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#### Paper:

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1	Socially informed dispersal in a territorial cooperative breeder							
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### **19 ABSTRACT**

Dispersal is a key process governing the dynamics of socially and spatially structured
 populations, and involves three distinct stages: emigration, transience, and settlement. At each
 stage, individuals have to make movement decisions, which are influenced by social,
 environmental, and individual factors. Yet, a comprehensive understanding of the drivers that
 influence such decisions is still lacking, particularly for the transient stage during which free living individuals are inherently difficult to follow.

26 2. Social circumstances such as the likelihood of encountering conspecifics can be 27 expected to strongly affects decision making during dispersal, particularly in territorial species 28 where encounters with resident conspecifics are antagonistic. Here we analyzed the movement 29 trajectories of 47 dispersing coalitions of Kalahari meerkats (*Suricata suricatta*) through a 30 landscape occupied by constantly monitored resident groups, while simultaneously taking into 31 account environmental and individual characteristics.

32 3. We used GPS locations collected on resident groups to create a geo-referenced social 33 landscape representing the likelihood of encountering resident groups. We used a step-34 selection function to infer the effect of social, environmental and individual covariates on 35 habitat selection during dispersal. Lastly, we created a temporal mismatch between the social 36 landscape and the dispersal event of interest to identify the temporal scale at which dispersers 37 perceive the social landscape.

4. Including information about the social landscape considerably improved our
representation of the dispersal trajectory, compared to analyses that only accounted for
environmental variables. The latter were only marginally selected or avoided by dispersers.
Before leaving their natal territory, dispersers selected areas frequently used by their natal
group. In contrast, after leaving their natal territory, they selectively used areas where they

43	were less likely to encounter unrelated groups. This pattern was particularly marked in larger
44	dispersing coalitions and when unrelated males were part of the dispersing coalition.
45	5. Our results suggest that, in socially and spatially structured species, dispersers gather
46	and process social information during dispersal, and that reducing risk of aggression from
47	unrelated resident groups outweighs benefits derived from conspecific attraction. Finally, our
48	work underlines the intimate link between the social structure of a population and dispersal,
49	which affect each other reciprocally.
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51	Keywords: Conspecific avoidance, informed dispersal, movement, social landscape, step
52	selection, Suricata suricatta
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### 68 INTRODUCTION

69 Dispersal of individuals is an important process governing the dynamics of spatially and 70 socially structured populations (Hanski 1999; Clobert et al. 2001). Through emigration and 71 immigration, dispersal can also affect local allele frequencies, alter the structure of existing 72 social groups, and lead to the formation of new groups (Clobert et al. 2001; Bateman et al. 73 2012). Improving our understanding of dispersal is therefore critical as we strive for better 74 forecasting of long-term population trends and viability (Bowler & Benton 2005; Ronce 2007). 75 The dispersal process can be divided into three phases: emigration from the natal 76 territory; transience; and settlement in a new territory (Bowler & Benton 2005). Each phase is 77 dependent on different individual, environmental, and social drivers, and our ability to 78 understand the mechanisms of dispersal relies on our ability to account for and understand the 79 factors affecting the movement of individuals in each of the three phases (Wiens 2001; Schick 80 et al. 2008; Clobert et al. 2009). Due to the difficulty of following individuals under natural 81 conditions, past studies have mainly focused on the causes and consequences of emigration 82 from the natal territory (Stephens et al. 2005; Cote & Clobert 2007, 2010). For instance, rates 83 of emigration have been shown to be density-dependent, to relax local competition, and to 84 favour inbreeding avoidance (Holekamp & Sherman 1989; Perrin & Mazalov 2000; Bowler & 85 Benton 2005; Bateman et al. 2012). In contrast, we have much less information about the 86 mechanisms and strategies that individuals use during the transient and settlement phases of 87 dispersal (Travis et al. 2012; Elliot et al. 2014). Empirical studies that focused on the transient 88 phase have concentrated on changes in movement characteristics and on the relationship 89 between movement trajectories and the surrounding environmental landscape. The combined 90 influence of the distribution and abundance of conspecifics - hereafter referred to as social 91 landscape – on decision making during transience has received relatively little empirical 92 attention, and related inferences are often based solely on theoretical work (Fletcher 2006;

