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Social interactions predict genetic diversification: an experimental manipulation in shorebirds

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Data Accessibility

Analyses reported in this article can be reproduced using the data provided by Charles Cunningham et al. (2018).

Abstract

Mating strategy and social behaviour influence gene flow and hence affect levels of genetic differentiation and potentially speciation. Previous genetic analyses of closely related plovers Charadrius spp. found strikingly different population genetic structure in Madagascar: Kittlitz’s plovers are spatially homogenous whereas white-fronted plovers have well segregated and geographically distinct populations. Here we test the hypotheses that Kittlitz’s plovers are spatially interconnected and have extensive social interactions that facilitate gene flow, whereas white-fronted plovers are spatially discrete and have limited social interactions. By experimentally
removing mates from breeding pairs and observing the movements of mate-searching plovers in both species, we compare the spatial behaviour of Kittlitz’s and white-fronted plovers within a breeding season. The behaviour of experimental birds was largely consistent with expectations: Kittlitz’s plovers travelled further, sought new mates in larger areas, and interacted with more individuals than white-fronted plovers, however there was no difference in breeding dispersal. These results suggest that mating strategies, through spatial behaviour and social interactions, are predictors of gene flow and thus genetic differentiation and speciation. Our study highlights the importance of using social behaviour to understand gene flow, although future work is needed to investigate the relative importance of social structure, intra- and between-season dispersal in influencing the genetic structures of populations.

Keywords: speciation, social network, mating opportunities, genetic structure, mating systems, spatial behaviour, dispersal, shorebird, gene flow, Madagascar
Introduction

How new species emerge despite homogenising gene flow is one of the most debated topics in evolutionary biology (Price, 2008; Futuyma, 2013). Although speciation is possible with continuous gene flow between lineages, this typically impedes speciation (Slatkin, 1987; Niemiller et al., 2008; Hereford, 2009; Matute, 2010; Feder et al., 2012). Understanding factors that affect gene flow is important beyond evolutionary biology; if local environments change abruptly or species suffer population or range contractions due to climate change, population fitness and productivity may decline unless genetic diversity is preserved within the extended population (Frankham, 1996; Arenas et al., 2012; Aitken & Whitlock, 2013). Sexual selection, typically more intense in polygamous than monogamous species, is often considered to facilitate speciation through variety of mechanisms via sexual conflict or intrasexual competition (Wilkinson and Birge, 2010; Gavrilets, 2014; Arnaqvist & Rowe, 2002; Ritchie, 2007). Greater gene flow creates more uniform population genetic structure, but it also maintains greater genetic diversity within the population (Aitken & Whitlock, 2013; Eberhart-Phillips et al., 2015). However, recent work suggests that the variance in mating success associated with strong sexual selection may also constrain speciation through promoting individual spatial movement, resulting in increased gene flow in polygamous species (Küpper et al., 2012; D’Urban Jackson et al., 2017).

Dispersal events typically increase gene flow, including natal and breeding dispersal, migration, as well as fine-scale movements that increase demographic connectivity within populations (Ronce, 2007; Burns & Broders, 2014; McGuire, 2013; Pilot et al., 2010). Many species of birds and mammals disperse to enhance mating opportunities and reproductive success; and access to mates, resources, and the avoidance of inbreeding are important in promoting sex specific dispersal
(Greenwood, 1980; Lenormand, 2002; Trochet et al., 2016). However, fine-scale continuous events, such as the social environment, spatial distribution and mate search behaviour, are often overlooked (Wey et al., 2015; Skrade & Dinsmore, 2010) in favour of rarer, large-scale dispersal events which cannot explain observed levels of gene flow alone (D’Urban-Jackson et al., 2017, Morinha et al., 2017). Individual movement patterns and space use strategies can influence social interaction as well as mating success, and hence gene flow (Duvall, 1997; Sih et al., 2009; McGuire, 2013). As well as affecting gene flow, the spatial distribution of individuals may in turn influence encounter rates influencing sexual competition (Tuni & Berger-Tal, 2012; D’Urban Jackson et al., 2017). This alteration of sexual selection patterns will in turn influence mating strategies (Oh & Badyaev, 2010), which provides feedback into movement patterns (Fromhage et al., 2016). Additionally, studies of social behaviour in birds, insects, and mammals have predicted higher levels of social interaction result in more gene flow, less speciation and higher extinction rates (Cockburn, 2003; Wilkinson & Birge, 2010; McGuire, 2013); suggesting gene flow may be reduced through limited social interactions.

