Across-year social stability shapes network structure in wintering migrant sparrows

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Abstract

Migratory birds often form flocks on their wintering grounds, but important details of social structure such as the patterns of association between individuals are virtually unknown. We analyzed networks of co-membership in short-term flocks for wintering golden-crowned sparrows (Zonotrichia atricapilla) across three years and discovered social complexity unsuspected for migratory songbirds. The population was consistently clustered into distinct social communities within a relatively small area (~7 ha). Birds returned to the same community across years, with mortality and recruitment leading to some degree of turnover in membership. These spatiotemporal patterns were explained by the combination of space use and social preference—birds that flocked together in one year flocked together again in the subsequent year more often than were expected based on degrees of home range overlap. Our results suggest that a surprising level of social fidelity across years leads to repeatable patterns of social network structure in migratory populations.

Keywords: Flocks, group living, MRQAP, social networks, spatial communities, temporal stability, winter ecology

Introduction

The social structure of animal populations—e.g. the size, composition and stability of social groups—is a fundamental aspect of social evolution (Alexander 1974). In birds, studies of breeding systems have shown that ecological conditions can favor different social structures ranging from simple pairs to cooperative breeding groups (Emlen 1982). The winter social structure of year-round resident birds has also been investigated, but to a lesser degree than for the breeding season (Ekman 1989; Kraaijeveld & Dickinson 2001; Aplin et al. 2012). In migratory birds, the most basic aspect of winter social structure is known for many species—e.g. territoriality versus flocking in social groups. However, in species that form flocks (defined here as temporary aggregations of individuals in the same place at the same time), we know almost nothing about dynamics of flock membership over space and time (see Myers 1983; Piper & Wiley 1990; Conklin & Colwell 2008 for notable exceptions).

Our lack of understanding of the winter societies of small-bodied birds is particularly surprising because these taxa were so crucial to the development of important theories in ecology. A large body of influential research on small-bodied birds in winter explored how food, predation and sociality interact to affect the evolution of optimal foraging (Stephens & Krebs 1986), sociality and optimal group size (Pulliam & Caraco 1984), energy management (Cuthill & Houston 1997), predator–prey interactions (Bertram 1978) and status signals (Rohwer 1975; Rohwer & Ewald 1981). For many of these topics, the pattern of group stability and the specific identities of group members matter. For example, the degree to which individuals form long-term associations could alter the dynamics of anti-predator behaviors and the form of cooperation involved (Croft et al. 2006; Micheletta et al. 2012). In addition, the pattern of social structure also has critical implications for the mechanisms by which intragroup competition is mediated by signals (Rohwer 1975).

In theory, the social structure of wintering birds could range from the small, highly stable groups observed in a variety of year-round resident birds (e.g. Ekman 1989) to short-term random associations with little or no structure (Myers 1983; Conklin & Colwell 2008). Between these two extremes, winter bird societies could also involve a complex mix of social stability and change in both space and time—often termed fission–fusion dynamics (Aureli et al. 2008). Migration poses an added challenge to across-year stability because individuals that winter together do not necessarily breed together (Ryder et al. 2011; Seavy et al. 2012), and thus long-term social bonds must bridge a break in contact between winter seasons. However, high levels of site fidel-
ity and long-term memory of individuals can promote social stability between neighbors in the breeding season (Goddard 1991), and the same type of stability could exist on the wintering grounds. Even in birds that switch flocks over short time scales (e.g. min, h), social stability can occur over longer time spans (e.g. days, seasons) if certain sets of birds tend to join flocks together more often than expected by chance. The challenge for researchers is to use observations of short-term dynamics to detect the underlying patterns of social associations as well as changes in social structure across time—a task that has become more tractable with recent advances in social network analysis (Wey et al. 2007; Croft et al. 2008; Pinter-Wollman 2014).

Social network analysis has recently emerged as a quantitative framework for understanding patterns of social structure in animals, and the number of studies documenting non-random social structure has exploded. Many of these studies have used network theoretical methods to detect social communities—clusters of individuals that associate with each other more often than expected by chance (Newman 2006) in populations of animals such as fish, birds and mammals (e.g. Wolf et al. 2007; Oh & Badyaev 2010; Mouri er et al. 2012). Here, we use social network analysis to determine the social structure of a wintering population of a migrant species, the golden-crowned sparrow (Zonotrichia atricapilla). Our goal was to determine whether flocks represent aggregations that arise from discrete social communities within the population, and whether long-term social preferences play a role in structuring such social communities.

We constructed networks of flock co-membership in three sequential non-breeding seasons. We show for the first time that the social lives of migrant birds in winter are characterized by complex community structure and that these social communities are remarkably stable across years despite the potentially destabilizing effects of recruitment, mortality and long-distance migration. We demonstrate that stable social associations among birds returning across years help shape these patterns of network structure.

