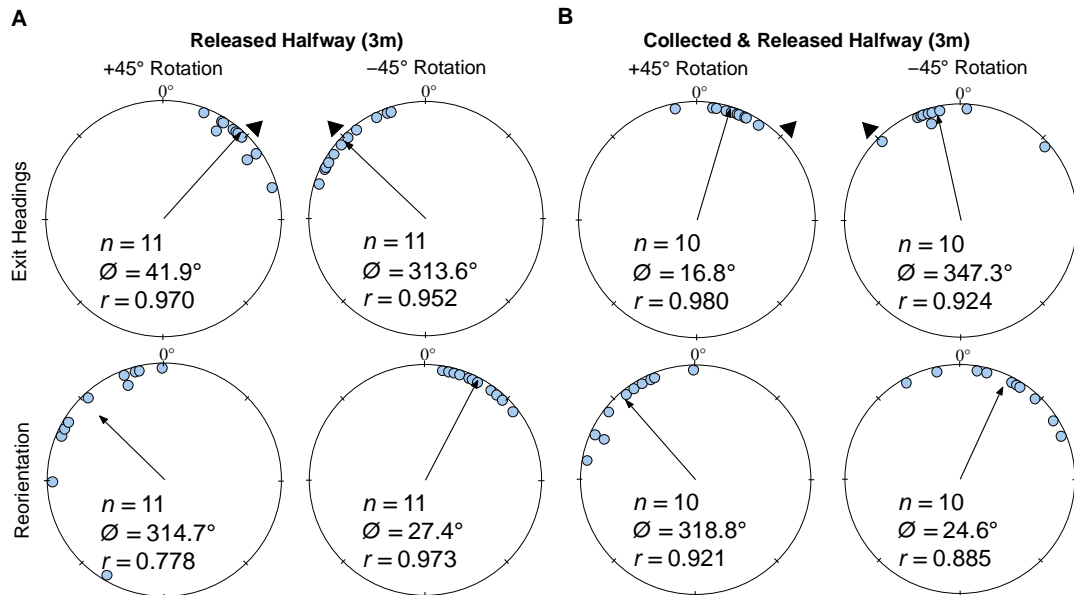
454

455

456

457

Both *Halfway Release* (~6m vector) and *Halfway Collection & Release* (~2m vector) foragers exhibited significant shifts (*Halfway Release*: Moore's Paired Test, $R=1.728$, $p<0.001$; *Halfway Collection & Release*: Moore's Paired Test, $R=1.380$, $p<0.005$) to the right of initial headings (mean \pm s.e.m.: $41.9\pm 4.9^\circ$ and $6.8\pm 4.3^\circ$; Figure 6AB) when the ambient e-vector was rotated clockwise ($+45^\circ$). Forager headings also significantly shifted (*Halfway Release*: Moore's Paired Test, $R=1.664$, $p<0.001$; *Halfway Collection & Release*: Moore's Paired Test, $R=1.07$, $p<0.05$) when the overhead e-vector was rotated counter-clockwise (-45°) with exit orientations to the left of initial headings (mean \pm s.e.m.: $-46.4\pm 6.3^\circ$ and $12.7\pm 8.5^\circ$; Figure 6AB). After exiting the filter in both conditions, foragers reoriented significantly back to the ambient lunar e-vector either to the left ($+45^\circ$: *Halfway Release*: Moore's Paired Test, $R=1.692$, $p<0.001$; *Halfway Collection & Release*: Moore's Paired Test, $R=1.600$, $p<0.001$) or right (-45° : *Halfway Release*: Moore's Paired Test, $R=1.604$, $p<0.001$; *Halfway Collection & Release*: Moore's Paired Test, $R=1.274$, $p<0.01$) of their exit heading direction. Shift magnitude was significantly higher in *Halfway Release* foragers compared to *Halfway Collection & Release* foragers tested at the same location 2m from the nest entrance (44.1° and 14.8° respectively; Watson-Williams F-test, $F_{(1,40)} = 29.105$, $p < 0.001$; Figure 6AB).



458

459

460

461

462

463

464

465

466

467

468

Figure 6. Circular distributions of *M. midas* headings during (A) *Released Halfway*, (B) *Collected & Released Halfway* conditions. Foragers in both conditions were tested at 2m from the nest with *Released Halfway* foragers having a long 6.0m vector and *Collected & Released Halfway* foragers having a 3.1m vector. Circular plots show the exit orientations of individual foragers from their initial headings while the reorientation represents the change in headings 1m after exiting the filter. Triangles denote $\pm 45^\circ$ e-vector rotation. The arrow denotes the length and direction of the mean vector. *n*, number of individuals; $\bar{\theta}$, mean vector; *r*, length of the mean vector.

