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RESEARCH ARTICLE

Social network analysis reveals context-dependent kin relationships in wild sulphur-crested cockatoos *Cacatua galerita*

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Abstract

- 1. A preference to associate with kin facilitates inclusive fitness benefits, and increased tolerance or cooperation between kin may be an added benefit of group living. Many species exhibit preferred associations with kin; however, it is often hard to disentangle active preferences from passive overlap, for example caused by limited dispersal or inheritance of social position.
- 2. Many parrots exhibit social systems consisting of pair-bonded individuals foraging in variably sized fission-fusion flocks within larger communal roosts of hundreds of individuals. Previous work has shown that, despite these fission-fusion dynamics, individuals can exhibit long-term preferred foraging associations outside their pair bonds. Yet the underlying drivers of these social preferences remain largely unknown.
- 3. In this study, we use a network approach to examine the influence of kinship on social associations and interactions in wild, communally roosting sulphurcrested cockatoos, *Cacatua galerita*. We recorded roost co-membership, social associations and interactions in 561 individually marked birds across three neighbouring roosts. We then collected genetic samples from 205 cockatoos, and conducted a relationship analysis to construct a kinship network. Finally, we tested correlations between kinship and four social networks: association, affiliative, low-intensity aggression and high-intensity aggression.
- 4. Our result showed that while roosting groups were clearly defined, they showed little genetic differentiation or kin structuring. Between roost movement was high, with juveniles, especially females, repeatedly moving between roosts. Both within roosting communities, and when visiting different roosts, individuals

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preferentially associated with kin. Supporting this, individuals were also more likely to allopreen kin. However, contrary to expectation, individuals preferred to direct aggression towards kin, with this effect only observed when individuals shared roost membership.

5. By measuring social networks within and between large roosting groups, we could remove potential effects of passive spatial overlap on kin structuring. Our study reveals that sulphur-crested cockatoos actively prefer to associate with kin, both within and between roosting groups. By examining this across different interaction types, we further demonstrate that sulphur-crested cockatoos exhibit behavioural and context-dependent interaction rules towards kin. Our results help reveal the drivers of social association in this species, while adding to the evidence for social complexity in parrots.

KEYWORDS

Cacatua galerita, kinship networks, parrots, population genetics, social network analysis, urban ecology

1 | INTRODUCTION

Living in groups can provide a range of benefits to all individual members (Brown & Brown, 1987). These benefits can be further enhanced by preferentially associating with specific others (Krause et al., 2002). For example, associations with individuals sharing characteristics such as age (Aplin et al., 2021), sex (reviewed by Ruckstuhl, 2007), body size (Croft et al., 2005; Pitcher et al., 1986) or rank (Smith et al., 2007) can reduce conflict (Dadda et al., 2005) and predation risk (Sorato et al., 2012). Moreover, associations with kin have been shown to provide fitness benefits that range from higher reproductive success (Silk et al., 2009) and inclusive fitness (Hamilton, 1964; Grafen, 1984; Levin & Grafen, 2019, though see Birch, 2017) to reduced stress levels (Gerlach et al., 2007) and increased longevity (Silk et al., 2010). Conversely, associating with kin can be disadvantageous if such associations increase the risk of inbreeding, or lead to competition between relatives (Evans et al., 2021). However, these costs can be partly mitigated via context-dependent social decision-making, for example avoiding kin during the breeding season while preferring them at other times (Evans et al., 2021). Yet while the costs and benefits of associating with kin have been extensively explored, the underlying processes leading to such patterns warrants further research.

Association with kin might reflect active choice, but alternatively might arise from passive mechanisms such as limited dispersal (Klauke et al., 2016) or an inheritance of social position (Ilany et al., 2021). There are two likely avenues to help disentangle active and passive drivers for social associations. The first is to examine this question in social systems exhibiting fission-fusion social dynamics, where larger groups or communities frequently split into subgroups of variable composition. In systems with such fluid group membership, sex, age and breeding status have all been shown to affect associations, with individuals sharing these characteristics being more associated

than expected by chance (Aplin et al., 2021; Chiyo et al., 2011; Zeus et al., 2018). When considering kinship however, the pattern is both more mixed and context-dependent. In species where individuals forms alliances, individuals jointly direction aggressions towards others tend to be more related than expected by chance (e.g. dolphins: Krützen et al., 2003; Parsons et al., 2003; monk parakeets: Dawson Pell et al., 2021). While a more general tendency to associate with kin has also been reported (e.g. Elephants Loxodonta africana: Chiyo et al., 2011; Geese Branta leucopsis: Kurvers et al., 2013; Giraffes Giraffa camelopardalis: Carter et al., 2013), other species show no such preference; for example, roosting patterns in big brown bats (Eptesicus fuscus, Metheny et al., 2008) are not affected by kinship. Perhaps most intriguingly, a recent comparative study investigating kinship, social networks and roost-switching behaviour in nine species of bats (Wilkinson et al., 2019), found that the preference to associate with kin was stronger in species with more roost-switching. This suggests a possible nonlinear relationship between social structure and kinship, where the benefits of associating with kin may not only be elevated in highly structured societies (e.g. cooperative breeding systems), but also be higher in social contexts where very few individuals are known to one another.