Travis et al. 2012; Gilroy & Lockwood 2016). Yet the social landscape is likely to exert an 93 94 important influence on the behaviour of dispersing individuals during the transient phase. 95 In territorial social species such as the lion (Panthera leo), the African wild dog 96 (Lycaon pictus) and the meerkat (Suricata suricatta), encounters with unrelated groups and 97 individuals are often antagonistic and costly particularly for the smaller party (McComb, 98 Packer & Pusey 1994; Creel & Creel 2002; Mares et al. 2011). Dispersers are typically 99 outnumbered by resident groups, and thus, they may avoid areas where encounters with 100 resident groups are likely to occur. Such avoidance may be particularly strong in smaller 101 dispersing coalitions, and shortly before settlement, when dispersers seek an exclusive 102 territory. At the opposite end, the search for potential mates and high quality habitats during 103 transience and settlement may bring dispersers closer to resident groups (Andreassen & Ims 104 2001; Glorvigen *et al.* 2012). Whether dispersers are attracted to resident groups is likely to be 105 related to whether or not they can ever join them. Thus, dispersing male meerkats would be 106 expected to be attracted to resident groups, females to avoid them (Stephens et al. 2005; Mares 107 et al. 2011). In species that aggressively defend their territory from unrelated conspecifics and 108 where encounters can have fatal consequences, reducing costs of aggression may outweigh the 109 benefits derived from conspecific attraction (Stamps 2001; Fletcher 2006). Irrespective of 110 which strategy (conspecific avoidance or attraction) dispersers adopt during transience, their 111 ability to assess the social landscape at different spatial and temporal scales can reduce costs 112 and increase dispersal success (Bonte et al. 2012). Scent marks and other signs left by residents 113 convey information about their distribution and status (Jordan, Cherry & Manser 2007; Mares 114 et al. 2011; Jackson, McNutt & Apps 2012) and may be used by dispersing individuals to 115 indirectly gain information of the social landscape through which they move, avoiding risky 116 direct encounters.

117 A long-term study of the Kalahari meerkat (Suricata suricatta) (Clutton-Brock & 118 Manser 2016) provides a unique opportunity to investigate the role of the social and 119 environmental landscape during the transient phase of dispersal in a social and territorial 120 species. Meerkats are territorial cooperative breeders living in groups of 5-40 individuals, and each group occupies an exclusive territory of 1-5 km<sup>2</sup> (Clutton-Brock et al. 1999; Jordan et al. 121 122 2007). Intruding meerkats, being dispersers or individuals from neighbouring groups, are 123 readily challenged and chased by territorial groups, and such clashes can have severe 124 consequences (Mares, Young & Clutton-Brock 2012). Groups are characterized by the 125 presence of a dominant pair that monopolizes most of reproduction; subordinate individuals 126 delay dispersal and help rearing their siblings. Older subordinate females are aggressively 127 evicted from their natal group, and they form same-sex dispersing coalitions (Clutton-Brock et 128 al. 1998). Encounters with the natal group after eviction typically result in aggressive 129 behaviour directed towards the evicted individuals (Young et al. 2006). Dispersing coalitions 130 either leave the natal territory and search for unrelated males and a place to settle, or they are 131 sometimes let to re-join their natal group after the dominant female has given birth (Clutton-132 Brock et al. 1998). Subordinate females are, however, typically evicted in successive breeding 133 attempts by the dominant female and eventually leave the area to form a new group (Clutton-134 Brock et al. 1998). Evictions happen year-round, but typically peaks between September and 135 March (hereafter referred to as the dispersal season). Dispersing coalitions may have to travel a 136 considerable distance to locate vacant territories and during this period experience costs of 137 increased predation pressure and aggression from other meerkat groups (Clutton-Brock et al. 138 1999). In contrast to females, subordinate male meerkats spontaneously leave their natal group 139 and prospect for extra-group mating opportunities. Such prospecting forays typically last few 140 days, and a small proportion of them culminates in the males' dispersal or disappearance

141 (Young, Spong & Clutton-Brock 2007), likely after encountering a coalition of unrelated142 dispersing females.

143 In this study, we aimed to investigate the effect of the social landscape on movement 144 behaviour and decision making during the transient phase of dispersal in a socially and 145 spatially structured species. To do so, we fitted GPS radio collars on dispersing coalitions of 146 Kalahari meerkats and followed their movements across territories occupied by constantly 147 monitored resident groups. We created environmental maps from satellite imagery to control 148 for the effect of habitat types, taken to represent different habitat quality, on movements. We 149 used step-selection models to quantify habitat and social selection of dispersers before and 150 after they left their natal territory. We regularly visited dispersers and collected information on 151 coalition composition to evaluate the effect of coalition size on selection of locations in the 152 social landscape during transience. Finally, we identified the temporal scale at which 153 dispersing coalitions perceived and reacted to the social landscape and investigated whether 154 they integrated information about the distribution of resident groups collected over different 155 weeks and months.

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## 157 **METHODS**

## 158 Study system and data collection

The study site was located at the Kuruman River Reserve (S 26.92289°, E 21.84048) and the adjacent ranch-land near Van Zylsrus, South Africa. The site includes a section of the fossil Kuruman riverbed; elsewhere, the landscape is typical of the Kalahari Desert and is characterized by sparsely vegetated sand dunes and dry pans (Fig. 1A). The climate is characterized by a dry season between April and November and a wet season between December and March, with 250 mm annual average precipitation. For further details on the study site see Mares *et al.* (2012) and Young (2003). 166 The local population of wild meerkats has been subject of a long-term study started in 1993, and all study animals were individually marked. In any given year, between 15 and 18 167 resident groups that inhabit an area of 50-60 km<sup>2</sup> were regularly monitored. Resident meerkat 168 169 groups were habituated to the close presence (< 1 m) of researchers who visited the groups 170 every two to four days to collect behavioural and life-history data. During group visits, which 171 took place early mornings and late afternoons, and lasted three to four hours, researchers 172 consistently collected GPS locations every 15 minutes using a handheld GPS unit. We used 173 this spatial information on resident groups to create social landscapes (further detailed below). 174 We mounted GPS radio collars on subordinate dispersing females few days prior to or 175 immediately after eviction. We used behavioural cues such as aggression from the dominant 176 female, restless behaviour, and social withdrawal to identify potential disperser and anticipate 177 the time of eviction. We captured and anesthetized meerkats following a standardized protocol 178 used at the Kalahari Meerkat Project (Jordan et al. 2007). All captures were approved by the 179 South African Department of Environment and Nature Conservation and were performed under 180 permit 'FAUNA 192/2014'.