466

467

468

Discussion

468

469

470

471

472

473

474

475

476

477

478

479

480

481

482

483

484

485

486

487

488

These results constitute the first instance of polarised moonlight use for homing and only the second reported instance of its use for orientation in any animal (Figure 7). *Myrmecia midas* foragers predictably altered their heading directions in response to experimental rotations in the ambient overhead lunar polarisation pattern. This ability to detect and attend to polarised moonlight persisted throughout the lunar cycle, with foragers attending to the pattern even under a crescent moon with ~20% lunar illumination. This indicates that polarised moonlight is detectable across the lunar month, making it a stable cue these ants can use when moving or updating their path integrator overnight.

While they can utilise the lunar polarisation pattern through the lunar cycle, foragers exhibited reduced heading shifts during the waning lunar phases, during which the moon's absence for a portion of the night leads to a cue gap overnight. These reductions in observed shifts are likely due to either decreased weighting or a degradation in the celestial compass due to periods when no celestial cues are alliable. The shift magnitude differences between also point to moonlight polarisation being continuously tracked throughout the overnight period, in line with this celestial cue being integrated into the navigator's path integrator. Further evidence of this cue's integration into the path integrator is illustrated in our *Halfway* testing conditions. Here, foragers with longer home vectors under a full moon responded almost fully to e-vector changes regardless of their distance to the nest, while short

489 vector distances led to less than half the e-vector shift, corresponding with
490 decreases in vector cue strength at small distances (Wystrach et al. 2015). These
491 changes align exactly with how *M. midas*, uses polarised sunlight as part of its
492 vector-based homing during evening and morning twilight (Freas et al. 2017b; Figure
493 7), suggesting that polarised moonlight is detected and integrated into its path
494 integrator along the same visual pathways as sunlight.