A second potential avenue is to examine variation in interaction decisions while controlling for their social opportunity. Here the assumption is that if individuals can recognise kin and are preferentially associating with them, they should also be more likely to direct affiliative behaviours towards relatives (e.g. grooming and food sharing) and be less likely to engage in aggressive interactions (reviewed by Hepper, 1986; Waldman, 1988). Supporting this, a meta-analysis of grooming in primates found a general positive correlation with kinship, although it also supported a larger role for direct reciprocity (Schino & Aureli, 2010). While the literature on social interactions is extensive, most studies have examined these questions in groups with high relatedness (e.g. in cooperative breeders, Madden

et al., 2012). There are relatively few examples of studies examining the influence of kinship on social decision-making in more open, fission-fusion species.

Here, we examined the influence of kinship on social associations and interactions in sulphur-crested (SC-) cockatoos, Cacatua galerita. Similar to many parrot species (Hardy, 1965; Power, 1966; Styche, 2000), SC-cockatoos live in large-scale fission-fusion societies, where stable night-roost aggregations of up to several hundred individuals (Lindenmayer et al., 1996) break-up during the day into smaller flocks that forage in a collective home range around the roost site. Foraging flocks have a fluid membership and are of variable size depending on activity, environment and season (Aplin et al., 2021; Noske et al., 1982). There is little evidence of any kin structuring within roosts in SC-cockatoos or any other parrot species (Wright & Wilkinson, 2001, but see Blanco et al., 2021). In addition to within roost dynamics, between-roost movements are common in SC-cockatoos (Aplin et al., 2021), although their predictors are largely unknown. The combination between withinand between-roost social dynamics confer a large choice of potential interaction partners, and previous work in this species has shown that SC-cockatoos form non-random social associations, with stable, long-term bonds beyond the pair (Aplin et al., 2021). However, little is known about the factors underlying association preferences or the predictors of social interactions such as allopreening or aggression.

Sulphur-crested cockatoos are urban persisters, and in such environments, populations are relatively habituated and often forage on the ground (Davis et al., 2017; Kirksey et al., 2018), giving a rare opportunity to conduct detailed behavioural observations. First, we habituated and marked 561 SC-cockatoos across five neighbouring roosting communities. We then collected genetic material for 393 individuals and performed an analysis of genetic relatedness for these five sites. Second, we collected detailed observations at three of these roost sites, and an intermediate regularly visited foraging site, as well as censusing roost co-membership. This resulted in 205 individuals where we had information on both kinship and social interactions. We then examined the correlation between kinship and sociality using social network analysis to compare kinship networks with social associations (foraging flock co-occurence) and three interaction networks—allopreening (an affiliative behaviour) and lowand high-intensity aggressive interactions. Finally, we performed a second analysis to examine behaviour at two scales: within and between roosting communities.

Previous work on this population of SC-cockatoos has revealed that individuals maintain long-term bonds outside of the pair, which presumably confer benefits to individuals (Aplin et al., 2021). Given the additional predicted benefits of interacting with kin, we predicted that these stronger bonds would often be with close kin. We further predicted that, if this represented an active decision (rather than, for example, delayed dispersal of offspring), SC-cockatoos would show a preference to associate with kin when visiting other roost groups, or when meeting kin from other roost groups at foraging sites. Finally, we predicted

that individuals would preferentially direct affiliative interactions towards kin and be less likely to direct aggressive interactions towards kin; we further predicted that this would hold when controlling for social opportunity. Our multi-step analytical approach removes the potential effect of spatial overlap and philopatry on any correlation between kinship and social associations, while allowing us to ask whether SC-cockatoos show behavioural or context-dependent social interaction rules towards kin that would infer both individual recognition and a degree of social complexity (Wilkinson et al., 2019).

2 | MATERIALS AND METHODS

2.1 | Study system

Sulphur-crested cockatoos are large (700-1200g), slow breeding, long-lived parrots. They are common across a large native range in Australia and Papua New Guinea, and populations appear to be increasing, including in human-modified areas (Davis et al., 2017). In the Sydney population, the majority of individuals are non-breeding, forming stable communal roosts of tens to hundreds of individuals, usually in parkland with remnant old growth trees, similar to previous descriptions at other sites (Lindenmayer et al., 1996). Tree hollows for nesting are a limited resource and breeding pairs (approximately 20%-40% of adults) defend nearby nest hollows year-round, but often roost with the rest of the group when not actively breeding (personal observation). Most individuals are site faithful, with a previous study on this population finding that birds used an average of two roost sites over a period of 5 years (Aplin et al., 2021). Within roosting communities, SCcockatoos exhibit fission-fusion dynamics during foraging, and individuals of different roosts associate when feeding on a variety of resources in the surrounding area (Aplin et al., 2021). In urban areas, resources include human derived and provisioned food (Kirksey et al., 2018; Klump et al., 2021), making birds relatively easy to habituate.