181 The GPS radio collars were composed of a 3.2 g stand-alone VHF beacon module 182 (Holohil Systems Ltd., Canada) and a 16.0 g stand-alone GPS module with integrated ZigBee 183 bidirectional communication capability powered by a 3.6 V <sup>1/2</sup> AA lithium battery (CDD Ltd., 184 Greece). We casted the two modules in epoxy resins and used a 4.5 mm wide cable tie coated 185 with soft heat-shrink tubing to attach the unit to the neck of the animals. Overall collar weight 186 was < 25 g, corresponding to 3-4 % of the animal body weight. Collars of this size and weight 187 do not affect meerkats (Golabek, Jordan & Clutton-Brock 2008). Only one individual was 188 equipped with a GPS radio collar in each dispersing coalition at any given time. We 189 programmed the collars to collect hourly GPS locations between sunrise and sunset as 190 meerkats sleep underground at night. We additionally fitted few individuals with VHF-only

191 radio collars (Biotrack, United Kingdom) weighing approximately 23 g, when deploying GPS

192 collars was not possible for logistic reasons. For coalitions fitted with VHF-only collars, we

193 manually collected GPS information at each visit (Supplementary Material S1).

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## **Dispersing coalitions**

We defined a dispersing coalition as a single female or multiple females, evicted from the natal group. We tracked each dispersing coalition by means of VHF telemetry every two to seven days to record coalition size, which included females and, when present, unrelated males. We defined a dispersal event as the collection of events starting at eviction and lasting until either return to the natal group, settlement in a new territory, or death before settlement. We did not include data collected after two weeks past successful settlement in a new territory as coalitions may show resident-like behaviour after settlement.

203 Within each dispersal season (September – March), we merged consecutive dispersal 204 events by the same dispersing coalition in a single combined dispersal event (Supplementary 205 Material S1). For example, if two sisters were evicted from the resident group, then accepted 206 back into the group, and evicted again shortly after, these two events were considered as one single dispersal event. This procedure reduced data dependence (two dispersal events of the 207 208 same coalition separated by a short period of time cannot be considered as independent) and 209 increased the data points of each combined dispersal event, which is a prerequisite for fitting 210 robust individual-level models for the step selection function (Craiu, Duchesne & Fortin 2008; 211 Fieberg et al. 2010; see below). During a three-year period, we monitored a total of 47 212 dispersing events, of which five were combined dispersal events consisting of two or three 213 consecutive events (Supplementary Material S1). Unless otherwise specified, all analyses were 214 performed using data from 47 dispersing coalitions.

We collected an average of 137 (range: 10 - 1114) GPS locations per dispersal event, which lasted on average 53 days (Supplementary Material S1). Twenty-seven coalitions rejoined their natal group after a variable duration of one (i.e. a coalition was evicted one day and allowed back to the group the following day) and 76 days. The remaining 20 coalitions did not return to their natal group and either dispersed for a minimum of six and a maximum of 284 days before settling in a new territory (n = 12 coalitions), dying (n = 3) or being lost (n = 5) before settlement.

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## 223 Creation of geo-referenced environmental layers

224 Within R 3.2.0 (The R Foundation for Statistical Computing), we used the rgdal package to 225 derive a geo-referenced digital elevation map (DEM) from Advanced Spaceborne Thermal 226 Emission and Reflection Radiometer (ASTER) images (Fig. 1B). We further used NASA's 227 Landsat 7 satellite images at 30x30 meters resolution to create a geo-referenced habitat map 228 following an unsupervised classification approach using the function *kmeans* (Fig. 1A). Given 229 the expected difference in reflectance between the vegetated sandy dunes and the dry 'pans', 230 this unsupervised approach was adequate. Following ground-truthed data at 80 randomly 231 selected locations, we merged the five classes (cfr. Fig. 1A) that resulted from the unsupervised 232 classification into three main habitat classes; namely "pans", "sandy matrix", and "transition" between the former two classes. This procedure allowed us to achieve an accuracy of 98% for 233 234 pans (1 in 45 pan locations was misclassified as transition), of 73% for sandy matrix (four in 235 15 matrix locations were misclassified as transition), and of 47% for transition. Pans were 236 never misclassified as matrix and vice versa (for further details see Supplementary Material 237 S2). The map obtained following this approach was very similar to vegetation maps for the 238 study site used by (Bateman et al. 2015) and Google Earth imagery (Google Inc. 2012) thus 239 confirming the reliability of the method (Fig. 1). Both ASTER and Landsat 7 images used are

available through the United States Geological Survey agency (http://www.usgs.gov).