495

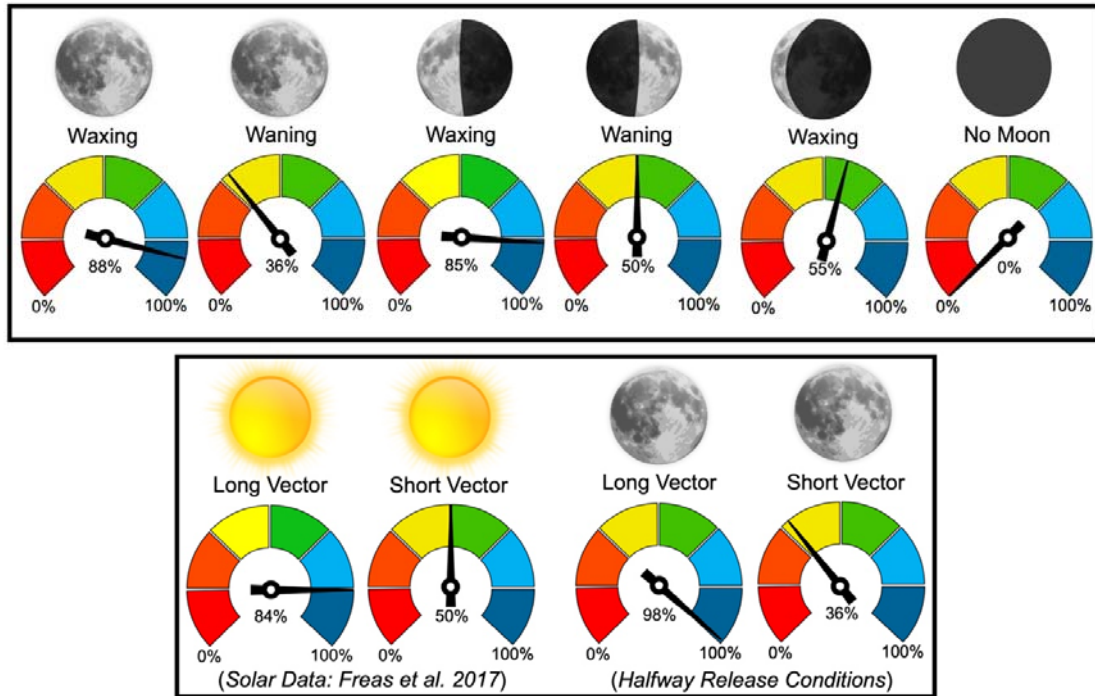
496 *Moonlight vs. Sunlight*

497 The nocturnal bull ants *Myrmecia pyriformis* and *Myrmecia midas* are known to use
498 the solar polarised light pattern during the twilight periods (Reid et al. 2011; Freas et
499 al. 2017ab), yet both species are active after twilight, when solar polarisation cues
500 are absent (Reid et al. 2011, 2013; Freas et al. 2017ab). The observed true nocturnal
501 navigation in these animals could be driven by the moon's presence with *M.*
502 *pyriformis*, showing more foraging activity on 'full-moon' nights (Reid et al., 2013). *M.*
503 *midas* exhibits a high level of overnight activity, with almost half of foragers returning
504 before the morning twilight. *M. midas* also navigates through heavily canopied forest
505 habitat where the moon may be occluded but its polarisation pattern across the sky
506 remains unobstructed. Thus, *M. midas* makes for an interesting species to
507 characterise lunar and solar polarised light detection.

508 While we cannot compare solar and moonlight polarisation navigation in
509 outbound ants (outbound foraging is highly correlated with evening twilight when the
510 solar light would overpower any moonlight polarisation pattern), striking similarities
511 occur when comparing solar and moonlight polarisation navigation in ants homing to
512 the nest. In the solar polarisation study (Freas et al. 2017b), inbound foragers tested
513 during morning twilight at 4–6 m from their nest altered their paths under the filter
514 for almost the full 45° solar e-vector manipulation (–45° rotation: –41.16°/ +45°
515 rotation: 34.13°; Figure 7) but only compensated for around half of the rotation when
516 tested 1–2 m from the nest (–45° rotation: –24.86°/ +45° rotation: 19.73°). We see
517 the same pattern with polarised moonlight with foragers exhibiting near full shifts 5m
518 from Nest 1 (–45° rotation: –41.4°/ +45° rotation: 38.4°) and half shifts 2m from Nest
519 2 (–45° rotation: –27.1°/ +45° rotation: 23.4°; Figure 7).

520 We also observed consistent slight under-estimation in the shifts even when
521 foragers had a long vector. Observations of ants after the filter was placed overhead
522 suggest that heading updates are not immediate, occurring only after the ant travels
523 along its original heading a few centimetres (~5cm). This means that even if the ant
524 fully shifts its heading, the delay will cause our measurements at filter exit to slightly
525 underestimate each individual's position since we measure the angle from where the
526 lunar cue changes (as filter was placed overhead) and not the position at which the
527 ant altered its heading.

528



529
530
531
532
533
534
535

Figure 7. Mean shift magnitudes, $+45^\circ$ and -45° combined, reported as percentages of the 45° e-vector rotation (100% = 45°). Vector data for solar polarised light is reported from Freas et al. 2017, Royal Society Open Science. Sun and moon images are public domain art accessed through wiki commons (<https://commons.wikimedia.org/>).

Moonlight and the Path Integrator

536
537 We also see the same full heading changes to both solar- and moonlight-
538 polarisation-pattern rotations when foragers were released halfway to the nest and
539 tested 2m with a larger vector (Solar: -45° rotation: -35.77° / $+45^\circ$ rotation: 39.42° ;
540 Moonlight: -45° rotation: -46.4° / $+45^\circ$ rotation: 41.9° ; Figure 7). In the *Halfway collect*
541 *& Release* condition, this shift significantly decreased at the same testing location,
542 indicating that the vector length and not the testing site dictated this adjustment in
543 cue weighting. These findings suggest that under the filter, foragers use any
544 available celestial and terrestrial cues that are still visible, yet the weighting of the
545 polarisation pattern appears to change in accordance with the vector state
546 (Burkhalter 1972; Narendra 2007) and not its test location close to the nest tree,
547 which could be a potentially highly salient landmark. This leads to several interesting
548 implications. First, these ants weight the polarisation cue more highly and perform
549 larger heading shifts when their current path integrator distance is longer. These
550 increased shift magnitudes align with the hypothesis that longer accumulated
551 vectors, ants increase the weighting given this cue (Burkhalter 1972; Narendra 2007;
552 Wystrach et al. 2015; Freas et al. 2017b). Secondly, these ants are using polarised
553 moonlight precisely the same way they use solar polarisation, meaning that just as
554 polarised moonlight is likely integrated into the forager's path integrator throughout
555 the night. Moonlight polarised light is likely processed through the same visual

556 pathways as polarised sunlight, meaning that these ants can use the same
557 underlying neural architecture for polarised solar light and polarised moonlight cues.
558 The only limiting factor would be the navigator's detection threshold (at least 20%
559 lunar luminance), which can be accomplished through the specialised eye
560 morphology of these ants (Zeil et al. 2014).