2.2 | Habituation, marking and morphometric data collection

Birds were habituated to human observers at suburban parks close to three focal roosts (Balmoral Beach BA, Clifton Gardens CG and Northbridge Golf Club NB), and at the nearest two neighbouring roosts (Sydney Botanic Gardens BG and Ivanhoe Park MA; Figure 1a) Over 2–3 weeks, individuals were attracted to the ground at the same time each day with small amounts of sunflower seed, a commercially available food commonly eaten by SC-cockatoos in urban and agricultural areas (Noske et al., 1982). The observer would remain as close as possible to the foraging flock until the birds began to forage on the ground around the observer and birds could be lightly touched on the back without eliciting an

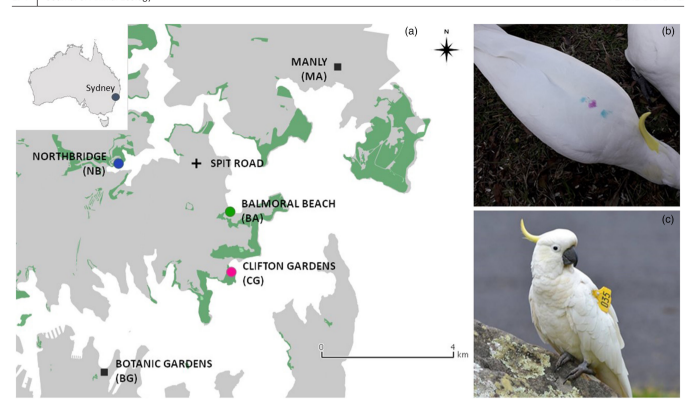


FIGURE 1 Study locations and examples of markings. (a) Map of sampling locations. Water surfaces are represented in white, hard surfaces (e.g. buildings, roads) in grey and vegetation (e.g. parks, bush-land) in green. Coloured circles represent focal roosts (green: BA, pink: CG, blue: NB) where social observations were conducted; grey squares are nearest neighbouring roosts (MA, BG) where birds were also marked and additional genetic samples were taken. The black cross represents an intermediate foraging site where birds from the three focal roosts mixed on a regular basis while being fed by locals, and social data was also collected. The map was constructed in QGIS (QGIS Development Team, 2009), using OpenStreetMap (OpenStreetMap contributors, 2017). (b) Example of marking with temporary dye, 'Teal-Pink-Teal', the marks remain visible for approximately 3 months, (c) example of a wing-tagged individual 035, named 'Shakespeare'.

adverse response. Once birds were habituated, they were temporarily and non-invasively marked using non-toxic fabric dye (Marabu Fashion Spray, MARABU GMBH), with each individual receiving a unique mark of one to three colours placed on the middle of the back (Figure 1b). This method resulted in 536 uniquely marked individuals across the five roosts. In addition, 17 birds present in our study area had already been caught, ringed and wingtagged as part of a previous study (Figure 1c, Aplin et al., 2021; Davis et al., 2017), and 8 birds were uniquely recognisable due to morphological variation (e.g. healed injuries). This resulted in 561 recognisable individuals, which we estimate to be >90% of the local population.

Individuals were opportunistically sexed and aged by eye colour (brown = juvenile <7 years; red = adult female; black = adult male). Finally, feather samples were collected without capture by plucking feathers from the back of habituated individuals. Once collected, feather samples (n = 1-10 feathers per individual) were stored in paper envelopes until ready for processing. The feather tip (1–2 mm; n = 351) was separated for genetic analysis (see below). This produced successful data for 244 samples; these were used to confirm morphometric sexing for adults, to assign sex for juveniles, and for relatedness analyses. All procedures were approved by the ACEC (ACEC Project No. 19/2107).

2.3 | Measuring social associations and interactions

We collected intensive social data at the three focal roost sites in northern Sydney and at one intermediate foraging site (Figure 1a) for 10 days in July (July 8–July 20) and 10 days in September (September 19–October 2) 2019. As for marking, individuals were attracted to the ground using small quantities of sunflower seeds scattered over a large area of approximately 385–500 m² to encourage ground foraging (e.g. of grass roots and shoots). Once the individuals were foraging on natural food sources, presence/absence scans (Altmann, 1974) were conducted every 10 min, where we recorded the identities of all individuals present within this study area in the park, both on the ground and in surrounding small trees.

Data was collected at the focal roosts over 3 h (July 2019) or 2.5 h (September 2019—time at each group adjusted to fit within daylight) each day. Time periods for presence scans at the intermediate foraging site were more opportunistic, depending on the feeding regime of the local resident (no additional feeding or interference was undertaken). Altogether, the final dataset consisted of 819 scans.

In addition to social association data, affiliative and aggressive interactions were recorded ad libitum between scans (Altmann, 1974).