Recent genetic analyses of closely related shorebirds, the Kittlitz’s plover Charadrius pecuarius and the white-fronted plover Charadrius marginatus, showed that they exhibit different population genetic structure throughout their breeding range in Madagascar: Kittlitz’s plover had a panmictic and homogenous population with no population structure detected, whereas the white-fronted plovers exhibited well-defined geographically distinct populations (Eberhart-Phillips et al., 2015). The life-history and ecology of these two species are very similar, e.g. both are small insectivorous ground-nesting shorebirds with modal clutch size of two eggs and precocial young, and these species often breed side by side in Madagascar (Zefania & Székely, 2013). However, their mating
systems are different: Kittlitz’s plovers are polygamous whereas white-fronted plovers are socially (and genetically) monogamous (Zefania et al., 2010; Maher et al., 2017). Parra et al. (2014) found that re-mating times were different between male and female Kittlitz’s plovers, whereas in white-fronted plovers the re-mating times were similar for males and females, demonstrating interspecific variation in mating opportunities and mate fidelity. The genetic data on population structure across a large geographic area (Eberhart-Phillips et al., 2015) and the experimental manipulation of mating opportunities in the field (Parra et al., 2014) provide a unique opportunity to explore the spatial and social processes through which sexual selection may influence gene flow within breeding seasons by using data that have not been analysed previously.

Here we investigate movement and interaction of experimental plovers, using spatial and network methodologies to analyse experimental data, to test two key predictions. First, due to differences in mating opportunities, we predicted more movement by polygamous Kittlitz’s plovers in order to find new mates compared with monogamous white-fronted plovers (Székely & Lessells, 1993; Küpper et al., 2012; Parra et al., 2014), specifically greater distance travelled over larger home ranges as well as higher dispersal distance. Second, in accordance with the first prediction and known population structure (Eberhart-Phillips et al., 2015), Kittlitz’s plovers should demonstrate greater spatial and social interaction with conspecifics than white-fronted plovers. Plovers have often been used as a behavioural model system to understand mating system evolution (Székely et al., 2006; Vincze et al., 2016; Maher et al. 2017), and testing these predictions using spatial and social interaction data will provide the link between population genetic study and diversification, and mating system variation using the Malagasy plovers as a case study.
METHODS

Study Species and Study Sites

Kittlitz’s and white-fronted plovers were investigated in southwest Madagascar. Kittlitz’s plovers were studied between 6 February and 13 May 2010 in Andavadoaka (22° 02’ S, 43° 39’ E, Fig. 1) where approximately 300 Kittlitz’s plovers breed around alkaline lakes (J.E. Parra, S. Zefania, & T. Székely, unpublished data). Fieldwork with the white-fronted plover was carried out between 1 April and 23 June 2011 at Lake Tsimanampetsotsa National Park (24° 3’ S, 43°44’ E, Fig. 1), a large alkaline lake (15 km × 0.5 km), surrounded by sandy beaches, short grass, and saltpans. Approximately 150 white-fronted plovers breed around the lake (J.E. Parra, unpublished data).

In the field, nests were searched for on foot or from hides by spotting incubating parents returning to their nest. In total, 18 Kittlitz’s plover pairs (36 individuals) and 14 white-fronted plover pairs (28 individuals) were captured with funnel traps placed on their nests (Fig. 1). The differing sample sizes reflect the maximum number that was possible to catch with the resources available (J.E. Parra, S. Zefania, & T. Székely, unpublished data). Nest search, trapping and behavioural observations followed standard protocols that have been adopted in previous publications (Carmona-Isunza et al. 2015, Vincze et al. 2016; D’Urban-Jackson et al. 2017; Maher et al. 2017). The traps were continuously monitored until a parent entered the trap and sat on the eggs, and then it was removed immediately to reduce stress and the risk of injury. All adults were ringed with an individual colour ring combination and a numbered SAFRING metal ring from the University of Cape Town, South Africa. Study birds were differentiated from other ringed individuals by using green permanent marker (Pilot Supercolour) on the individual’s white belly.
The mate removal protocol of Székely et al. (1999) was followed to experimentally create unmated sexually-active individuals. This experimental treatment ensured that a mate-searching phase was included within the movement of all individuals, which would not have been possible with purely observational study. Briefly, both parents were trapped, ringed, measured and a blood sample was taken for sex determination since the adult plumage is sexually monomorphic in both species (see below and Supplementary Information). One parent was then selected at random (since the sex was not known until after the experiment) for release at the capture location immediately; and the other parent was taken into captivity (see below). In both Kittlitz's and white-fronted plovers, both the male and female incubate the eggs (Hockey et al., 2005; Urban et al., 1986). Eggs of experimental birds were translocated to nearby conspecific nests with eggs at a similar developmental stage. Only pairs incubating two eggs (modal clutch size in both species) were manipulated. Trapping locations for both species were distributed evenly over an area of similar size (Fig. 1).

Removed plovers were transported to a nearby purpose-built aviary as detailed in Parra et al. (2014). Captive plovers were measured and then released after their former mate either found a new mate, or was not seen in the study sites for at least 12 days. Time in captivity was comparable for white-fronted plovers (number of days in captivity: 8.0 ± 1.71 days, N = 14, we provide mean ± SD unless stated otherwise) and Kittlitz's plovers (7.12 ± 2.57 days, N = 18). Although captive plovers appeared to lose a small amount of body mass during their time in captivity (2.77 ± 0.51 g in Kittlitz's plover, and 0.73 ± 0.22 g in white-fronted plover), many remated shortly after release.
indicating salubrious condition. The experiment was approved by the Malagasy authorities – see additional information including the Ethical note below and in Supplementary Information.