**Disentangling the effects of space use and sociality**

A major difficulty in analyses of social networks is to understand whether social communities arise as a consequence of shared spatial preferences, preferential social associations between some individuals or both (Pinter-Wollman et al. 2014). By definition, associations among individuals are determined on the basis of close spatial proximity, so the key question is whether an association reflects shared preference for the same space, or whether the animals use the same space because they prefer to associate with each other. One approach to teasing apart some of the influences of spatial preference from social preference is to compare empirical networks against spatially explicit null models, e.g. by building simulated networks that randomly group individuals into short-term flocks based on the overlap in their broader patterns of use of space (Ramos-Fernández et al. 2006; Best et al. 2014). This null model approach is conservative for testing the effects of social preferences because it cannot account for the fact that space use patterns themselves can reflect social preferences. For example, a lack of difference in structure between an empirical network and networks generated by a spatially explicit null model cannot reject the possibility that social processes were important in determining individual home ranges in the first place. However, demonstrating that empirical networks exhibit significantly more structure than predicted from a spatially explicit null model can provide evidence that social preferences among certain individuals plays some role in structuring the community. Thus, this approach identifies the minimum contribution of social preferences above and beyond their role in determining the patterns of overlapping home ranges of individuals. A complimentary approach is to use network regression methods to determine whether specific social attributes (e.g. familiarity) predict patterns of associations in flocks independently of space use patterns. While this approach still suffers from the potential influence of social preference on space use, it can help identify specific social factors that influence network structure. In this study, we employ both null model and network regression approaches to determine how social preference shapes social network structure across time.

**Methods**

**Constructing social networks**

Sparrows arrive at our study site (The University of California, Santa Cruz Arboretum) in October–November and depart for their breeding grounds, likely in disparate areas along the Alaskan coast (Seavy et al. 2012), in March–April each year. Our study spanned three non-breeding seasons: January–March 2010 (Season 1), October 2010–February 2011 (Season 2) and October 2011–April 2012 (Season 3). Each year, we captured birds using baited traps and attached individually unique combinations of color bands. In Season 2, we did not band any birds between October–December 2010. We censused short-term flocks (defined as a group of individuals found within a single 5 m radius) by identifying the color-banded individuals in each flock. Most censuses were conducted while the flock was foraging in short grass and more rarely when they were foraging on shrubs and trees. The birds are habituated to people and we typically observed birds from a distance of about 10 m. In most cases, we left the flock once all individuals were identified before we lost sight of the flock. We also noted the location where the flock was first observed, using an aerial photograph with 10 x 10 m grids (Figure 1) as a map. To ensure independence of our data points, our samples included flocks censused at least 20 min apart, as preliminary analysis suggested that flock membership often changes within this time frame. We also included flocks censused less than 20 min apart as independent data points if they contained no more than one individual in common. We only included flock observations that occurred away from feeders used for trapping. A minority of observations (≤ 10%) in Seasons 2 and 3 were conducted on days when feeders were placed elsewhere within our study site at some point during the day and excluding these observations did not affect our results.

We included in our analysis only birds with confirmed band combinations and those banded prior to the beginning of the census period for each season. Thus, birds banded early in the season (October–December) were included in Season 1 because flock censuses began in January, but early-banded birds were excluded in other seasons when flock observations began in October. Inclusion of individuals banded later in the season would cause us to underes-
Figure 1 Social communities and their spatial distribution across years. (a, c, e) Social networks of flock co-membership in each season. Each node, representing an individual bird, is assigned to a social community, denoted by node color and colored bubbles. Edge widths are proportional to the association index. Edges connecting nodes in different social communities are drawn in red. Node placement is determined by a force-directed algorithm from the igraph package (Csárdi & Nepusz 2006), which tends to place strongly associated nodes closer together. (b, d, f) Social community home ranges for each season overlaid on an aerial photo and grid of the study area. The filled areas and outlines represent 50% and 90% utilization density estimates respectively (see Appendix S2 for methods). Each cell of the study grid is 10 x 10 m. (d) For visual clarity, we excluded the minor community of three individuals in Season 2 (shown in white) from the home range plot.
timate the strength of their network relationships as prior to banding they would not be identified even if present in a flock. We also omitted from the network analysis transient individuals that were observed fewer than three times, but the major results are not affected by their removal. For each season, we calculated the Simple Ratio association index (Cairns & Schwager 1987) for each pair of individuals, which ranges from 0 for pairs never seen in the same flock and 1 for pairs always seen in the same flock. We then constructed social networks using the individuals as nodes and association index as edge weights. The flock composition data are available on the Dryad Digital Repository (doi: 10.5061/dryad.d3m85).