Additionally, we used Google Earth to manually digitize and export the main channel of theKuruman River (Fig. 1).

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## 244 Creation of geo-referenced social layers

245 We used GPS location data collected on resident groups to create a geo-referenced social 246 landscape (Fig. 1C). Because GPS locations on resident groups were collected every 15 247 minutes throughout a visit and were therefore highly spatially and temporally correlated, we 248 computed the harmonic mean of all locations collected during one visit resulting in one GPS 249 location per visit. For each resident group, we created a kernel utilization density (KUD) map 250 with the function kernelUD in the adehabitatHR package (Calenge 2006). We reduced the 251 reference smoothing parameter  $h_{ref}$  until the smallest possible contiguous territory (i.e., one 252 single polygon) was obtained (Kie et al. 2010; Fieberg & Börger 2012). This procedure was 253 required because  $h_{ref}$  typically over-smooths the data, inflating territory size. We used GPS 254 data collected during a time interval of 30 days prior to eviction of the dispersing coalition of 255 interest to create the KUD maps for each resident group. We excluded resident groups with 256 less than 10 locations collected over the 30-day interval to reduce the likelihood of unreliable 257 KUD estimation. We multiplied KUD pixel values by group size and summed KUD raster 258 maps of each resident group to create a contiguous social landscape comprising the distribution 259 and abundance of all resident groups for the given 30-day interval. Low pixel values 260 represented non-risky area characterized by a small likelihood of encountering resident groups, 261 while high pixel values represented risky areas. We assigned NA values to pixels with no 262 information on resident groups. For those dispersing events lasting more than two weeks, we 263 recalculated the social landscape with a two-week sliding window. This procedure allowed us 264 to create a dynamic social landscape. A social landscape was created using data on resident

groups collected within 30 days prior eviction. The dispersing coalition of interest was then
'allowed to' move through this social landscape for two weeks (days 0 to 15 of the dispersal
trajectory). If the dispersal event lasted more than two weeks, a second social landscape was
created using data collected on resident individuals within 30 days prior day 15 of the dispersal
event. The same dispersing coalition was then 'allowed to' move through this new social
landscape for two weeks (days 16 to 30 of the dispersal trajectory); and so forth (see
Supplementary Material S3 for a dynamic representation of the social landscape).

272 To investigate at what temporal scales dispersing individuals perceive the social 273 landscape, we repeated the same process for alternative time intervals of 45, 60, 75, 90, 105, 274 120, 150, 180, 240 days. In a final analysis, which we used as control treatment, we created a 275 temporal mismatch between the social landscape and the dispersal events. For this, we created 276 the social landscape by shifting back the beginning of each time interval by 180 days compared 277 to the time of eviction of the dispersing coalition of interest. Because we assumed such a social 278 landscape to be of little relevance for dispersers, we expected no consistency in their selection 279 and avoidance of presumed risky areas.

280

## 281 Statistical Analysis

282 We used a step-selection function (SSF) to infer the effect of social and environmental 283 covariates on habitat selection during dispersal (Fortin et al. 2005). Within this framework, the 284 decision of a coalition to use a given location is conditioned on environmental and social 285 characteristics at alternative locations (Fortin et al. 2009) that the coalition could have reached 286 within an hour (the sampling interval between consecutive relocations). For each observed (i.e. 287 used) step along a dispersing coalition path, we created ten alternative steps; the end of these 288 steps represented alternative locations that the coalition could have chosen. A step is here 289 defined as the vector between two consecutive relocations spaced one hour apart. We created

290 alternative steps by randomly drawing angles from a continuous uniform distribution  $U(0, 2\pi)$ 291 and steps from a normal distribution  $N(\mu,\sigma)$  with  $\mu$  equal to two times the mean empirical step 292 length of monitored dispersing coalitions, with  $\sigma$  equal to one step length standard deviation. 293 We chose  $\mu$  to allow alternative relocations to be far from the relocation used by the focal 294 coalition, and thus to be characterized by different environmental and social factors. Dispersing 295 meerkats are bound to sleeping burrows, with new burrows being used as the dispersal process 296 progresses. While steps between consecutive burrows can be expected to be directional (i.e. 297 small turn angles), steps in the vicinity of sleeping holes are more circuitous (i.e. equally 298 distributed turn angles). As we could not distinguish between these two types, drawing angles 299 from a uniform distribution appeared more parsimonious.

Following this matched case-control design (Fortin *et al.* 2005), we built SSF using
conditional logistic regression to compare used (scored 1) and alternative (scored 0) locations.
The SSF took the general form

 $w(\mathbf{x}) = \exp \left(\beta_1 x_1 + \beta_2 x_2 + \dots + \beta_{n-2} x_{n-2} + \beta_{n-1} x_1 : x_2 + \beta_n x_1 : x_3\right)$ 