The number for each type of interaction is presented in Table S2. Finally, we censused roost co-membership on three separate occasions (July 17th–21st, September 15–20, September 30–October 3, 2019) at the three focal roosts. The observer stood under the roost trees pre-dawn and attracted individuals down to the ground with sunflower seeds as soon as the birds began to become active, recording all marked individuals present. Roost counts were solely used to assign birds to a particular roost. Assignments were made if birds had been observed sleeping at one roost site at least twice (n=153 individuals for whom roost memberships could be classified). Across the three focal roosts, marking location corresponded to roost membership for 125 of 153 birds (Pearson's Chi-squared test, $\chi^2=178.67$, df=4, p<0.001). Marking location was, therefore, used as proxy for roost membership for birds where roost was not known.

2.4 | Genetic analysis

2.4.1 | DNA analyses

DNA was processed by Diversity Arrays Technology (DArT), using their proprietary restriction-enzyme-based genome complexity reduction platform called DArTseq (Cruz et al., 2013; Kilian et al., 2012). Four combinations of restriction enzymes were tested and the PstI-SphI combination was selected. After digestion, DNA was processed following Kilian et al. (2012), using two different adaptors corresponding to two different restriction enzyme overhangs. Both adaptors contained an Illumina flow cell attachment sequence, and the PstI-compatible adapter also contained a sequencing primer sequence and a barcode region. A PCR was performed on the resultant libraries as follows: initial denaturation at 94°C for 1 min, then 30 cycles of 94°C for 20 s, 58°C for 30 s and 72°C for 45 s, and a final extension step at 72°C for 7 min. After PCR, equimolar amounts of the libraries underwent a c-Bot (Illumina) bridge PCR followed by single end sequencing for 77 cycles on an Illumina Hiseq2500. Additional details on this platform can be found in Kilian et al. (2012) and Cruz et al. (2013).

The resultant sequence data were processed using the proprietary DArT Pty Ltd analytical pipelines. Poor quality sequences were removed, and stringent selection criteria were applied to the barcode region to de-multiplex the sequence reads (minimum barcode Phred score was set at 30, while the minimum whole-read Phred score was set at 10). Sequences were then truncated to 69 bp and clustered with a Hamming distance threshold of 3. Low quality bases from singleton tags were corrected based on collapsed tags. These sequences were used for SNP calling using the proprietary DArTsoft14 pipeline based on DArT PL's C++ algorithm. True orthologous SNP variants were discriminated from paralogous sequences by assessing various parameters, including sequence depth, allele count and call rate. We calculated SNP error rates before and after filtering – that is, the number of SNP mismatches between replicate pairs over the total number of SNPs that were not missing in both

replicates. Analysis was based on the 13 replicate samples (Table S3) using R functions from Mastretta-Yanes et al. (2015).

2.4.2 | SNP filtering

The DArTseq SNPs were filtered based on data quality, missing data and linkage using the R package dartR 1.0.5 (Gruber et al., 2018). The initial dataset comprised 58,740 SNPs. Loci that were genotyped in >80% of individuals were removed (i.e. call rate). DArT runs 30% of the samples in replicate in independent libraries and sequencing runs. Loci were removed that were not 100% reproducible, based on the consistency of the locus measured across these technical replicates. SNPs with a read depth < 10 and a minor allele frequency < 0.01 were removed. Putatively sex-linked markers based on SNPs that were mostly homozygous in males and mostly heterozygous in females, and/or SNPs that aligned with the Z-chromosome of the Finch genome were removed. Only one SNP was retained per DArTseq marker to remove potentially-linked SNPs. After implementing these filters, the dataset comprised 12,447 SNPs.

Initially, we did not filter for Hardy Weinberg Equilibrium (HWE) as the dataset likely contains numerous family groups, violating the HWE assumption of random mating. However, as a conservative approach, we subsequently re-analysed a dataset with HWE filtering implemented to check for consistency. We performed the HWE test using the R package pegas 0.13 (Paradis, 2010), implementing 1000 Monte Carlo replicates. Loci with a *p*-value <0.05 based on the exact *p*-test were removed (removing a further 3507 SNPs).

Finally, before further downstream analysis, one of each of the replicate samples (with the least amount of missing data) were removed from the dataset. We then excluded one potential hybrid individual and one individual with >60% missing data; all other individuals had <15% missing data. This resulted in 393 individuals for the relatedness analysis, of which 205 belonged to the 5 roosts considered in our study.

2.4.3 | Relatedness analysis

We compared eight relatedness estimators to determine which one was the most appropriate for the SC-cockatoo SNP dataset (Wang, 2011). This included seven estimators implemented in the COANCESTRY software package (Wang, 2011), and a maximum likelihood kinship coefficient, calculated using the R package SNPRelate 1.14 (Zheng et al., 2012). To assess the suitability of these relatedness estimators, we simulated 100 parent-offspring pairs, 100 full-sibling pairs, 100 half-sibling pairs, 100 cousin pairs and 100 unrelated pairs and assessed the correlation between the relatedness estimates and true relatedness values. These simulations were performed in COANCESTRY using the empirical SNP allele frequencies and missing data. We compared the estimates using base R functions and the R package related 1.0 (Pew et al., 2015). After selecting the most reliable relatedness estimator based on the simulation

data, we performed a relatedness analysis on the empirical dataset. The reliability of the estimator was verified by assessing how well it approximated two known parent-offspring relationships included in the dataset.