Detecting social communities in empirical and simulated networks

We used a modularity-optimization community detection approach to detect social communities (Newman 2006)—groups of individuals that are tightly connected with each other through flock co-membership. Modularity (Q) refers to the weighted proportion of edges that occur within a pre-defined group, minus the expected proportion of such edges if edges were distributed randomly in the network. Modularity-optimization community detection refers to a class of methods to search for the groupings of nodes that result in the maximum modularity value (Q$_{max}$). The value of Q$_{max}$ is used as a metric of how discretely a network is divided into distinct clusters, and we later statistically compare empirical Q$_{max}$ values to those of null models. We tried several different community detection algorithms using the R package “igraph” (Csárdi & Nepusz 2006). We used the method of Clauset et al. (2004) for our final analysis because it consistently yielded the highest Q$_{max}$ value (i.e. the optimal community division) among these community detection methods. This method allowed us to assign each individual to a social community for each season.

We used a bootstrapping technique to account for sampling error in our observed networks (Lusseau et al. 2008). We resampled flocks with replacement up to the number of flocks observed (Table 1) to create a re-sampled network. We calculated Q$_{max}$ for 1000 re-sampled networks to generate bootstrapped confidence interval for our empirical measure of modularity. We used the same bootstrap procedure to calculate a novel index of the robustness of community assignment, r$_{community}$ whose value is 1 when all bootstrap replicates result in the exact same community assignment as the empirical result, and 0 when community assignments in bootstrap replicates are random with respect to empirical network (see Appendix S1).

We tested whether the observed community structure was greater than expected by chance using two null models. The first null model (hereafter “Random Flock Model”) assumed that flock associations were random, but controlled for the observed sizes of flocks and number of times each bird was seen. In effect, this model simulated flocks as random aggregations of individuals occurring at separate time points such that any individual could join any flock, but with several constraints: the total number of flocks, number of individuals in each flock and the number of flocks an individual joined all matched the empirical dataset. We conducted this randomization using the “Swap” algorithm (Bejder et al. 1998; Whitehead et al. 2005) with 2*`v “swaps” for each run of the randomization procedure (v = total sum of individuals across all flocks). Preliminary analysis indicated that this was sufficient to ensure that the flock matrix had been randomized. We then applied the community detection methods as described above to measure the modularity (Q$_{max}$) of each network generated from a randomized flock matrix and then repeated this process 1000 times to produce a null distribution for comparing the empirically observed modularity measures.

Our second null model, the “Spatial Flock Model,” incorporated information about home ranges under the assumption that birds flock randomly with those that share the same space. First, we estimated the home range of each bird using minimum convex polygons based on whether the individual had been recorded in each 10 x 10 m grid cell on the map. In each iteration of the simulation, we constructed flocks using the observed flock sizes and grid cell locations from the empirical data, and drew flock members at random from among all individuals for which that grid cell location was part of their home range. Flocks were simulated to occur at separate times such that each individual could join any flock within their home range. We then used this flock membership matrix to construct social networks and measure modularity as above. We repeated this procedure 1000 times to produce a distribution of modularity values.

We compared the modularity values generated by the Random Flock Model and the Spatial Flock Model with the empirical modularity value with bootstrap confidence intervals for each year.

Testing for effects of prior social associations on network structure

Having found that the observed social networks exhibited more discrete community structure than predicted by our null models, we tested whether a specific form of social preference—social stability across years—could explain

<table>
<thead>
<tr>
<th>Season</th>
<th># Flocks observed</th>
<th>Mean # banded individuals per flock (SD)</th>
<th>N</th>
<th># Communities (# indiv. per community)</th>
<th>Q$_{max}$</th>
<th>R</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>77</td>
<td>3.3 (1.9)</td>
<td>31</td>
<td>3 (17, 9, 5)</td>
<td>0.43</td>
<td>0.15</td>
</tr>
<tr>
<td>2</td>
<td>340</td>
<td>2.2 (1.7)</td>
<td>43</td>
<td>4 (17, 12, 11, 3)</td>
<td>0.49</td>
<td>0.19</td>
</tr>
<tr>
<td>3</td>
<td>430</td>
<td>2.0 (1.4)</td>
<td>27</td>
<td>3 (14, 7, 6)</td>
<td>0.43</td>
<td>0.17</td>
</tr>
</tbody>
</table>

N = Total number of individuals in the network; Q$_{max}$ = modularity; R = The proportion of total edge weights that links nodes in different communities (denoted in red in Figure 1).

Table 1. Basic statistics for social networks
how social communities arise. For each of three comparisons
(Season 1 vs. Season 2, Season 2 vs. Season 3, Season 1 vs.
Season 3), we constructed matrices of association indices be-
tween all possible dyads of individuals that were present in
both years. We then conducted Mantel Tests to determine
whether social associations were consistent across years.