**Behavioural Observations**

Both the immediately released and the captive plovers released from the aviary were searched for within the study area every day in the field, using a car and mobile hide, after release. When an experimental plover was found, the coordinates of its location were taken with a handheld GPS receiver (Garmin e-Trex H). In addition, we collected 30 minute behavioural samples of one of the two species, the white-fronted plover, by recording the behaviour of experimental plovers every 30 seconds (see details in Parra et al., 2014) immediately after a resighting. The identity of other experimental plovers the focal individual interacted with during this time was also recorded. Although attempted, it was not possible to complete 30 minute behavioural samples for every resighting due to logistical survey limitations, i.e. if the focal individual flew off. Behavioural categories included social interactions such as fighting, courting and copulation. Two observers (M.B. and J.E.P.) collected the behavioural records, and both sampling methodology and behavioural categories were standardized between the observers. Since adults are sexually monomorphic in both species (Urban et al., 1986; Hockey et al., 2005) we used molecular sex typing to determine the sex of individuals (dos Remedios et al., 2010). Molecular sexing was carried out in NERC-Biomolecular Analysis Facility at the University of Sheffield (for details see dos Remedios et al., 2010; Parra et al., 2014, Supplementary Information).

**Home Range and Movement Analyses**
The R package adehabitatHR (Calenge, 2006) was used to calculate the home ranges of individual plovers using the kernel method (Worton, 1989) using every observed sighting of each individually marked plover (termed ‘relocations’ henceforward). First, the utilization distributions (UD) of 24 white-fronted (total relocations 327; mean 13.63 ± 5.75, min 6, max 26) and 32 Kittlitz’s plovers (total relocations 512; mean 16.0 ± 6.53, min 6, max 28) were calculated; 2 white-fronted and 4 Kittlitz’s individuals were not included in the UD analyses because they had less than 6 relocations (Calenge, 2006). The kernel smoothing parameter, $h$, was optimised by the least-square cross validation (LSCV) method (Gitzen & Millspaugh, 2003). For several individuals the LSCV did not converge (Seaman & Powell, 1998), hence in order to produce a UD for every experimental individual, smoothing parameter limits were set beyond which the ad hoc method was used (Worton 1995; Calenge, 2011; Kie, 2013). These limits were set by eye to ensure there was not unrealistic fragmentation or over-smoothing of home ranges. The home range was then calculated from the UD as the area within which the probability of locating an individual is equal to a specified value (Worton, 1989, Calenge, 2011). To include the mate searching area as well as the core use area, a 90% home range was used in the analysis (Fig. 1, Fig. S1) as it provided the largest reliable home range size (Börger, 2006).

Second, plover movement was investigated using step lengths of individuals (Marsh and Jones, 1988; Turchin, 1998; Zeller et al., 2012), calculated with the R package adehabitatLT (Calenge, 2006). Step lengths, calculated as the distances between consecutive points (Fig. 1, Fig. S1), were summed and then divided by the number of relocations to infer the mean step length for each individual Kittlitz’s plover (grand mean step duration: 1.88 ± 0.77 days) and white-fronted plover (1.04 ± 0.68 days). Third, breeding dispersal was investigated as the net distance between
territories (Fig. 1, Fig. S1. This was calculated from the distance between the first nest location in
the original territory, i.e. the capture point; and the centroid point of the core-use area, i.e. the
secondary territory. The core-use area was calculated as the 50% home range using the same
technique used to find the 90% home range size (above).

Generalized linear models (GLMs), with Gaussian error structure and identity link function, were
used to test whether species, sex and their interaction predict the spatial behaviour of plovers (i.e. home range size, mean step length, and distance from previous territory). Log transformation was
used for each response variable to normalise the data. Two models were fitted for each predictor variable, one basic model with fixed factors of species, sex, and species * sex interaction; and another model with additional control variables including number of days tracked, number of relocations, and captivity (i.e. released immediately after capture in the field, or released from captivity). ‘Number of relocations’ was not included in the GLM analyses of mean step length as it was used in the calculation of the variable. The models were compared using an Analysis of
Deviance test, and in all cases the more complex model did not improve the fit of the basic model
(see Supplementary Information), and so the basic models were retained.

Spatial Interaction Analyses

Spatial interaction between experimental plovers was estimated, using the Utilisation Distribution
Overlap Index (UDOI) with the R package adehabitatHR (Calenge, 2011), as a proxy for
behavioural connectivity within plover populations: greater overlap between home ranges indicates higher levels of space sharing and greater opportunity for social interaction, and potentially, increased gene flow through the population. The UDOI is an estimate for space use
sharing between individuals (Fieberg & Kochanny, 2005; Chynoweth et al., 2015), which utilises the UD described in the previous section. Thus, UDOI indices were calculated between the UD of individual plovers monitored during the study period. UDOI values range from 0.0 to 2.0, a value less than one indicates less overlap than expected whereas a value above one indicates higher overlap than would be expected relative to uniform space use (Fieberg & Kochanny, 2005). All interactions were temporally constrained, so that interactions between experimental individuals that had no temporal overlap in relocations were not included in the analysis. Although the two species had different numbers of individuals for the interaction analyses (32 Kittlitz’s and 24 white-fronted plovers) and so UDOI could not be directly compared, we calculated two further characteristics using UDOI (see below): the (i) interaction network density, and (ii) relative spatial overlap between sexes. These measures are suitable for comparison as they describe overall network structure and are not affected by group size (Wey et al., 2008).