304 where  $\beta$ 's are selection coefficients associated with covariates and biologically meaningful 305 interaction terms (Fortin & Fortin 2009). Steps with higher SSF scores w(X) are more likely to 306 be chosen by the animals (Fortin *et al.* 2005), and  $\beta = 0$  indicates absence of 307 selection/avoidance (Forester, Im & Rathouz 2009). The environmental covariates considered 308 in the analyses included ELEVATION (continuous), DISTANCE TO RIVER (continuous), 309 and the binary habitat class PAN (coded 1 if a dispersing coalition was on a pan and 0 if 310 outside). We did not include the other two habitat classes (sandy matrix and transition) to avoid 311 collinearity with pans. The social variables were RISKINESS (the continuous social landscape 312 values), and NATAL (coded 1 if a dispersing coalition was within and 0 if outside the territory 313 boundary of the natal group). Because we anticipated that dispersing coalitions behave 314 differently within and outside the territory of the natal group, we included a RISKINESS by

315 NATAL interaction in our model. We also added dispersing COALITION SIZE as part of an 316 interaction term with RISKINESS to test whether coalitions of different size reacted differently 317 to the social landscape. Because each used and its ten alternative locations were assigned the 318 same coalition size, coalition size cannot appear in the model as a main effect (Fortin & Fortin 319 2009). We further defined a CLUSTER as the unique identifier for each independent dispersal 320 event, and a STRATUM as the unique identifier for each combination of used and alternative 321 locations. We used CLUSTER and STRATUM as grouping variables in the analyses (Fortin et 322 al. 2005). We implemented the above model in the *clogit* function in the *survival* package to 323 obtain population-level estimates of the selection coefficients associated with each covariate. 324 We calculated alternative SSF models with RISKINESS values obtained at each time intervals 325 (30, 45, ..., 240 days) to detect the temporal scale at which dispersing coalitions showed the strongest response to the social landscape (cfr. Fig. 3). We reported  $\beta$  coefficients associated 326 327 with all covariates only for the time interval that showed the strongest response. 328 In a second step, we used  $\beta$  coefficients obtained from conditional logistic regression 329 fitted to each coalition independently (Fieberg et al. 2010) to make inference on the effect of 330 additional coalition-specific covariates on step selection. We used linear regression to test the 331 effect of coalition size on estimated  $\beta$  coefficients, which were treated as response variables

332 (Thurfjell, Ciuti & Boyce 2014). Because some coalitions either never left the territory of their

atal group or almost only roamed outside of it (Supplementary Material S1), it was not

possible to fit a single model including the variable NATAL to each single dispersing

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events (e.g. all observed and alternative locations in a stratum have the same NATAL value).

coalition. This was due to matrix singularities resulting from a coalition experiencing the same

337 We therefore ran two separate models; one with data collected when the dispersing coalitions

338 were within the territory of their natal group, and one when they were outside the natal

territory. To obtain robust coalition-specific estimations, we only retained dispersing coalitions

340 with a minimum of 30 used locations (see Craiu et al. 2008). Of the 47 monitored dispersing 341 coalitions, 30 coalitions met this criterion while roaming within the territory of their natal 342 group and 15 while outside (Supplementary Material S1). Like above, we run these two models 343 only for the time window at which RISKINESS showed the strongest response. 344 In an additional analysis, we used a generalized additive mix model in the mgcv 345 package (Wood 2004) to find the shape of any nonlinear relationship of RISKINESS over time 346 (days since eviction) without imposing any specific parametric form. We averaged 347 RISKINESS values on a daily basis to reduce autocorrelation issues. We allowed for 348 differences between coalitions that returned to the natal group and coalitions that settled, while 349 controlling for the effect of coalition size and the presence of unrelated males within a 350 coalition. We included coalitions identity as random term.

351

### 352 **RESULTS**

353 Movement patterns during dispersal were largely dependent on coalition size. Of the 47 354 dispersing coalitions monitored, the coalitions that settled (n = 12) were typically larger (mean 355 = 2.3 females or 4.7 females and males) than the coalitions that returned (n = 33 coalitions; 356 mean = 1.8 females or 1.9 females and males; Wilcoxon W = 116, p = 0.14 and W = 39.5, p < 0.14357 0.01, for females or females and males respectively) (Supplementary Material S4). Average 358 maximum displacement from the site of capture was 2'263 m (range 434 - 10'742 m). 359 Coalition size showed a positive relationship with the log-transformed maximum distance 360 travelled ( $F_{1.46} = 6.14$ , p = 0.017) after correcting for differences in dispersal time.

After eviction from their natal group, dispersing coalitions selected for low-lying slacks between sand dunes and avoided pans (Table 1). Selection coefficients for PANS in four coalitions appeared implausibly large for an exponential model (e.g. -17.68; Table 2) and were likely due to matrix singularities. We therefore ran two alternative models: in one model, we 365 excluded these four coalitions, and in the other model we retained all coalitions but removed 366 PANS from the model. Results of coefficients of selection (i.e., ß values) remained unchanged. 367 Dispersing coalitions neither selected nor avoided locations closer to (respectively, further 368 away from) the fossil Kuruman riverbed. Dispersing coalitions reacted differently to the social 369 landscape depending on whether they were inside or outside the territory of the natal group 370 (RISKINESS:NATAL interaction term:  $\beta = -0.055$ , robust SE, 0.014, p < 0.01; Table 1). When 371 they were still within the territory of their natal group, they selected for locations characterized 372 by a higher likelihood to encounter the natal group. When they roamed outside the territory of 373 their natal group, they avoided locations characterized by a high likelihood of encountering 374 unrelated resident groups. These results thus support the hypothesis that dispersing coalitions 375 move in a way to minimize interactions with unrelated territorial groups.