While significant correlations in association matrices
would suggest social preference for familiar individuals, this
could simply reflect spatial fidelity to home ranges and not
to familiar flock mates. To determine if social preferences
based on past social experience could be detected after con-
trolling for the degree of home range overlap, we used a net-
work regression approach called MRQAP (multiple regres-
sion quadratic assignment procedure) in which a dependent
matrix is regressed against multiple independent matrices
of the same size (Krackhardt 1988). Here, we use the matrix
of association indices between each pair of focal individu-
als (all individuals observed across two consecutive years)
as the dependent matrix, and the two independent matrices
consisted of the association indices in the previous year and
home range overlap between each pair of focal individuals.
We calculated the degree of home range overlap for each of
these pairs of birds as \( 2C/(A+B) \), where \( A \) and \( B \) are number
of \( 10 \times 10 \) m grid cells included in the home range of each in-
dividual bird (estimated by minimum convex polygons), and
\( C \) is the number of grid cells included in both home ranges.
This analysis was conducted using the package “statnet” in R
(Handcock et al. 2003).

Having shown that individuals flock with the same birds
across years, we were interested in whether these long-term
social ties play a role in shaping the community structure of
the social network. For example, strong ties between individ-
uals that return to the same social community across years
could generate community clusters in the network. To test
this idea, we measured how well-connected an individual
was within its own social community using a normalized
score of within-community node strength \( z_i \) (Guimerá &
Amaral 2005). If \( K_i \) is the sum of association indices of indi-
vidual \( i \) to other individuals in its own social community, \( s_j \),
\( K_{si} \) is the average of \( K_i \) over all the nodes in \( s_j \), and \( \sigma_{K_{si}} \)
is the standard deviation of \( K \) in \( s_j \), then:

\[
z_i = \frac{K_i - K_{si}}{\sigma_{K_{si}}}
\]

We used an ANOVA to compare the within-community
strength of individuals that returned across years with those
that were included in the network for the first time. Note
that in Seasons 2 and 3, we only included birds that were
previously banded (see above). Thus, we are comparing the
within-community strength of individuals that returned to
the population for a second year with those that returned
three times or more.

Results

*Microgeographical community structure*

In this study, we defined *flocks* as temporary aggregations
of individuals in the same place at the same time. In contrast,
*social communities* are clusters of individuals that are tightly
connected within the social networks constructed from pat-
terns of flock co-membership throughout the season. Thus,
each social network represents the cumulative pattern of flock
associations over the course of several months, and social
community structure represents the partitioning of the
population into clusters of individuals that flock more of-
ten with each other than expected. In each of three seasons,
the network could be partitioned into three main social com-
munities (Figure 1, Table 1; Season 2 network has an addi-
tional 4th community of three individuals). The assignments
of individuals to communities were generally robust to sam-
pling error: pairs of individuals in the same community in the
empirical network were usually in the same communities in
bootstrap replicate networks (Appendix S1, Figure S1
Figure S2; Season 1 \( r_{community} = 0.82 \); Season 2 \( r_{community} = 0.76 \);
Season 3 \( r_{community} = 0.82 \)). The communities had spatially
overlapping home ranges with relatively discrete core areas,
though the degree of overlap varies among pairs of clusters
and across years (Figure 1).

Modularity differed significantly between the Random
Flock Model, the Spatial Flock Model and the empirical net-
work in each of the three years (Figure 2; all ANOVA com-
parisons \( P < 0.001 \); statistical results shown in Table S1). The
modularity of the empirical network was much greater than
expected from the Random Flock Model, indicating that the
observed network structure was highly non-random. The
Spatial Flock Model, which incorporated information about
individual home ranges, also was significantly more modu-
lar than the Random Flock Model, demonstrating that spa-
tial segregation of home ranges per se leads to some level of
community structure. However, empirical networks had still
higher modularity values than the Spatial Flock Model; evi-
dence that the population is more discretely divided into so-
cial communities than expected from the pattern of home
range overlap. Thus, flocks are unlikely to be random ag-
grations of birds that share home ranges, and these ob-
servations suggest a role for social preferences above and
beyond the role that sociality plays in determining home
ranges. However, it remains unclear from these results alone
whether the community structure arises from short-term pre-
ferences (e.g. preference to continue flocking with the
same group over the course of a day) or long-term dynamics.
We now show that these social preferences stem from long-
term familiarity that persists across years.

*Individuals prefer to flock with the same individuals
across years*

Patterns of social associations were strikingly consistent
across years. Mantel tests revealed very high correlations in
association indices between pairs of individuals that returned
across consecutive years (Season 1 vs. Season 2: \( n = 20 \) birds,
Mantel \( r = 0.63 \), C.I. = 0.49–0.71, \( P < 0.001 \); Season 2 vs. Sea-
son 3: \( n = 15 \) birds, Mantel \( r = 0.73 \), C.I. = 0.63–0.84, \( P < 0.001 \).
The consistency in association patterns also continued across
three winters (Season 1 vs. Season 3: \( n = 8 \) birds, Mantel \( r = 0.72 \),
C.I. = 0.50–0.84, \( P < 0.001 \). Such consistent associations
between individuals across years could be due to high fidel-
ity to home ranges across years, social preferences that persist
across years or both. Our network regression (MRQAP) anal-
ysis showed that associations between pairs of individuals
were significantly affected by both the degree of home range overlap and the association strength of that pair in the previous year (Table 2; see Figure S1). However, the effect of previous association on predicting flocking patterns in subsequent years was approximately two to four times stronger than the effect of home range overlap (Table 2). Thus, sparrows clearly exhibit social preference for flocking with familiar individuals with whom they flocked in the previous season.