The spatial interaction network was produced using the R package igraph (Csardi & Nepusz, 2006). For each species an interaction matrix was created of UDOI weighted ties (edges) between individuals (nodes) where UDOI was positive. Thus, a node represents an individual plover, and edges represent its spatial interactions between individuals within the sampled population. Network density was then calculated as the proportion of potential edges, i.e. all of the possible interactions, which were observed in the network, i.e. UDOI greater than 0 (Wey et al., 2008). Standard errors for the species interaction network densities were calculated using 9,999 network bootstraps (Snijders & Borgatti, 1999), and two sample t-tests were carried out to test whether interaction networks significantly differed between species.
To test whether the spatial interaction network functioned as a suitable proxy for behavioural connectivity, a social interaction matrix was created using behavioural observations of experimental white-fronted plovers after release. Edges were weighted by the number of 30 second intervals in which either courtship or fighting behaviour was recorded with other experimental individuals, during the 30-minute observation period taken after each relocation. The observed interaction matrix was then compared to the UDOI matrix using a partial Mantel test, utilising the R package ecodist (Goslee & Urban, 2007) with the distances between territories, i.e. the centroid point of the 50% home range core-use area (see above), of individuals included as the control matrix. Data were not available to create a behavioural interaction matrix for Kittlitz’s plover.

As overlap size is dependent on the number of conspecific experimental individuals within the study area, it is not possible to compare overlap, i.e. the UDOI value, directly between groups of differing network sizes, unlike network density. Hence sex-specific interactions were investigated separately for each species; total spatial interaction between individual plovers and either conspecific males, or females, was calculated. GLMs with Gaussian error structure and identity link function were fitted with sex as a predictor variable, and response variables of total UDOI between the focal bird and (i) all individually marked males, and (ii) all individually marked females involved in the study. The response variables, Y, were transformed to a normal distribution by adding one and then log transforming, i.e. ln(Y+1).

Spatial analysis was carried out in R (R Core Team, 2015) and ArcGIS 10.4 (Esri, Redlands, CA, USA), and spatial data was converted between them using R packages maptools (Bivand & Lewin-Koh, 2016) and rgdal (Bivand et al., 2015). Figures were produced using the R packages ggplot2
(Wickham, 2009) and igraph (Csardi & Nepusz, 2006). Additionally, functions from the R packages plyr (Wickham, 2011) and sp (Bivand et al., 2013) were used during the analysis.

Both experiments were approved by the Ministry of Environment, Forests and Tourism of the Republic of Madagascar (Research permit No: 053/11/MEF/SAP/SCB of 11 March 2011 and 132/10/MEF/SAP/SCB of 6 May 2010) and Madagascar National Parks (No: 398-10/MEF/SAP/SCB of 18 May 2011). Blood sampling was also covered by these research permits. The blood transport permit was approved by Service de la Gestion de la Faune et de la Flore, Direction de la Valorisation des Resources Naturelles, Ministère de l'Environnement et des Forêts Madagascar (authorization number 080N-EA06/MG11).

Kittlitz’s and white-fronted plovers are common breeding birds in much of Africa and Madagascar and not considered threatened by the IUCN (BirdLife International, 2014).

The experiment was designed to reduce adverse effects on local plover populations and all necessary precautions were taken to ensure their welfare was suitably protected. Captive plovers were monitored daily and kept under standard conditions (see Parra et al., 2014) to reduce their stress levels. In addition, translocated eggs coped with the natural breeding conditions of local clutches in the two plover populations. Although monitoring the augmented clutches was beyond the scope of the experiment, nest checks suggest that at least 33.3% and 19.4% of augmented nests survived until hatching in the Kittlitz’s plover (N = 36 nests) and the white-fronted plover (N = 20 nests), respectively. Survival in these nests appeared to be higher than for unmanipulated nests (13.4% and 8.9%, based on N = 101 Kittlitz’s plover nests and N = 56 white-fronted plover nests, respectively; J.E. Parra et al., unpublished data).
RESULTS

Home Range Size and Movement

Kittlitz’s plovers had significantly larger home ranges (9.02 ± 8.21 ha, N = 32 plovers) than white-fronted plovers (3.27 ± 4.74 ha, N = 24 plovers; Table 1), although home range sizes did not differ between males and females (Figure 2, Table 1). Kittlitz’s plovers also had a higher mean step length (223.8 ± 194.1 m, N = 34 plovers) than white-fronted plovers (94.0 ± 117.3 m, N = 26 plovers), and a marginally significant species * sex interaction suggests sex-difference between the two species (Table 1). Although the mean step duration was longer in Kittlitz’s plover (two sample t-test: t_{54} = 3.84, p < 0.001), days of tracking (duration) did not explain significant variation in mean step length (Table S2).