561

562 *Polarised Moonlight and Lunar Phase*

563 Foragers showed clear evidence of detecting and employing polarised moonlight
564 when homing to the nest across the lunar cycle, even on waxing crescent moon
565 nights. This aligns with polarised moonlight's use in dung beetles, with individuals
566 able to maintain their straight-line paths under quarter and crescent moon e-vectors
567 (Dacke et al. 2004). Furthermore, the lack of a shift-magnitude reduction between full
568 and crescent nights suggests no reduction in detection. While we could have
569 continued to test with smaller portions of the moon surface illuminated, a reduction in
570 shift magnitude could result from either physiological limits in detection or 'decisional'
571 processes in how much weight to accord the cue. Behavioural responses and
572 physiological limits cannot be untangled behaviourally, and detection thresholds
573 would require intracellular recordings under dim polarised light. Finally, the lack of
574 shifts with *No Moon* foragers indicates that these navigators do not fall back on
575 memories of the evening or morning solar e-vector when presented one overnight.

576 One unexpected finding was the reduction in shift magnitude under waning
577 moons relative to waxing moons. When we first tested foragers under *Waning Full*
578 *Moons* and *Waning Quarter Moons* they showed clear evidence of attending to the
579 polarisation cue, but shift magnitudes were significantly smaller compared to the
580 *Waxing Moon* conditions. This reducing in shift magnitude suggests that polarised
581 moonlight was being detected but it was weighted weakly, perhaps due to the
582 celestial compass incorporation into the path integrator being interrupted This
583 observed reduction in shift magnitude likely results from the period overnight in
584 which the waning moon was absent from the night sky, meaning foragers could not
585 attend to a consistent polarisation pattern throughout the night. The moonlight
586 pattern only becomes visible in the sky once the moon reaches -18° below the
587 horizon; thus, waning-moon nights present periods when there are neither solar nor
588 lunar cues available to maintain the compass, potentially degrading this estimate.
589 This suggests that when the moon is waxing and present throughout the overnight
590 period (at least until moonset), *M. midas* foragers are continuously tracking it and
591 integrating this compass cue into their path integrator. When there are periods
592 overnight when the moon and its polarisation pattern are absent, it is either weighted
593 weakly or it becomes misaligned with the true homing direction since the celestial
594 compass incorporation into the path integrator has been interrupted with overnight
595 periods with no detectable celestial cues.

596

597 *Conclusions*

598 In the first evidence of polarised moonlight being used to navigate to a goal location,
599 inbound *M. midas* foragers detect and respond predictably to rotations of the

600 moonlight e-vector orientation under a filter and reorient back to the ambient e-vector
601 after filter exit. This ability occurs across lunar phases, suggesting that polarised
602 moonlight is a detectable cue throughout the lunar month. Heading changes due to
603 polarised moonlight align with responses to polarised sunlight as part of the path
604 integrator during solar twilight. This indicates that polarised moonlight is likely
605 detected and integrated into the ant's path integrator for inbound homing along the
606 same visual pathways as polarised sunlight. Reductions in heading shifts due to
607 differences in PI vector lengths, and periods without access to polarised light
608 patterns suggest that these animals are able to weight the information provided by
609 celestial polarised light. In so doing these foragers can cater their navigational
610 decisions proportionate to more closely match the reliability of available navigational
611 information.

612

613 **Acknowledgments**

614 *Funding*

615 This project was funded by a Macquarie University Research Fellowship
616 (MQRF0001094), by Macquarie University, and by an ARC Discovery Grant
617 (DP200102337).