Re-assembly of same communities across years

The social community structure of the network remained stable across years despite the fact that a significant proportion of the individuals in the social network failed to return across years (36% from Season 1 to 2, 65% from Season 2 to 3, 74% from Season 1 to 3; Table 3). The consistent community structure across years was the result of social preferences that persisted across years between pairs of individuals that returned to the population. Individuals returning to the population across consecutive years were significantly more likely than not to re-join the social community located in the same area of the study site (Table 3; binomial test with expected probability = 1/N\textsubscript{community}; Season 1 vs. Season 2: 18 of 20 returning birds, P < 0.0001; Season 2 vs. Season 3: 15 of 15 returning birds, P < 0.0001). Consistency in community membership also persisted across a two-year interval: 7 of the eight individuals that were in both the Season 1 and Season 3 network also returned to the same social community (Table 3; P = 0.003). Birds that were observed in two consecutive seasons appear to have acted as the cores for social communities—they had stronger connections with other members of their social communities compared to other birds in the network (Figure 3; ANOVA: Season 2, F\textsubscript{1,41} = 6.5, P = 0.015; Season 3, F\textsubscript{1,25} = 13.2, P = 0.001). Because these networks exclude birds that are banded for the first time during the observation period (i.e. birds likely in their first winter seasons), our analysis shows that birds continue to become more central in their communities even after three or more seasons.

Discussion

We discovered previously unsuspected levels of social complexity for a wintering migrant songbird. Golden-crowned sparrows exhibit a highly non-random social structure in which three main communities occur within a small area (size of study site ~ 7 ha) each year. Short-term flocks were dynamic in composition but consisted of different subsets of a larger stable social community of individuals. Strikingly, we found that birds that flocked together often during one season were more likely to flock together again in the following year, above and beyond what would be expected by the extent of overlap in their home ranges. To our knowledge, this is the first evidence of such social stability across years in a wintering population of migratory birds. Across-
year social stability between birds also leads to repeatable community structure of the population across years, even with large degrees of turnover in the population due to mortality and recruitment.

Our findings enrich our understanding of winter sociality of birds in the genus *Zonotrichia*, a particularly well-studied group with respect to social dynamics of non-breeding birds. One detailed longitudinal study (Piper & Wiley 1990) found that white-throated sparrows (*Zonotrichia albicollis*) tended to return to the same location across years, as we observed here, and that individuals also showed across year consistency in home range size. However, because the study did not explore the patterns of flock associations we do not yet know whether white-throated sparrows show the same patterns of network structure observed for golden-crowned sparrows. *Zonotrichia* sparrows, including golden-crowned sparrows, have also played a central role in the development of the theory of “badges of status” as a means of mediating social competition (Rohwer 1975; Rohwer & Ewald 1981; Watt 1986; Chaine et al. 2011, 2013). Much of this work assumed that these birds live in fluid flocks where individual recognition is not possible. Our findings suggest that the social context in which badges are used may be more nuanced than early theory assumed, a point we discuss more fully below.

Wintering golden-crowned sparrows show fission–fusion social dynamics that are similar to those found in some mammalian societies such as elephants (Wittmeyer et al. 2005), equids (Sundaresan et al. 2006), sea lions (Wolf et al. 2007), dolphins (Connor et al. 2000), and some primates (Smuts et al. 1987). As with these systems, sparrow populations consist of social communities that subdivide to form temporary foraging flocks. Similar social dynamics have recently been described in a few systems that are generally not

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**Table 2.** Results of network regression (MRQAP) analysis comparing the effects of (1) previous year’s association strength and (2) extent of home range overlap on the association strength between pairs of individuals in each of two seasons, seasons 2 and 3.

<table>
<thead>
<tr>
<th>Season</th>
<th>Previous Year Association</th>
<th>Home Range Overlap</th>
<th>Full Model</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
<td><em>P</em></td>
<td>Estimate</td>
</tr>
<tr>
<td>2</td>
<td>0.24</td>
<td>&lt;0.001</td>
<td>0.13</td>
</tr>
<tr>
<td>3</td>
<td>0.45</td>
<td>&lt;0.001</td>
<td>0.12</td>
</tr>
</tbody>
</table>

**Table 3.** Comparisons of community membership across years to determine the number of individuals that returned to the same versus different communities.