Contrary to expectations, the distance between the former territory and the new territories was not different between Kittlitz’s plovers and white fronted plovers, nor did it differ between males and females (Table 1). However, Kittlitz’s plovers were found to have greater variation in their breeding dispersal distances than white-fronted plovers (F-test: F_{31,23} = 0.290, p = 0.003, Figure 2).

Spatial Interaction

Kittlitz’s plovers were more spatially interconnected than white-fronted plovers. The density of the Kittlitz’s plover spatial association network (0.742 ± 0.093 [SE], N = 32 plovers) was significantly higher (two sample t-test: t_{54} = 4.399, p < 0.001, Figure 3) than that of the white-
fronted (0.284 ± 0.047 [SE], N = 24 plovers). Additionally, the white-fronted spatial association network was significantly correlated with the observed behavioural interaction network (partial Mantel test: $r_m = 0.351$, $p < 0.001$) controlling for distance between territories.

The spatial interactions of Kittlitz’s plovers were less sexually structured than those of white-fronted plovers (Figure 3). In Kittlitz’s plover, an individual’s spatial overlap with both males and females was not predicted by the sex of the interacting individual indicating a lack of sex-specific spatial interactions (GLMs, males: $t = 1.633$, $p = 0.113$; females: $t = 1.341$, $p = 0.190$; Figure 4; Table 2). In contrast, male white-fronted plovers had more spatial interaction with females than males (GLM: $t = 4.137$, $p < 0.001$; Figure 4). Likewise, female white-fronted plovers had a larger amount of spatial interaction with males than females (GLM: $t = 5.652$, $p < 0.001$; Figure 4; Table 2).

**DISCUSSION**

The analyses of experimental data that have not been presented previously showed that spatial movement and interaction of unmated individuals varies between closely-related species. These results augment the analyses of Parra et al. (2014) that reported different mating times, courtship behaviour and pair bonds between the two Malagasy plover species. The work presented here provided two novel results; species differences in both spatial behaviour and inferred social interactions with conspecifics.

Firstly, both mean step length and home range size were larger in Kittlitz’s than in white-fronted plovers. This result showed that the polygamous Kittlitz’s plovers exhibit less restricted
movements than the socially (and genetically) monogamous white-fronted plover. The smaller home ranges in white-fronted plover may be due to more restricted mate search behaviour and/or to more limited movements of experimental birds once they found a mate. We believe both explanations are likely and more work is needed to disentangle the movements of un-mated and mated individuals possibly by recording the movements of radio-tagged individuals. Furthermore, although field observations were carried out as consistently as possible between the two species, the relocations were not necessarily uniformly spaced between the two species.

Contrary to expectation, new Kittlitz’s and white-fronted plover territories were similar distances from their previous territories, and hence the breeding dispersal distances do not seem to explain differences in gene flow within these populations of plovers. However, the other measures of spatial distribution and search effort did align with the observed genetic structure, and for future studies we recommend using territorial or social metrics alongside distance metrics where possible to understand fine-scale spatial patterns. Other studies have also found social interactions and spatial behaviour explaining gene flow, in addition to or in absence of dispersal (Burland et al., 1999; Pilot et al., 2010; McGuire et al., 2013). Although migration did not predict genetic differentiation in shorebirds (D’Urban-Jackson et al., 2017) it remains important to consider the role between-season movement plays, such as natal dispersal (Ronce, 2007; Mabry et al., 2013), which is not investigated in this study. Further work is needed to distinguish the relative importance of natal versus breeding dispersal in generating gene flow (Wey et al., 2015). Importantly, although there was no difference in the dispersal distance between the species, Kittlitz’s plovers showed greater variation in their dispersal distances: the furthest Kittlitz’s plover dispersed was 2202m, compared to 462m in white-fronted plover. This demonstrates the capability
to disperse greater distances within a breeding season, and deserves further study in the context of other forms of dispersal.

Secondly, consistent with our predictions, Kittlitz’s plovers were more spatially interconnected than white-fronted plovers. The Kittlitz’s plover spatial interaction network density was significantly higher, and therefore birds likely interact with higher numbers of conspecific experimental individuals. The spatial association network of white-fronted plover correlated with the observed social interaction network, indicating that results from the spatial interaction network may also be considered in the context of a social network.