376 Avoidance of risky locations outside the territory of the natal group persisted through 377 all time windows used to determine the social landscape, but the strength of risk avoidance 378 peaked at time windows between 60 and 75 days (Fig. 2). When we introduced a 180-day 379 mismatch between the social landscape and the time of eviction (see methods), we observed no 380 differences in the degree of selection of risky locations for areas inside and outside the territory 381 of the natal group (Fig. 2). This suggested that the social landscape is of relevance only within 382 a time intervals of a few months but is not informative over longer periods. Overall, adding 383 information about the social landscape to the baseline model that only accounted for 384 environmental variables improved model performance (log-likelihood ratio test for nested models:  $\chi^2 = 261$ , p < 0.01, Supplementary Material S5). 385

Analysis at the coalition level did not support the hypothesis that smaller dispersing coalitions avoided areas where encounters were likely to occur more than larger coalitions. Although not statistically significant, there was an indication that selection for high-likelihoodof-encounter areas decreased as dispersing coalition size increased (Fig. 3); both within (slope 390 = -0.013, SE = 0.009, p = 0.14) and outside (slope = -0.12, SE = 0.11, p = 0.31) the territory of the natal group. Overall, results obtained at the coalition level (Table 2) largely supported the results obtained at the population level (Table 1) showing that when dispersing coalitions were within the territory of their natal group, they selected for areas characterized by a higher likelihood to encounter the natal group (Table 2, Fig 3).

As time after eviction progressed, coalitions that settled increasingly used areas where they were less likely to encounter conspecifics (F = 17.2, p < 0.01), whereas we could not detect any relationship with time for coalitions that eventually re-joined their natal group. The presence of unrelated males was associated with the use of areas characterized by a lower likelihood to encounter resident conspecifics (t = -3.4, p < 0.01), both for coalitions that rejoined their group and for coalitions that settled elsewhere, while coalition size had no statistically significant effect (t = -1.1, p = 0.3).

402

## 403 **DISCUSSION**

Our study showed that, in a population of wild Kalahari meerkats, the movements of dispersing 404 405 individuals were influenced by the social landscape, i.e. the distribution and abundance of 406 resident conspecifics. Importantly, the reaction to this social landscape, depended on whether 407 dispersers were inside or outside the territory of their natal group. Before leaving their groups' 408 territory, dispersers appeared to maximise their chance of being in the same area as their natal 409 group. After leaving their natal territory, they selectively used areas where they were less likely 410 to encounter other unrelated groups, and increasingly so when unrelated males were part of the 411 dispersing coalition. This contrasting use of the social landscape suggests that dispersing 412 meerkats can acquire information on the distribution and abundance of conspecifics and use 413 this information to minimize antagonistic contacts with unrelated territorial groups.

414 Seeking proximity of the natal group despite the possibility of aggression, is consistent 415 with existing evidence of the advantages of group living and delayed dispersal in wild Kalahari 416 meerkats (Clutton-Brock et al. 1999; Clutton-Brock & Manser 2016). The tendency for 417 dispersing individuals to 'shadow' their natal group after being evicted was more pronounced 418 for smaller than for larger dispersing coalitions, suggesting that small coalitions may suffer 419 higher costs or gain smaller benefits by leaving their natal area. Associated costs may be direct 420 costs such as increased mortality due to reduced antipredator vigilance and dilution effect in 421 smaller groups (Delm 1990), or fitness costs associated with reduced recruitment rate in small, 422 newly formed groups (N. Maag pers. comm.).

423 Outside the territory of the natal group, moving through areas characterized by a small 424 likelihood of encountering unrelated resident conspecifics is likely to be beneficial since it 425 reduces antagonistic interactions and increase dispersal success. In the effort to avoid resident 426 groups, however, dispersers may move through suboptimal and resource-poor areas (Palomares 427 et al. 2000). While resident meerkat groups have been shown to concentrate along the edges of 428 the dry riverbed and surrounding flats habitats (Bateman et al. 2015), the dispersing coalitions 429 monitored during this study avoided such habitats (here referred to as pans). The riverbed and 430 pans offer more shelters (bolting holes) than the surroundings (Manser & Bell 2004); 431 avoidance of these safer habitats suggests that avoidance of resident groups may be a stronger 432 driver of habitat selection during dispersal than predation pressure. Such strategy may be 433 favoured by a relative low predation pressure; we only recorded one case of predation by a bird 434 of prey (on a single dispersers) and two meerkats were hit by cars. As food availability can 435 mask the effect of predation and conspecific aggression on habitat selection we urge further 436 investigation. In the Kalahari rainfall is highly variable both between and within years and it 437 affects food availability with consequences on meerkats growth and body condition (English, 438 Bateman & Clutton-Brock 2012). Changes in food availability may be partly responsible for

some of the variations in habitat selection observed among individual dispersing coalitions. This in light of the fact that the confidence intervals for the  $\beta$  coefficients of the habitat features considered were only lightly different from zero (which would represent no selection) and hence more susceptible to changes in selection or avoidance following alteration of the characteristics of the habitat itself.