<table>
<thead>
<tr>
<th>Season 1 Membership</th>
<th>Season 2 Membership</th>
<th>Season 3 Membership</th>
</tr>
</thead>
<tbody>
<tr>
<td>Green</td>
<td>Yellow</td>
<td>Blue</td>
</tr>
<tr>
<td>Green</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Yellow</td>
<td>1</td>
<td>8</td>
</tr>
<tr>
<td>Blue</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>New*</td>
<td>4</td>
<td>9</td>
</tr>
<tr>
<td>Proportion New</td>
<td>0.33</td>
<td>0.53</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Season 1 Membership</th>
<th>Season 2 Membership</th>
<th>Season 3 Membership</th>
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<tr>
<td>Yellow</td>
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<tr>
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<td>0</td>
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<tr>
<td>White</td>
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<td>New*</td>
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<tr>
<td>Proportion New</td>
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<td>0.33</td>
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</table>

<table>
<thead>
<tr>
<th>Season 1 Membership</th>
<th>Season 2 Membership</th>
<th>Season 3 Membership</th>
</tr>
</thead>
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<td>Blue</td>
</tr>
<tr>
<td>Green</td>
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<td>0</td>
</tr>
<tr>
<td>Yellow</td>
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<td>2</td>
</tr>
<tr>
<td>Blue</td>
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</tr>
<tr>
<td>New*</td>
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<td>4</td>
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<tr>
<td>Proportion New</td>
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<td>0.67</td>
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</table>

Community colors follow Figure 1 for each community, we also note the number of individuals that did not return across seasons and the number of new immigrants that joined the community.

* New denotes birds that are included in the network for the first time in Season 2 (top table) or Season 3 (bottom two tables).
considered as highly social, such as sharks (Mourier et al. 2012). Our study adds to the growing evidence that fission-fusion dynamics are widespread across animal societies (Aureli et al. 2008).

The structure of social networks is strongly influenced by both social preference among individuals and potentially non-social factors such as the spatial segregation of individuals in the population (Pinter-Wollman et al. 2014). For example, if habitat is heterogeneous and some home ranges cluster together, then social communities could arise as an emergent property of foraging patterns (Ramos-Fernández et al. 2006) – i.e. communities could consist of individuals that simply prefer to forage in the same locations without any benefits from being social. Similarly, individuals may form temporary social groups with other individuals with which they share space, even when the benefits of sociality do not depend on the identity of the individuals. Alternatively, network structure could be driven by purely social factors – individuals may seek out certain flock mates (e.g. familiar individuals or kin) because they gain additional benefits. In many cases, both social and spatial factors are likely to contribute to social structure. The challenge is to develop methods to tease apart the relative contributions of social versus non-social processes that lead to social network structure (Aureli et al. 2008; Pinter-Wollman et al. 2014).

In this study, we used two complimentary approaches to demonstrate social factors are involved in structuring the network: (1) comparing empirical networks with null models that incorporate information on individual space use patterns (i.e. our Spatial Flock Model), and (2) using network regression (MRQAP) to test whether social factors (i.e. previous association) influence network structure independent of spatial patterns. The null model approach demonstrated that the empirical networks exhibit levels of community structure that cannot be explained by patterns of space use alone. The network regression tests showed that a large part of this unexplained variation in association patterns could be explained by previous social experience – birds prefer to flock with the same individuals across years.

While we detected a clear role of across-year social preference in social network structure in our sparrow population, it is important to note that social preferences could be even more important than our statistical tests indicate. This is because social factors could also influence the patterns of individual space use – birds may occupy a home range because they prefer to flock with other individuals that are found in that location. While the Spatial Flock Model captured the baseline level of community structure expected if flocks were composed of random sets of individuals that share home ranges, it did not account for the potential influence of social preference on home ranges. Development of more sophisticated models that incorporate changes in space use and social associations across finer temporal scales may help assess the influence of social preference on space use patterns. Statistical tests based on such spatiotemporal models may reveal that social preferences play an even bigger role in the social structure of wintering birds.

Across-year social preference could contribute to stability in social structure if returning individuals segregate into discrete groups based on past familiarity and form the cores of social communities in future years. Supporting this suggestion, we found that birds returned to the same social communities across years, and that birds observed in consecutive years had stronger ties within those communities. As a result, the overall social structure remained stable despite substantial turnover of individuals in the population due to mortality and recruitment: Comparing across three seasons, each community was composed of largely different individuals between Seasons 1 and 3 (64–86% new members; Table 3). In effect, social communities outlast the lives of their individual members. This type of turnover in community membership is known to promote the long-term stability human social networks (Palla et al. 2007), and it could explain the across-year stability of social communities in golden-crowned sparrows. Understanding the processes of formation and maintenance of social communities in golden-crowned sparrows will require more detailed data on behavioral and temporal aspects of flock associations and locations at finer time scales within seasons.