In Kittlitz’s plover, home ranges of both males and females overlapped with several other experimental individuals. The high levels of spatial interaction suggest flocking behaviour; Kittlitz’s plovers exhibit complex gregarious social behaviour where individual plovers join to flock for feeding and resting; even members of breeding pairs join flocks (Urban et al., 1986; Hockey et al., 2005), but the relative numbers of paired and un-paired individuals within these flocks is not currently known. This greater degree of sociality increases the potential for high levels of gene flow across a population; however, a recent study of genetic structure in the social, but monogamous, red-billed chough Pyrrhocorax pyrrhocorax found strongly segregated populations (Morinha et al., 2017), suggesting both mating opportunities and social interaction are needed to facilitate high levels of gene flow. Flocking behaviour may facilitate gene flow through lower energy costs associated with mate searching due to high densities, and reduced risk of predation while searching (DeRivera et al., 2003; Kasumovic et al., 2007).
Kittlitz’s plover’s exhibit uni-parental brood care, whereas white-fronted plovers are biparental (Zefania & Székely, 2013), and this brood care strategy may allow them to interact more frequently with other members of the population due to less time spent on parental care. In contrast, white-fronted plovers exhibit greater philopatry and are less social when searching for a mate, and males and females had few interactions with conspecifics, inherently leading to strongly spatially structured populations. This suggests polygamous plover species have a plastic, flexible social structure which spreads over a broad geographical range (Küpper et al., 2012; Eberhart-Phillips et al., 2015), whereas monogamous plovers exhibit social rigidity with few social interactions within a restricted home range. A recent genetic analysis of 79 geographically distinct populations of 10 plover species provided consistent results with the latter argument, since polygamous plovers exhibited less geographic differentiation than monogamous ones (D’Urban Jackson et al., 2017).

Additionally, interactions between plovers were not sexually structured in Kittlitz’s plovers, but were in white-fronted. We found that a significant difference in spatial overlap between the sexes was only found in white-fronted plovers; overlap with the opposite sex was significantly higher for both males and females than same-sex overlaps. Small home range overlap with same-sex individuals, combined with the previous results of fewer interactions and less movement, suggests strong territoriality in white-fronted plovers (Ostfeld, 1986). This fits in with the expectation that pair bonding and bi-parental care will generally see an individual be more fixed in its range of movement (Fricke, 1986; Sommer, 2003). In line with the latter argument, female pied flycatchers Ficedula hypoleuca exhibit similar restricted mate searches as a consequence of competition with conspecifics for nest sites (Slagsvold et al., 1988).
An explanation of the spatial patterns observed in white-fronted plover may be the costs of finding nest sites leading to high territoriality (Brashares & Arcese, 1999). Strong competition for breeding vacancies would prevent formation of a floating population of single plovers as any paired plover has a high probability of losing their breeding status if they leave a nesting territory in an attempt to find a new mate. Therefore, if the costs of searching are related to defending a nesting site, males and females should stay together to protect a territory and spend less time searching for a mate to reduce the risk of losing both their nest site and breeding status (Ens et al., 1996).

Consistently, white-fronted plovers exhibit high breeding site-fidelity and territory retention within and between years (Lloyd, 2008). Consequently, monogamy and biparental care associated with territoriality are probably the best strategy to maximize reproductive success in white-fronted plovers as demonstrated in other shorebird species (Lessells, 1984; Gratto et al., 1985). Conversely, breeding sites do not seem to be limited for Kittlitz’s plovers, they can breed with nests of different pairs 10 – 30 metres apart (Urban et al., 1986; Hockey et al., 2005). Hence, the observed tight sexually-structured spatial behaviour in white-fronted plovers may be indicative of territoriality, resulting in low gene flow through a population due to confined search behaviour and low interaction rate.

In conclusion, we found different spatial movements and inferred social interaction patterns in unmated individuals of closely related plover species exhibiting different breeding systems. These findings may have important implications for the role of spatial interaction in gene flow and speciation; as well as how spatial behaviour and social interactions are shaped by competition, mating opportunities and territoriality. Taken together, the different spatial behaviour and social structure in two sympatric plover populations are consistent with molecular results obtained from
populations along the west coast of Madagascar (Eberhart-Phillips et al., 2015). Our study demonstrates that spatial and network analyses provide valuable tools in investigating, and quantifying, how social interaction, competition and mating strategies impact on gene flow and speciation. In particular, we emphasise the need for social and/or territory metrics to be used in accordance with distance metrics when investigating genetic structure. Future investigations with detailed movements of focal species, in an explicit phylogenetic framework, are needed to fully understand the roles of mating system and social interaction, as well as the relative importance of intra- and between-season movements in speciation.

References


Table 1. General linear models of home range size, mean step length and dispersal from previous territory (response variables) of male and female Malagasy plovers. Figures in bold indicate statistically significant relationships. The data were natural log transformed prior to the analyses.