618 *Conflicts of interest*

619 The authors declare no conflicts of interest associated with this work.

620 *Ethics*

621 There are no state or federal governmental regulations guiding the research of
622 invertebrates in Australia.

623

624

625 **References**

- 626 Burkhalter, A. (1972). Distance measuring as influenced by terrestrial cues in *Cataglyphis*
627 *bicolor*. In *Information Processing in the Visual System of Arthropods* (ed. R. Wehner),
628 pp. 303-308. Berlin: Springer.
- 629 Dacke, M., Byrne, M. J., Baird, E., Scholtz, C. H., & Warrant, E. J. (2011). How dim is dim?
630 Precision of the celestial compass in moonlight and sunlight. *Philosophical*
631 *Transactions of the Royal Society B: Biological Sciences*, 366(1565), 697-702.
- 632 Dacke, M., Byrne, M. J., Scholtz, C. H., & Warrant, E. J. (2004). Lunar orientation in a beetle.
633 *Proceedings of the Royal Society of London. Series B: Biological Sciences*, 271(1537),
634 361-365.
- 635 Dacke, M., Nilsson, DE., Warrant, E. et al. Built-in polarizers form part of a compass organ in
636 spiders. *Nature* 401, 470–473 (1999). <https://doi.org/10.1038/46773>
- 637 Dacke, M., Nilsson, D. E., Scholtz, C. H., Byrne, M., & Warrant, E. J. (2003). Insect orientation
638 to polarized moonlight. *Nature*, 424(6944), 33-33.
- 639 Dacke, M., el Jundi, B., Smolka, J., Byrne, M., & Baird, E. (2014). The role of the sun in the
640 celestial compass of dung beetles. *Philosophical Transactions of the Royal Society B:*
641 *Biological Sciences*, 369(1636), 20130036.

- 642 el Jundi, B., Warrant, E. J., Byrne, M. J., Khaldy, L., Baird, E., Smolka, J., & Dacke, M. (2015).
643 Neural coding underlying the cue preference for celestial orientation. In Proceedings of
644 the National Academy of Sciences (Vol. 112, Issue 36, pp. 11395–11400). Proceedings
645 of the National Academy of Sciences.
- 646 Foster, J. J., Kirwan, J. D., El Jundi, B., Smolka, J., Khaldy, L., Baird, E., ... & Dacke, M. (2019).
647 Orienting to polarized light at night—matching lunar skylight to performance in a
648 nocturnal beetle. *Journal of Experimental Biology*, 222(2), jeb188532.
- 649 Freas, C. A., & Cheng, K. (2019). Panorama similarity and navigational knowledge in the
650 nocturnal bull ant *Myrmecia midas*. *Journal of Experimental Biology*, 222(11),
651 jeb193201.
- 652 Freas, C. A., & Cheng, K. (2022). The basis of navigation across species. *Annual review of*
653 *psychology*, 73, 217-241.
- 654 Freas, C. A., Narendra, A., & Cheng, K. (2017a). Compass cues used by a nocturnal bull ant,
655 *Myrmecia midas*. *Journal of Experimental Biology*, 220(9), 1578-1585.
- 656 Freas, C. A., Narendra, A., Lemesle, C., & Cheng, K. (2017b). Polarized light use in the
657 nocturnal bull ant, *Myrmecia midas*. *Royal Society open science*, 4(8), 170598.
- 658 Freas, C. A., Plowes, N. J., & Spetch, M. L. (2019). Not just going with the flow: foraging ants
659 attend to polarised light even while on the pheromone trail. *Journal of Comparative*
660 *Physiology A*, 205(5), 755-767.
- 661 Freas, C. A., Wystrach, A., Narendra, A., & Cheng, K. (2018). The view from the trees:
662 nocturnal bull ants, *Myrmecia midas*, use the surrounding panorama while descending
663 from trees. *Frontiers in psychology*, 9, 16.
- 664 Gál, J., Horváth, G., Meyer-Rochow, V. B., & Wehner, R. (2001). Polarization patterns of the
665 summer sky and its neutral points measured by full-sky imaging polarimetry in Finnish
666 Lapland north of the Arctic Circle. *Proceedings of the Royal Society of London. Series*
667 *A: Mathematical, Physical and Engineering Sciences*, 457(2010), 1385-1399.
- 668 Greiner, B., Cronin, T. W., Ribí, W. A., Wcislo, W. T., & Warrant, E. J. (2007). Anatomical and
669 physiological evidence for polarisation vision in the nocturnal bee *Megalopta genalis*.
670 *Journal of Comparative Physiology A*, 193, 591-600.
- 671 Herzmann, D., & Labhart, T. (1989). Spectral sensitivity and absolute threshold of
672 polarization vision in crickets: a behavioral study. *Journal of Comparative Physiology A*,
673 165, 315-319.
- 674 Homberg, U., & Paech, A. (2002). Ultrastructure and orientation of ommatidia in the dorsal
675 rim area of the locust compound eye. *Arthropod Structure & Development*, 30(4), 271-
676 280.
- 677 Horváth, G., & Varjú, D. (2004). Polarized light in animal vision: polarization patterns in
678 nature. Springer Science & Business Media.
- 679 Horváth, G., Lerner, A., & Shashar, N. (2014). Polarized light and polarization vision in animal
680 sciences (Vol. 2). G. Horváth (Ed.). Berlin: Springer.
- 681 Jander, R. (1957) Die optische Richtungsorientierung der Roten Waldameise (*Formica rufa* L.)
682 *Zeitschrift für vergleichende Physiologie* Bd, 40, 162-238.
- 683 Klotz, J. H., & Reid, B. L. (1993). Nocturnal orientation in the black carpenter ant
684 *Camponotus pennsylvanicus* (DeGeer)(Hymenoptera: Formicidae). *Insectes Sociaux*,
685 40, 95-106.
- 686 Labhart, T., & Meyer, E. P. (1999). Detectors for polarized skylight in insects: a survey of
687 ommatidial specializations in the dorsal rim area of the compound eye. *Microscopy*
688 *research and technique*, 47(6), 368-379.