2.5 | Social network analyses

We considered four social networks: social associations, low- and high-intensity aggression networks and allopreening. All social analyses were done using RStudio (version 1.4.1103; R version 4.0.4, R Core Team, 2021).

2.5.1 | Social associations

To create social networks for foraging associations, we created a group by individual matrix, where individuals were assigned to the same group if observed in the same presence scan. Given that estimates for rarely observed individuals may be inaccurate (Farine & Whitehead, 2015), we thresholded our data following the example of Aplin et al. (2013). In our case, removing individuals with fewer than 22 observations meant there was no significant relationship between number of observations and the unweighted degree. We then used the simple ratio index (SRI; Whitehead, 2008) to create an undirected network for each time period (July and September 2019) where edges between individuals were scaled between 0 (never observed in the same group) and 1 (always observed together). The association matrices for each period were highly correlated (Mantel test. n = 72, r = 0.63, p < 0.001). Therefore, association data for both study periods were pooled together for further analysis, resulting in a social network of 254 individuals. We tested for evidence of nonrandom preferred associations by examining whether the coefficient of variation (CV) of the observed SRI were significantly higher than permuted values. Permutations were conducted in the R package ASNIPE (Farine, 2013), permuting individuals in close spatio-temporal proximity, keeping rows and columns constant.

2.5.2 | Aggression networks

To create aggression networks, we first *post hoc* split behavioural sequences of aggressive interactions into two distinct types of behaviours that clearly differed in risk: low intensity (no body contact, hereafter LI-) and high intensity (body contact and chases, hereafter HI-). One might expect different underlying decision rules to these interactions, especially in interactions with kin, as only high-intensity interactions involve a risk of injury. For LI-aggressions this resulted in $N_{\rm int}=1457$ interactions, and for HI-aggressions, this gave $N_{\rm int}=91$ interactions (Table S2). For each type of aggression, we then calculated an undirected interaction matrix that additionally controlled for social opportunities to interact. For the LI-aggressions, edge weights were obtained by calculating for each dyad the number of

LI-aggressions divided by the total number of scans in which individuals were associated. For HI-aggressions, we calculated the proportion of HI-aggressions among all agonistic interactions between a given dyad, therefore controlling for the overall tendency of two individuals to engage in aggressive interactions.

2.5.3 | Allopreening networks

Allopreening events were recorded *ad libitum* over the observation period (n = 26). Within an allopreening event, preenings were mostly reciprocated. They also occurred in trees, making it challenging to record the initiator of the event. To create affiliative networks, we therefore constructed undirected networks where edge weights were obtained by calculating for each dyad the number of preening interactions divided by the sum of all preening events for both individuals. As observations of allopreening events were relatively rare (Table S2), we did not threshold the network.

2.6 | Statistical analysis of kinship and social networks

2.6.1 | Genetic differentiation between roosting communities

To examine whether there was any genetic differentiation between roosts, we performed two analyses: the first included five roosting communities, located within a 10 km radius (BA, CG, NB, BG, MA—Figure 1). The second focused on the three focal roosting communities (BA, CG, NB—Figure 1) located within a 4-km radius. For each analysis, we used mantel tests in the R package VEGAN (Oksanen et al., 2013), to correlate relatedness matrices with a spatial matrix of roost membership. This was performed using roost co-membership assigned for individuals at the three focal roosts by roost counts.

2.6.2 | Influence of kinship on social interactions

Level 1: study population

To test whether associations, LI- and HI-aggressive interactions and/ or affiliative interactions were influenced by relatedness, we compared kinship with these measures in separate models using multiple regression quadratic assignment procedures (MRQAPs) via the ASNIPE package (Farine, 2013), where the significance was ascertained using permutations (Farine & Carter, 2022). We additionally included roost co-membership and sex- and age-similarity as covariates in all models. In all MRQAP analyses, we used a continuous relatedness matrix. However, because we considered it unlikely for individuals to recognise second- or third-degree relatives, we also re-ran the same analysis on a binary matrix, specifying whether or not a given dyad were first degree relatives, that is, that their relatedness was above or equal to 0.36 (Figure S4; Tables S4-S7).

Level 2: Within vs between roost interactions

To test whether the correlation between relatedness and our social measures was mostly driven by within-roost or between-roost interactions, we conducted a second analysis where we (i) subset all edges to within-roost only and restricted randomisations to within the same roost and (ii) subset all edges to those occurring between individuals of different roost membership. Due to an increased likelihood of type II errors with this reduced dataset, we did not do this analysis for HI-aggression or allopreening. For all analyses, an estimate of type II errors (calculated following Foroughirad et al., 2019) are presented in Section S1.