<table>
<thead>
<tr>
<th></th>
<th>Home range size</th>
<th>Mean step length</th>
<th>Dispersal from previous territory</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
<td>t value</td>
<td>Estimate</td>
</tr>
<tr>
<td></td>
<td>(0.241)</td>
<td>(&lt;0.001)</td>
<td>(0.171)</td>
</tr>
<tr>
<td>Species</td>
<td>-1.264</td>
<td>-3.374</td>
<td>-0.598</td>
</tr>
<tr>
<td></td>
<td>(0.375)</td>
<td>(0.001)</td>
<td>(0.259)</td>
</tr>
<tr>
<td>Sex</td>
<td>-0.074</td>
<td>-0.211</td>
<td>0.315</td>
</tr>
<tr>
<td></td>
<td>(0.352)</td>
<td>(0.833)</td>
<td>(0.241)</td>
</tr>
<tr>
<td>Species * Sex</td>
<td>-0.121</td>
<td>-0.225</td>
<td>-0.647</td>
</tr>
<tr>
<td></td>
<td>(0.537)</td>
<td>(0.823)</td>
<td>(0.366)</td>
</tr>
</tbody>
</table>

General linear models using Gaussian error structure and identity link function were fitted separately to home range size, mean step length and dispersal from previous territory. These models were compared with models including additional control variables; these were found not to improve the model fit, and hence the more basic models were sufficient (see Supplementary Information). Standard errors and p-values are in presented in parentheses for the estimates and t-values, respectively. ‘Female’, ‘white-fronted’, and ‘white-fronted female’ were used as reference levels.
Table 2. General linear models of total spatial overlap of the focal individual with males, and females (response variables) in two species of Malagasy plovers. Figures in bold indicate statistically significant relationships. The data were natural log transformed prior to the analyses.

<table>
<thead>
<tr>
<th></th>
<th>Male overlap</th>
<th>Female overlap</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
<td>t value</td>
</tr>
<tr>
<td><strong>White-fronted</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>0.191</td>
<td>3.061</td>
</tr>
<tr>
<td></td>
<td>(0.062)</td>
<td>(0.006)</td>
</tr>
<tr>
<td>Sex</td>
<td>0.365</td>
<td><strong>4.137</strong></td>
</tr>
<tr>
<td></td>
<td>(0.088)</td>
<td>(&lt;0.001)</td>
</tr>
<tr>
<td><strong>Kittlitz’s</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>0.733</td>
<td>7.531</td>
</tr>
<tr>
<td></td>
<td>(0.097)</td>
<td>(&lt;0.001)</td>
</tr>
<tr>
<td>Sex</td>
<td>0.232</td>
<td>1.633</td>
</tr>
<tr>
<td></td>
<td>(0.142)</td>
<td>(0.113)</td>
</tr>
</tbody>
</table>

General linear models using Gaussian error structure and identity link function were used to analyse spatial overlap. As inter-species analysis was not appropriate due to differing sample sizes, the only explanatory variable included was Sex, with Female the reference factor level. Estimate standard errors and t values’ corresponding p-values are in parentheses.
Supplementary Information

Molecular sexing

A small blood sample was taken from each adult's brachial wing vein, by puncturing, collecting drops of blood (25 µl approx.) in capillary tubes, and storing this in Eppendorf tubes of Queen's Lysis Buffer. DNA was extracted from blood samples using the ammonium acetate extraction method (Parra et al., 2014). For molecular sex typing, Z- and W-chromosome-specific genes were amplified via polymerase chain reaction (PCR) using Z-002B/Z-002D primers. For additional certainty in sex assignment, W-chromosome-specific Calex-31 primers, developed in the genus Charadrius, were utilized (Parra et al., 2014). PCR amplification was conducted on a DNA Engine Tetrad 2 Peltier Thermal Cycler under the following conditions: 95 °C for 15 min, followed by 35 cycles of 94 °C for 30 s, 56 °C for 90 s, 72 °C for 60 s with a final extension of 60 °C for 30 min. Samples were visualized on an ABI 3730 automated sequencer. IR Dye-labelled tailed primers separated the products of Z-002B/Z-002D primers into either one (ZZ) or two (ZW) bands, indicating male or female, respectively. The W-specific Calex-31 product only appeared as one band indicating female. Images were scored using GeneMapper software version 4.1 (Applied Biosystems, Foster City, CA, U.S.A.). To maximize reliability, all samples were sexed using two markers. Additionally, for 8% of the total samples (11 Kittlitz's plover and 10 white-fronted plover individuals) molecular sexing was repeated; in all cases, repetitions concurred with the original results.

Table S1. Home range sizes of male and female Malagasy plovers.
Two GLMs were used to analyse home range size using Gaussian error structure and identity link function. The first including only Sex, Species, and their interaction; and the second including a number of additional control variables (above). The two models were tested and including control variables was not found to improve the fit of the model (Analysis of Deviance, F = 2.371, df_{large} = 52, df_{small} = 49, p = 0.082). Duration refers to the number of days that a plover was tracked in the field; captivity, whether or not the bird was held until its previous mate re-mated; and number of relocations, the number of GPS recordings taken. Reference factor level, estimate standard errors and t values’ corresponding p values are in parentheses.