444 Our results did not support the hypothesis that larger dispersing coalitions are 445 competitively stronger than smaller coalitions and therefore less likely to avoid areas 446 characterized by a high likelihood of encountering unrelated resident groups. This may be due 447 to the fact that no matter how large a dispersing coalition may be, it will inevitably be 448 substantially smaller than a resident group, which averages 15.5 individuals (Bateman et al. 449 2013), and therefore consistently avoid it. Our results suggest that rather than influencing 450 movement patterns during transience, coalition size influenced the final outcome of the 451 dispersal event – with smaller coalitions returning to the natal group and larger coalitions 452 settling in new territories. Movements during transience were instead influenced by the 453 presence of unrelated males and increasing time after eviction. Both factors caused dispersing 454 coalitions to avoid areas with a high likelihood to encounter resident groups, suggesting that 455 newly formed groups seek vacant or little used territories to settle. Mates thus not only 456 influence dispersal patterns at emigration and settlement (Cote & Clobert 2007; Davidian et al. 457 2016; Gilroy & Lockwood 2016), but also during the transient phase. In fact, evidence 458 suggests that, in meerkats, the association with unrelated males is one of the key drivers 459 promoting the transition between the consecutive phases of dispersal (N. Maag et al. in 460 review).

Results from our study also provide important insight into the temporal scale at which dispersing coalitions perceive and react to the social landscape. The underlying assumption of our analysis is that the GPS locations used to model the distribution of resident groups directly 464 correlate with cues of their presence and that dispersing coalitions perceive and react to such 465 cues. These may be direct encounters or indirect cues such as scent markings. Our analysis 466 showed strongest avoidance of areas characterized by a high likelihood to encounter unrelated 467 resident groups when the social landscape was modelled using GPS data collected on resident 468 groups during the 60/75-day interval preceding the dispersal trajectory. Using GPS locations 469 collected during shorter or longer time periods resulted in decreasing avoidance. We speculate 470 that a social landscape based on information collected during a too short period (< 30 days) 471 does not accurately represent the risks perceived by dispersing coalitions. The same applies to 472 social landscape based on information collected long before the dispersal event took place, as 473 suggested by our mismatch design (180-days timescale). These results suggest that care must 474 be taken in the temporal offset for the information used to create the social landscape.

475 The relative roles of direct and indirect cues in shaping dispersers' own representation 476 of the social landscape remains to be determined (Creel et al. 2005; Broekhuis et al. 2013). The 477 ability to predict risks related to distribution and abundance of conspecifics based on indirect, 478 long-lasting olfactory cues seems a parsimonious mechanistic explanation for a species that 479 relies on scent marking for between- and within-group signalling and communication (Jordan 480 et al. 2007; Mares et al. 2011; Bateman et al. 2015). While we have no direct evidence on how 481 long such cues may last for in meerkats under specific environmental conditions, reaction to 482 olfactory cues of several weeks to months has been documented for mammalian scent marks 483 (20-25 days: Helogale parvula; 65 days: sniffer dogs; 40-100 days: Mesocricetus auratus; 180 484 days: Micricebus coquerely) (Apps, Weldon & Kramer 2015). Our results suggest similar time 485 windows to be plausible for meerkats, but further investigation is required.

486 Our work, which focused on understanding the effects of the social landscape on
487 movement behaviour and decision making during the transient phase of dispersal, provides
488 empirical evidence for one of the "four areas where research effort should be directed", as

489 identified by Clobert et al. (2009) in their seminal review paper. Our results support the 490 concept of socially informed dispersal (Clobert et al. 2009), where individuals gather social 491 information during their movements. Empirical evidence if this applies also during the 492 transience phase is critically lacking. The ability of dispersers to assess the topology of the 493 social landscape influences how and where they move, with possible consequences on the 494 settlement process. The latter can in turn influence the dynamic and persistence of structured 495 populations. While we are only scratching the surface in understanding the proximate 496 mechanisms and demographic consequences of dispersal, our results are an important step 497 towards a better understanding of socially informed dispersal in a spatially structured species. 498

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- 513 DATA ACCESSIBILITY STATEMENT Data supporting the results of this manuscript will
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- 515

AUTHORS' CONTRIBUTION GC, AO and NM conceived the study, NM collected data on
dispersing groups, TCB led the collection of data on resident groups, GC and LB performed
analysis, GC wrote the first version of the manuscript, and all authors contributed substantially
to revisions.

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683 Figure 1: Environmental and social geographic layers for the study area along the fossil 684 Kuruman River, South Africa. The site was divided in five discrete habitat types 685 subsequently merged into three: pans, transition and sandy matrix (A). For comparison, Google 686 Earth images have been inserted (framed inserts). A digital elevation model for the study area 687 ranging between 890 and 960 meters above sea level was derived from ASTER imageries (B). 688 A kernel utilization density (KUD) map for each resident group was calculated reducing the 689 reference smoothing parameter h<sub>ref</sub> until the smallest possible contiguous territory (grey 690 polygon) was obtained. Dots represent collected GPS locations (C1). KUD pixel values were then multiplied by group size. Light (yellow) colors represent frequently used regions (C2). 691 692 KUD raster maps of each resident group were summed together to create a social landscape comprising all resident groups (C3). Example trajectories of a dispersing coalition that settled 693 694 (dark dotted line) and a dispersing coalition that returned to the natal group (grey dashed line) 695 are shown (C3). In all panels, the main river channel (solid blue line) are depicted for spatial 696 reference.