The existence of fission-fusion flock dynamics, stable social relationships and stable community structure in golden-crowned sparrows has important implications for sociality in non-breeding contexts. For example, forming stable groups with familiar individuals could provide a number of benefits through active food sharing (Carter & Wilkinson 2012) and improved coordination in anti-predator behavior (Croft et al. 2006; Micheletta et al. 2012). The partitioning of the popula-

**Figure 3** Birds that were seen in the previous year has stronger ties within their social communities than birds not previously seen in both (a) Season 2 and (b) Season 3. Shown are box-and-whiskers plots. Statistics are reported in the main text.
tion into small social communities can also facilitate cooperation via reciprocity (Trivers 1971; van Doorn & Taborsky 2011) and should favor the evolution of individual recognition as a means to identify flock-mates and prevent invasion of cheaters (Pagel & Dawkins 1997). While interactions between less familiar individuals are expected to involve conventional signals that mediate conflict [e.g. badges of status: (Rohwer 1975; Maynard Smith & Parker 1976)], individual recognition should help settle contests in small groups (Pagel & Dawkins 1997). We previously confirmed that badges of status influence the outcomes of competition between pairs of unfamiliar golden-crowned sparrows (Chaine et al. 2011, 2013). The existence of stable groups where individual recognition could be favored would suggest that badges may be used in interactions during community formation or between communities, and that individual recognition might be more important within communities. A thorough exploration of the function and evolution of status signals will require a shift in perspective to context-dependent signaling directly tied to the dynamics of social organization.

Acknowledgments — Special thanks to the UCSC Arboretum for use of an indoor office space for banding and other logistic support. We thank D. B. McDonald for consulting on the design of our observation protocol during the early part of this work, and especially with network theory training to D.S. We also thank many field assistants that conducted observations that informed our understanding of this system. Independent research by J. Garcia, D. Moldoff, A. Roth, K. Tjernell and C. Troll laid the groundwork for this study. We also thank W. Piper and two anonymous reviewers for feedback that improved this manuscript. This work was funded by a UCSC Special Research Grant to B.E.L. and NSF-DDIG (IBN-0309215) grant to A.S.C. and B.E.L. A.S.C. received support from ANR-JCJC NetSelect and is part of the LABEX-TULIP (ANR-10-LABX-41).

Authorship All authors contributed to research design. AS, JA, OJ, IML and BL collected data. DS conducted analyses and wrote the first draft of the manuscript, and DS, AS and BL contributed substantially to revisions.

References


Supporting Information

Supporting Information follows.
Supporting Information

Across-year social stability shapes network structure in wintering migrant sparrows

Daizaburo Shizuka, Alexis S. Chaine, Jennifer Anderson, Oscar Johnson,

Inger Marie Laursen & Bruce E. Lyon

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Supplemental Methods
Table S1: Comparisons between the distributions of modularity ($Q_{\text{max}}$) in the null models and empirical network.

<table>
<thead>
<tr>
<th>Season</th>
<th>$F$</th>
<th>$d$</th>
<th>$P$</th>
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<tr>
<td>Random Flock vs. Spatial Flock</td>
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<td>2</td>
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<tr>
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<td>49102</td>
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<td>&lt;0.001</td>
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<td></td>
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<tr>
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<tr>
<td>3</td>
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<tr>
<td>Random Flock vs. Empirical</td>
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The distributions for the empirical networks are generated by bootstrapping. $F$, Cohen’s $d$ and $P$-values are based on two-way ANOVA.
Appendix S1: Assessing the robustness of community assignments

Supplemental Methods

In our empirical networks, individuals are assigned to each community based on the modularity optimization method of Clauset et al. (2004). However, it is important to assess how robust these community assignments are to sampling error in our flock observations. Lusseau et al. (2008) outlined a general framework for assessing the confidence of community assignments using a bootstrapping technique. However, they did not provide details of how one would quantitatively assess the robustness of community assignments. Here, we expand on the bootstrapping approach to provide a quantitative measure of confidence in community assignments for a given network.

A major complication to assessing the confidence of community assignments through bootstrapping is that there is no clear way to determine which community in a ‘resampled network’ corresponds with which community in the empirical network. To circumvent this problem, we shift our focus from the community assignment at the individual level (i.e., ‘individual A belongs to community X’) to the dyadic level—i.e., how confident are we that two individuals that are assigned to the same community actually belong together? This measure of confidence in community assignment would be high if pairs of individuals that are assigned to the same community in empirical networks are often assigned to the same community when the observed data are resampled through bootstrapping.

We use the same boostrapping technique as described in the main text for generating a confidence interval for our empirical measure of modularity—for each year,
we resampled flocks with replacement up to the number of flocks observed to create a ‘resampled network’. We then used the same modularity optimization method used in the empirical network (Clauset et al. 2004) to assign individuals to social communities in this resampled network. Repeating this procedure 1,000 times, we can then generate a matrix of values $P_{ij}$, which is the proportion of resampled networks in which each pair of individuals are assigned to the same community, given that both individual were included in the sample. This matrix of $P_{ij}$ values can then be represented as a network in which edge widths represent the pairwise confidence in community assignments over all pairs of individuals (Figure S1).