3 | RESULTS

3.1 | Social organisation

In northern Sydney, 561 birds were individually marked or recognisable, and genetic material was collected from 205 birds. We sampled 164 birds from the three focal roost sites (Figure 1a; BA: n = 68; CG: n = 55; NB: n = 41), comprising 63 females and 72 males, of which 83 were adults, 45 juveniles and 37 birds of unknown age (Figure 2). At two additional roost sites, we sampled 41 birds (BG: n = 17; MA: n = 24), comprising 12 females and 23 males; 18 adults, 17 juveniles and 6 birds of unknown age. Of the eight relatedness estimators compared, the SNPRelate maximum likelihood kinship coefficient was most appropriate for this SNP dataset based on simulated dyads (Figure S1), having the highest correlation with the expected relatedness values (cor = 0.999; p < 0.001).

Overall, there was a weak tendency for individuals to be more related within than between roosting sites, but this was only present when considering all 5 roosts (focal roosts: Mantel Test, n = 164,

r = -0.003, p = 0.60; all five roosts: Mantel Test, n = 205, r = 0.011, p = 0.049). The higher relatedness within – compared with between - roosts was not solely due to delayed dispersal in juveniles; within each roost site, the number of adult-adult close-kin dyads was higher than expected within each population under random mixing (t-test, t = 2.28, df = 82, p = 0.025). In addition, juveniles tended to be observed at more roosting sites than adults (Figure S2A; females estimate $\pm 95\%$ confidence interval: ES = 0.553 [0.063-1.04], p = 0.02, males: ES = 0.739 [0.256-1.222], p<0.001; p-values calculated using Tukey adjustments for multiple testing). Juvenile females, but not juvenile males, tended to be less faithful to a specific site than adults of the respective sex, where faithfulness was defined as the proportion of observations at an individuals assigned roost (Figure S2B; females: ES = -0.107 [-0.216-0.002], p = 0.06, males:ES = -0.058 [-0.151-0.035], p = 0.37, p-values calculated using Tukey adjustments for multiple testing).

Analysis of association patterns revealed strong evidence for non-random associations (Median CV random = 1.26, CV obs = 1.25, p = 0.003), suggesting that individuals had preferred and non-preferred associates. While the social association network was clearly clustered by roosting community (Figure 2a), individuals also exhibited extensive foraging associations with individuals from different roost groups (Figure 2b).

3.2 Relatedness and social networks

When the entire study population was considered, kinship positively predicted social association strength (Figure 2a,b, ES = 0.097, p < 0.001). As expected, associations were also stronger between members of the same roost and with individuals of similar age (Figure 2, roost co-membership: ES = 0.097,

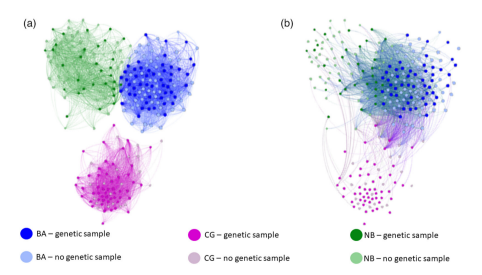


FIGURE 2 Association network of sulphur-crested cockatoos across three roosting locations, with (a) representing associations between individuals of the same roost and (b) representing only associations between individuals of different roosts. Networks are thresholded to only include individuals with at least 22 observations. Nodes are coloured by roost (blue: BA, pink: CG, green: NB), and the shade of each colour illustrates whether a genetic sample has been collected (dark shades) or not (lighter shades). Edges between nodes represent association strength and are thresholded to above 0.15 for visual clarity.

p<0.001, age: ES = 0.006, p = 0.012), but not of the opposite sex (sex: ES = 0.002, p = 0.18). The results were similar when considering only within-roost edges (kinship: ES = 0.11, p = 0.005; sex: ES = 0.006, p = 0.24; age: ES = 0.018, p<0.001; Figures 2a and 3a). When sub-setting the dataset to edges between individuals of different roost locations (Figure 2b), the effect of kinship on association strength was still present (ES = 0.086, p = 0.001), but there with no effect of sex or age (sex: ES = -0.001, p = 0.58; age: ES = 0.0003, p = 0.87). Finally, results were similar whether kinship was considered as a continuous or binary measure (close-kin or not; Table S4).

3.3 | Relatedness and aggressive interactions

Even after accounting for social opportunity, kinship was positively correlated with LI-aggression (ES = 0.41, p = 0.020, Figure 3b). LIaggressions ($N_{int} = 1457$, Table S2) also occurred more frequently between individuals belonging to the same roost than with individuals visiting from different roosts (roost co-membership: ES = 0.17, p < 0.001). There was also a trend for age similarity to predict LIaggressions (ES = 0.020, p = 0.09), but no effect of sex (ES = 0.01, p = 0.23). When considering only within-roost edges, the results were similar (kinship: ES = 0.78, p = 0.03; age: ES = 0.063, p = 0.08; sex: ES = 0.038, p = 0.31), as were the results when only considering close-kin dyads (Table S5). Between-roost LI-aggressions, however, were not influenced by kinship, age or sex (kinship: ES = -0.025, p = 0.41; age: ES = 0.002, p = 0.72; sex: ES = 0.004, p = 0.41), although there was a tendency to avoid directing LI-aggressions towards close-kin of different roost-membership (ES = -0.010. p = 0.08, Table S5).