**Table S2.** Mean step length of male and female Malagasy plovers.

<table>
<thead>
<tr>
<th></th>
<th>Model 1</th>
<th></th>
<th>Model 2</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
<td>t value</td>
<td>Estimate</td>
<td>t value</td>
</tr>
<tr>
<td></td>
<td>(0.171)</td>
<td>(&lt;0.001 ***)</td>
<td>(0.339)</td>
<td>(&lt;0.001 ***)</td>
</tr>
</tbody>
</table>
Two GLMs were used to analyse home range size using Gaussian error structure and identity link function. The first including only Sex, Species, and their interaction; and the second including a number of additional control variables (above). The two models were tested and including control variables was not found to improve the fit of the model (Analysis of Deviance, $F = 0.387$, $df_{\text{large}} = 56$, $df_{\text{small}} = 54$, $p = 0.681$). Duration refers to the number of days that a plover was tracked in the field; and captivity, whether or not the bird was held until its previous mate re-mated. Reference factor level, estimate standard errors and t values’ corresponding p values are in parentheses.

**Table S3.** Breeding dispersal of male and female Malagasy plovers.

<table>
<thead>
<tr>
<th></th>
<th>Model 1</th>
<th>Model 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
<td>t value</td>
</tr>
<tr>
<td><strong>Species</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(white-fronted)</td>
<td>-0.598</td>
<td>-2.310</td>
</tr>
<tr>
<td></td>
<td>(0.259)</td>
<td>(0.025 *)</td>
</tr>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(female)</td>
<td>0.315</td>
<td>1.307</td>
</tr>
<tr>
<td></td>
<td>(0.241)</td>
<td>(0.197)</td>
</tr>
<tr>
<td><strong>Captivity</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(yes)</td>
<td>0.091</td>
<td>0.474</td>
</tr>
<tr>
<td></td>
<td>(0.192)</td>
<td>(0.638)</td>
</tr>
<tr>
<td><strong>Duration</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.008</td>
<td>0.832</td>
</tr>
<tr>
<td></td>
<td>(0.009)</td>
<td>(0.409)</td>
</tr>
<tr>
<td><strong>Species*Sex</strong></td>
<td>-0.647</td>
<td>-1.767</td>
</tr>
<tr>
<td>(white-fronted female)</td>
<td>(0.366)</td>
<td>(0.083 .)</td>
</tr>
</tbody>
</table>

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Two GLMs were used to analyse home range size using Gaussian error structure and identity link function. The first including only Sex, Species, and their interaction; and the second including a number of additional control variables (above). The two models were tested and including control variables was not found to improve the fit of the model (Analysis of Deviance, $F = 0.525$, $df_{large} = 52$, $df_{small} = 49$, $p = 0.667$). Duration refers to the number of days that a plover was tracked in the field; captivity, whether or not the bird was held until its previous mate re-mated; and number of relocations, the number of GPS recordings taken. Reference factor level, estimate standard errors and t values’ corresponding p values are in parentheses.

**Figure legend**

**Figure 1.** Maps of study sites of Kittlitz’s and white-fronted plover in SW Madagascar, with the study area in the left panel and illustrative detail in the right. Dashed lines represent the breeding dispersal between the original nest capture sites (denoted by crosses), and secondary territories of male (white circles) and female (black circles) experimental plovers used in the spatial analyses. As an illustration of data used in analysis, the home ranges of three male (white fill, solid outline) and female (grey fill, dashed outline) Kittlitz’s plovers and five white-fronted male and female plovers are shown.

**Figure 2.** (a) Home range size, (b) mean step length and (c) dispersal from previous territory in two Malagasy plover species. The data were normalised using natural log transformations.

<table>
<thead>
<tr>
<th>Species*Sex</th>
<th>-0.242 (0.749)</th>
<th>-0.323 (0.748)</th>
<th>-0.161 (0.775)</th>
<th>-0.208 (0.836)</th>
</tr>
</thead>
</table>
Numbers of individuals are shown beneath categories. The lower and upper borders of the box are lower and upper quartiles, respectively; the horizontal bar is the median; and whiskers represent the lowest and highest observations apart from the outliers. Circles denote outliers that are between the first and third interquartile from the nearer edge of the box.

**Figure 3.** Spatial association networks of Kittlitz’s and white-fronted experimental plovers. Nodes represent adult males and females; vertices represent the amount of overlap (UDOI) of individual’s home ranges. The Kittlitz’s network was more interconnected than the white-fronted network, as the densities differed significantly (two sample t-test, $t_{54} = 4.462, p < 0.001$).

**Figure 4.** Total spatial overlap of individual home ranges with conspecific experimental males or females in white-fronted and Kittlitz’s plover, quantified using the utilization distribution overlap index (UDOI). Inter-species comparison of UDOI size is not appropriate due to differing sample sizes, but the relationship between male and female overlap within species can be compared. The lower and upper borders of the box are lower and upper quartiles, respectively; the horizontal bar is the median; and whiskers represent the lowest and highest observations apart from the outliers. Circles denote outliers that are between the first and third interquartile from the nearer edge of the box.

**Figure S1.** Typical spatial data collected. The home range size is the area of the 90% home range (shaded grey). The mean step length was calculated as the total distance between consecutive relocation points (solid circles), divided by the number of points. The breeding dispersal (dashed line) was taken to be the distance between territories; between the original nest location (hollow circle, moved for visualisation purposes) and the centroid of the 50% core use area (hollow square).