We can use a network of $P_{ij}$ values to test our confidence in the empirical community assignments using the assortativity coefficient (Newman 2003). The assortativity coefficient measures the degree to which edges occur between nodes of similar type versus those of different types. Here, we use the community assignments in the empirical network as the node type. Thus, high levels of assortativity would indicate that the probability of being in the same community in bootstrap replicates is highest for pairs of nodes that were in the same community in the empirical network—evidence that our empirical community assignments were robust to sampling error. Because edges in our network are weighted, we based our measure confidence in community assignment on a weighted assortativity coefficient for undirected networks (Farine 2014). We denote the proportion of the total network edge weights ($P_{ij}$) that occur between nodes of empirical community $x$ (i.e., nodes of the same color in Figure S1) as $e_{xx}$, and $e_{xy}$ is the proportion of total network edge weights that connect nodes of types $x$ and $y$ (i.e., nodes of different color in Figure S1). The value $a_x = \sum_y e_{xy}$ represents the proportion of edges
that connect nodes of community $x$ to a different community, $y$, for all values of $y$ (i.e. all possible communities). Thus, if edge weights are distributed randomly among nodes of different communities, then the expected proportion of edge weights occurring between nodes within community $x$ is $(a_x/2)^2$. Here, $a_x$ is multiplied by $\frac{1}{2}$ because undirected edges will be counted twice (e.g., an edge that connects community $x$ to community $y$ also connects community $y$ to community $x$). Then, the degree to which community assignments in the empirical network and bootstrap networks are consistent can be calculated as

$$r_{\text{community}} = \frac{\sum_x e_{xx}^2 - \sum_x \left(\frac{a_x}{2}\right)^2}{1 - \sum_x \left(\frac{a_x}{2}\right)^2}$$

Thus, $r_{\text{community}}$ represents the degree to which the proportion of edge weights ($P_{ij}$) occurs between nodes assigned to the same empirical community, compared to the random expectation. This value is scaled such that $r_{\text{community}}$ equals 1 when the empirical community assignments are perfectly robust—i.e., when all bootstrap replicates result in the exact same community assignments as the empirical network. Conversely, $r_{\text{community}}$ equals 0 when community assignments in bootstrap replicates are completely random with respect to the original empirical communities. The value of $r_{\text{community}}$ can be negative if nodes assigned to different empirical communities are more often assigned to the same community in bootstrap replicates.
Supplemental Results

Our community assignments in the empirical networks were generally robust to sampling error. Figure S1 visualizes the proportion of bootstrap replicates in which two individuals were assigned to the same community, and Figure S2 summarizes the edge weights as box-and-whisker plots. Individuals that were assigned to the same community in empirical networks (i.e., nodes that have the same color in Figure S1) were often assigned to the same community in resampled networks, while individuals assigned to different empirical communities were rarely in the same community in resampled networks (Figure S2). Thus, each network had large $r_{\text{community}}$ values (Season 1 $r_{\text{community}} = 0.82$; Season 2 $r_{\text{community}} = 0.76$; Season 3 $r_{\text{community}} = 0.82$).
Figure S1: Aggregated bootstrap networks of golden-crowned sparrows in Seasons 1 through 3 (a-c). Nodes represent individuals and edges represent $P_{ij}$, the proportion of bootstrap replicates (‘resampled networks’) in which two nodes were assigned to the same community. Nodes are colored according to their original community assignments in the empirical networks (Figure 1 in main text). Nodes that belong to the same community in the empirical networks are also likely to be assigned to the same community in resampled networks, demonstrating the robustness of community assignments. We used a force directed algorithm to determine node placement (see Figure 1). Edges with values <0.05 (pairs that are in the same community in less than 5% of bootstrap replicates) have been trimmed for ease of display. These edges were included in our analyses.
Figure S2: Box-and-whisker plots showing the distributions of $P_{ij}$, proportion of bootstrap replicates in which pairs of nodes assigned to the same community in bootstrap replicate networks. (a-c) In each season (Season 1, 2 and 3 respectively), the gray plot represents $P_{ij}$ values for node pairs assigned to the same community in empirical networks, and white plot represents values for pairs assigned to different communities in empirical networks.
Appendix S2: Visualizing the spatial structure of social communities

Supplemental Methods

To visualize the spatial structure of social communities—e.g., whether or not communities occupy discrete areas within the study site—we estimated community home ranges and overlaid these on our aerial maps of the study site (Figure 1). We assigned each flock to a community based on which community had the most members represented in the flock. If equal numbers of individuals from multiple communities occurred in the flock, the flock was assigned to both communities. We then used the locations where the flock was first located as a data point. We used a kernel home range estimation method to visualize the home range (90% utilization density) and core areas (50% utilization density) of each social community.
REFERENCES CITED