Similar to LI-aggressions, HI-aggressive interactions ($N_{\rm int}=91$, Table S2) occurred more frequently between individuals belonging to the same roost, rather than between residents and individuals visiting from different roosts (roost co-membership: ES = 0.01, p<0.001). Kinship (ES = 0.055, p = 0.03) also positively predicted HI-aggression (Figure 3c), but there was no clear effect of age or sex similarity (age: ES = 0.003, p = 0.17; sex: ES = 0.003, p = 0.09). Due to low sample size, we did not examine the interaction between kinship and roost membership on HI-aggressions.

3.4 | Relatedness and allopreening interactions

Observations of allopreening events were scarce, with 26 allopreening interactions involving 41 individuals (Table S2). Yet even so, individuals were observed engaging in allopreening events with multiple partners. Allopreening interactions were more likely to be observed between stronger social associates (ES = 0.191, p < 0.001, Figure 3), related individuals (ES = 0.275; p = 0.008), individuals of different roost membership (ES = -0.015, p = 0.02) and tended to be more frequent between members of the opposite sex (ES = 0.012, p = 0.08).

4 | DISCUSSION

By combining relatedness data with extensive observations of social networks and interactions at multiple levels, our study gained an unprecedented insight into effects of kinship on the social lives of a wild parrot species. First, we collected genetic material and built kinship networks for five neighbouring roosting communities in a 25 km² urban and suburban area of Sydney. Our results showed a very weak genetic differentiation between these roost sites, suggesting extensive gene flow. This was supported by our repeated observations of between-roost movements, especially for juvenile females. Second, we used social network analysis to show that despite well-mixed social networks and a low degree of kin structuring, individuals preferentially associated with close kin. Supporting this, individuals were also more likely to allopreen kin, an important affiliative behaviour in parrots (Morales Picard et al., 2020). Contrary to expectations, aggressive interactions were also more likely to occur between kin within roost sites; intriguingly this was absent from between-roost interactions. Finally, we showed that this preference to associate with kin could be observed both within roost sites, and in encounters with individuals from other roosting sites.

A previous study in this system has shown that individuals maintain long-term preferred social associations (Aplin et al., 2021). Our work extends those findings to reveal one driver of these association choices, with kinship predicting social associations within roost sites, suggesting an active preference to associate with kin. Perhaps even more interestingly, we found that kinship was also positively correlated with association strength between individuals of different roost groups, suggesting that individuals continued to recognise and prefer kin even after dispersal. One benefit of such kin-based between-roost associations may be facilitated emigration into, or interactions with, groups containing closely related individuals (Van Hoof, 2000). For example, in cooperatively breeding brown jays Cyanocorax morio, males tend to disperse into groups where male relatives are already established (Williams & Rabenold, 2005). In the case of the brown jay, the formation of alliances between related males may facilitate the entrance into the new social group. A similar behaviour was also described for invasive monk parakeets, where male siblings exhibited parallel dispersal, then often cooperated when building a new nest and breed (Dawson Pell et al., 2021). Cooperative breeding or the formation of alliances has not been observed in SC-cockatoos, yet the observed preening interactions both between kin and individuals of different roosts is an intriguing indicator of the potential importance of maintaining relationships across groups.

In birds, preening is often described as having an essential role in maintaining pair bonds, such as mate bonding and partner retention across breeding seasons (Gill, 2012), or parental cooperation (Kenny et al., 2017). While our study does indeed identify allopreening between members of different sexes, it was not restricted to putative mates. Rather, individuals were observed engaging in preening events with multiple partners, and these interactions were positively with both kinship and social association strength. Allopreening

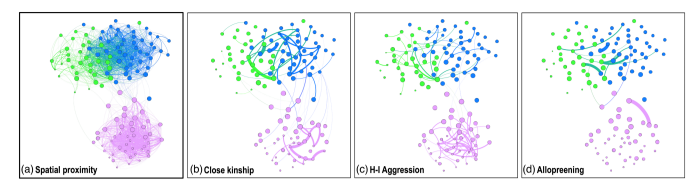


FIGURE 3 Social networks across three roosting communities. (a) Spatial association network of cockatoos where edge width represents association strength. Edges are thresholded to above 0.15 for visual clarity. (b) Association network subset to show edges between individuals with close kinship. (c) Association network subset to show edges between individuals that engaged in HI aggressions. (d) Association network subset to show edges between individuals that exchanged in allopreening. In all networks, node size represents average association strength, nodes are coloured by roost (blue: BA, pink: CG, green: NB) and networks are thresholded to only include individuals with at least 22 observations.