699 Figure 2: Dispersing coalitions step selection coefficients associated with the distribution 700 of resident groups. The beta coefficients show the departure from baseline social landscape 701 values that dispersing coalitions select when moving within the territory of the natal group. 702 This means that when dispersing coalitions are outside the territory of the natal group they 703 select for areas characterized by a lower likelihood of encountering resident conspecifics. 704 Discrete social landscapes (real scenario) were calculated using GPS relocation data collected on resident groups during alternative time intervals (30, 45, ..., 240 days) prior to emigration 705 706 by the dispersing coalition of interest. For comparison, a temporal mismatch between the social 707 landscape and the dispersal events was created by shifting the time intervals used to create the 708 social landscape back in time by 180 days (mismatch scenario).

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Table 1: Population-level selection coefficients. Coefficients and robust standard errors (S.E.) from a step selection function analysis where each observed location was conditioned on 10 alternative locations. Pans: factorial (whether a location is inside a pan or in the surroundings). Natal: factorial (whether a coalition is inside or outside the territory of the natal group). Riskiness: continuous values associated with the social landscape (range:  $x \rightarrow 0$  to max(x)). Elevation and distance to river are measured in meters. Coal size: Maximum coalition size. Removal of distance to river did not change the results.

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		coefficient	S.E.	p-value
740	ELEVATION	-0.020	0.008	0.011
741	PANS (inside)	-0.259	0.131	0.049
741	DISTANCE TO RIVER	0.000	0.000	0.440
742	NATAL (outside)	-0.314	0.163	0.163
	RISKINESS	0.059	0.009	< 0.01
743	NATAL(outside):RISKINESS	-0.055	0.014	< 0.01
711	COAL SIZE:RISKINESS	-0.005	0.003	0.090
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**Table 2: Individual-level selection coefficients.** Coefficients and standard errors of
movement steps during dispersal regressed against environmental and social variables. For a
description of variables see Table 2.

### Inside NATAL

COALITION ID	COAL SIZE	ELEVATION	PANS (inside)	DISTANCE TO RIVER	RISKINESS	COALITION ID	COAL SIZE	ELEVATION	PANS (inside)	DISTANCE TO RIVER	RISKINESS
	β						β				
1	1	-0.085	-16.354	-0.004	0.402	1	1	-0.093	-17.523	0.000	0.065
2	2	-0.037	-15.668	0.002	0.056	4	6	-0.025	0.390	0.000	-0.315
3	3	0.025	-16.995	-0.003	0.066	5	6	-0.003	-1.579	0.000	-0.027
4	6	-0.012	0.208	-0.004	0.043	8	2	-0.001	0.403	0.000	0.004
5	6	0.040	-1.583	0.000	0.003	31	1	-0.044	0.179	0.000	0.029
6	4	0.065	-0.112	0.000	0.100	10	6	-0.061	-0.163	0.000	0.019
7	3	0.103	-1.643	0.001	0.384	32	4	-0.013	-0.867	0.000	0.021
8	2	-0.027	0.925	0.000	0.042	33	3	-0.065	0.323	0.000	-0.021
9	1	0.005	-0.962	0.000	0.076	34	4	-0.018	0.093	0.000	-0.326
10	6	-0.010	-0.235	0.000	0.061	16	3	0.029	-0.092	0.000	0.053
11	2	-0.083	-17.720	0.000	0.160	35	6	-0.102	-0.354	0.000	-3.831
12	4	-0.028	-0.707	0.000	0.098	17	3	0.061	-1.244	0.001	-0.056
13	2	-0.086	-0.855	0.002	0.012	19	9	0.030	1.082	-0.001	-0.012
14	2	-0.005	-0.037	0.001	0.060	22	3	-0.166	-0.059	0.002	-0.121
15	5	-0.032	0.027	0.000	0.106	30	1	-0.068	-0.490	0.000	-0.010
16	3	-0.018	0.210	0.001	0.077						
17	3	-0.037	-0.407	-0.001	0.070						
18	3	-0.078	-0.120	0.000	0.092						
19	9	0.007	0.105	0.000	0.053						
20	2	-0.042	0.096	0.001	0.043						
21	1	-0.043	-1.288	0.002	0.194						
22	3	-0.037	-0.869	0.002	0.058						
23	2	-0.160	0.001	0.001	0.025						
24	2	0.139	-1.251	0.001	0.161						
25	1	0.044	0.268	-0.001	0.115						
26	1	-0.070	-0.283	0.000	0.060						
27	6	0.064	-0.326	0.000	-0.045						
28	2	-0.178	1.088	-0.001	0.004						
29	1	0.022	0.337	0.000	0.063						
30	1	0.012	0.101	-0.001	0.061						