- 689 Lebahardt, F., & Ronacher, B. (2014). Interactions of the polarization and the sun compass in
690 path integration of desert ants. *Journal of Comparative Physiology A*, 200, 711-720.
- 691 Narendra, A. (2007). Homing strategies of the Australian desert ant *Melophorus bagoti* II.
692 Interaction of the path integrator with visual cue information. *Journal of Experimental*
693 *Biology*, 210(10), 1804-1812.
- 694 Narendra A, Kamhi JF & Ogawa Y. 2017. Moving in dim light: behavioural and visual
695 adaptations in nocturnal ants. *Integrative and Comparative Biology* 57: 1104-1116.
- 696 Perez, S., Taylor, O. & Jander, R. 1997. A sun compass in monarch butterflies. *Nature* 387, 29.
697 <https://doi.org/10.1038/387029a0>
- 698 Reid, S. F., Narendra, A., Hemmi, J. M., & Zeil, J. (2011). Polarised skylight and the landmark
699 panorama provide night-active bull ants with compass information during route
700 following. *Journal of Experimental Biology*, 214(3), 363-370.
- 701 Reid SF, Narendra A, Taylor RW & Zeil J. 2013. Foraging ecology of the night-active bull
702 ant, *Myrmecia pyriformis*. *Australian Journal of Zoology* 61: 170-177.
- 703 Rost, R., & Honegger, H. W. (1987). The timing of premating and mating behavior in a field
704 population of the cricket *Gryllus campestris* L. *Behavioral Ecology and Sociobiology*, 21,
705 279-289.
- 706 Ugolini, A., Galanti, G., & Mercatelli, L. (2013). Do sandhoppers use the skylight polarization
707 as a compass cue?. *Animal Behaviour*, 86(2), 427-434.
- 708 Warrant, E., & Dacke, M. (2016). Visual navigation in nocturnal insects. *Physiology*, 31(3),
709 182-192.
- 710 Wehner, R., & Müller, M. (2006). The significance of direct sunlight and polarized skylight in
711 the ant's celestial system of navigation. *Proceedings of the National Academy of*
712 *Sciences*, 103(33), 12575-12579.
- 713 Wehner, R. and Srinivasan, M.V. (2003). Path integration in insects. In: Jeffery KJ (ed) *The*
714 *neurobiology of spatial behaviour*. Oxford University Press, Oxford, pp 9–30.
- 715 Wystrach, A., Mangan, M., & Webb, B. (2015). Optimal cue integration in ants. *Proceedings*
716 *of the Royal Society B: Biological Sciences*, 282(1816), 20151484.
- 717 Zeil J, Ribí WA, Narendra A. 2014 Polarization vision in ants, bees and wasps. In *Polarized*
718 *light and polarization vision in animal sciences*, 2nd edn (ed. Gábor Horváth), pp. 41–60.
719 Berlin, Germany: Springer Series in Vision Research.