is generally understudied in wild parrots (but see Diamond & Bond, 1999; McCulloch, 1981; Taylor, 2002). In captivity, allopreening patterns of nine parrot species were positively correlated to the time a dyad spent in close proximity (Morales Picard et al., 2020). Furthermore, a preference to preen related, rather than unrelated individuals, has been described in juvenile birds (e.g. corvids: Fraser & Bugnyar, 2010; e.g. captive parrots: Garnetzke-Stollmann & Franck, 1991), where it is hypothesised to help build bonds with important social partners. The number of preening events in our dataset was low, and conclusions need to be drawn with care. Yet our results suggest that SC-cockatoos may gain some direct or indirect benefits from building and maintaining strong social bonds with kin, even after dispersal. Future work should seek to quantify what benefit these between-roost kin relationships give to individuals.

Somewhat surprisingly, we found that SC-cockatoos directed aggression towards individuals within rather than outside their roost. Furthermore, after social associations were controlled for, individuals also preferred to aggress kin with which they shared roost co-membership. A preference to direct aggression towards familiar individuals could potentially reflect strategies for improving dominance (Hobson et al., 2014, 2021), especially since 93% of all observed aggressive interactions involved only displacements that bear no direct risks of injuries for the individuals involved. However, the increased aggression between kin was in complete contrast to our prediction of higher tolerance. To note, all juveniles in our dataset were already independent of adults (2 to 7 years of age). One potential explanation therefore might come from the aggressive eviction hypothesis (Marsh, 1988; Wahlström, 1994), which states that increased aggression can act to drive emigration. For example, in western bluebirds Sialia mexicana, dispersal and dispersal distance are influenced by aggressions from kin (Aguillon & Duckworth, 2015). Although this hypothesis was formulated for natal emigration, it seems reasonable to expect increased intragroup aggression to also drive secondary dispersals, and even shortterm movements between social groups. Unfortunately, we did not have a large enough sample size to classify and distinguish between

aggressive interactions between offspring-parents and between siblings. However this hypothesis would be consistent with our observation that this preference to aggress kin is either lost, or perhaps reversed, when individuals interact with those of different roost membership.

Overall, the social system of SC-cockatoos allowed us to disentangle association preferences from spatial overlap, with well-mixed social networks within clearly-defined roosting groups coupled with regular associations with individuals of different roosts. We found that, despite exhibiting low overall kin structuring, individuals prefer to associate and allopreen with their kin both within and between roosting groups. This, therefore, suggests that (1) individuals are able to recognise kin, including siblings, and actively prefer them and (2) that kin-recognition is retained over a considerable period of time, including after dispersal. In passerine birds, cross-fostering experiments have demonstrated that kin recognition is often learned in the nest (Sharp et al., 2005). Yet the typical clutch size for SCcockatoos is one or two chicks per year, and juveniles appear to be largely independent at 1 year of age, suggesting that kin recognition must also occur across broods. In zebra finches between-brood recognition is achieved through olfactory cues (Krause et al., 2012). However, given the evidence for extensive vocal learning in parrots (Wright & Dahlin, 2018), and the existence of parent-facilitated vocal signatures in parrotlets Forpus passerinus (Berg et al., 2012, 2013), it seems most likely that kin recognition in SC-cockatoos involves vocal learning.

Whatever the mechanism of kin recognition, preferred associations with kin are often thought to benefit individuals in a number of ways, including through inclusive fitness, increased tolerance and cooperation. What the benefits are in this case are not obvious – kin do not cooperate to breed, and rather than exhibiting increased tolerance, individuals actually appear to be more likely to aggress kin, including with high-intensity aggressions that can carry risk of injury. While we were unable to ascertain the drivers, our study did reveal that interactions with related individuals are context-dependent, suggesting the existence of interaction rules that vary with the

behavioural context and on roost membership. More investigation of these dependencies may therefore be a fruitful avenue to identify the benefits of kin relationships in this system. Perhaps most intriguingly, our results suggest that roost membership is a key factor, with SC-cockatoos exhibiting preferred associations with individuals of other roosting groups that are often kin based, and appearing to maintain these relationships with affiliative behaviours such as allopreening. How individuals benefit from these extended networks remains to be further explored.

AUTHOR CONTRIBUTIONS

Julia Penndorf, Lucy M. Aplin, John M. Martin conceived the ideas and designed the study; Julia Penndorf, Lucy M. Aplin, Barbara C. Klump and John M. Martin collected social and morphometric data and collected feather samples. Kyle M. Ewart collated and analysed genetic data, and Julia Penndorf and Lucy M. Aplin performed analysis on sociality and kinship. All authors contributed to writing the manuscript and gave final approval for publication.

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CONFLICT OF INTEREST

The authors declare that no competing interests exist.

DATA AVAILABILITY STATEMENT

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182 Journal of Animal Ecology PENNDORF ET AL.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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