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Ageing in a collective: the impact of ageing individuals on social network structure

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Ageing affects many phenotypic traits, but its consequences for social behaviour have only recently become apparent. Social networks emerge from associations between individuals. The changes in sociality that occur as individuals get older are thus likely to impact network structure, yet this remains unstudied. Here we use empirical data from free-ranging rhesus macaques and an agent-based model to test how age-based changes in social behaviour feed up to influence: (i) an individual's level of indirect connectedness in their network and (ii) overall patterns of network structure. Our empirical analyses revealed that female macaques became less indirectly connected as they aged for some, but not for all network measures examined. This suggests that indirect connectivity is affected by ageing, and that ageing animals can remain well integrated in some social contexts. Surprisingly, we did not find evidence for a relationship between age distribution and the structure of female macaque networks. We used an agent-based model to gain further understanding of the link between age-based differences in sociality and global network structure, and under which circumstances global effects may be detectable. Overall, our results suggest a potentially important and underappreciated role of age in the structure and function of animal collectives, which warrants further investigation.

This article is part of a discussion meeting issue 'Collective behaviour through time'.

1. Introduction

The costs and benefits of an animal's behaviour toward members of their collective can shape lifespans, life-history evolution and the pace of senescence. For example, the support that cooperative breeding species receive from non-breeding helpers may lead to longer lifespans than those of related solitary species [1]. The transfer of enhanced ecological knowledge to close relatives during collective foraging may have contributed to the evolution of reproductive

senescence and prolonged post-reproductive lifespan (i.e. menopause) in female killer whales [2–4]. The formation of strong relationships between group mates has also been linked to enhanced individual longevity in a range of group-living taxa [5]. It is therefore clear that living in collectives can alter the adult ageing process. However, an important question is whether ageing, in turn, influences the behaviour of adults in collectives, and ultimately the structure of collectives themselves.

Growing evidence suggests that older adults differ in their social behaviours and social relationships from young adults [6–11]. One pattern that seems to be emerging across taxa is that older adults interact with fewer individuals than do younger adults, concentrating social relationships on close associates and kin [6,12–15]. Given that social networks are an emergent feature of association rules between individuals [16], shifts in patterns of social behaviour with age might not only affect who ageing individuals associate with directly (i.e. their direct connectedness), but could also affect higher-order network structure. Age-based changes in social behaviour could scale up to alter an individual's connections to the partners of their social partners (their indirect connectedness [17]) as well as the overall structure (topology) of the social network, both of which can have consequences for disease transmission [18–23], information transfer [24–26], the cohesive movement of groups [27,28] and many other eco-evolutionary dynamics [29,30]. Yet little attention has been given to understanding the impact of social ageing for the polyadic social world or the structure of the collective.

Understanding how ageing shapes an individual's indirect connectedness may be particularly relevant as such connections are tightly linked to processes that can directly influence fitness, including those described above. Declines in indirect connectedness with age may help limit exposure to disease [20–22], which might be beneficial in aged animals experiencing immunosenescence, but simultaneously could inhibit the transfer of important socio-ecological information [26,31], which could exacerbate pre-existing fitness declines in old age. In some cases, indirect connectivity may be an even more important predictor of fitness than direct connections dyadic associations [17,32]. Therefore, changes in indirect connections may be a particularly important component of the social ageing phenotype to investigate. Recent work has offered some glimpses into how measures of indirect connectedness can differ between young and old adults. In marmots and Barbary macaques, older adults have measures of indirect connectedness that suggest they have partners who are themselves not well connected [9,33] (but this is not the case in rhesus macaques, see [34,35]). Older marmots and rhesus macaques are less effective at reaching disparate nodes in the network compared to younger adults [34–36]. By contrast, older adults are more strongly embedded in cliques or clusters in their networks compared to younger adults in marmots [9], but not in either Barbary or rhesus macaques [33,35]. However, most research to date has compared differences in measures of indirect connectedness among adults of different age classes (e.g. old versus young), but has lacked the longitudinal data required to quantify how the social positioning of individuals changes across their lifetimes [9]. Such longitudinal analyses have the potential to reveal important patterns that might otherwise be masked by differences between individuals or cohorts [37].

Tracking within-individual changes in measures of indirect connectedness is essential for more firmly placing changes in sociality across the lifespan within the larger ageing phenotype and therefore understanding the causes and consequences of these patterns of social ageing [38].

Populations composed of a greater proportion of older (or younger) adults may also be structured in meaningfully different ways, and this could affect important processes such as communication and cooperation. For example, the loss of old individuals through age-related disease or trophy hunting can disrupt intergenerational flow of accumulated social and ecological knowledge, impeding collective movement and the ability to locate critical resources [4,39–42]. The age structure of a group can also regulate the behaviour of younger individuals [43,44], influencing aggression rates and social cohesion [45]. The impact of diminished cohorts of younger individuals on overall network structure is less well understood, but likely to have repercussions for network connectivity and cohesiveness given that younger adults are more socially active in many populations [8,10,12–14]. For example, the simulated removal of juvenile killer whales led to networks that were more fragmented than when random individuals were removed, suggesting an important role of young individuals in maintaining network cohesion [46]. Despite the established ecological and evolutionary importance of network structure [29,30], the underlying drivers of variation in network structure remain understudied [47–50]. Ageing, as an important process underlying patterns of individual-level variation in sociality, might therefore provide a window into how simple processes can generate complex network structures [51,52].

Using both empirical data and a theoretical model, we explore how social ageing of individuals relates to measures of indirect connectedness and overall network structure in a group-living primate, the rhesus macaque (*Macaca mulatta*), which is an emerging model in social ageing research [38,53]. As female rhesus macaques grow older, they show clear changes in their patterns of direct connectedness: they reduce the size of their social networks and focus their social effort on a few important partners, including close kin [6]. Despite this, females do not reduce the rate at which they engage in social interactions as they age, indicating that although their networks get smaller, older females continue to invest the same amount of time into fewer relationships [6].

Given these previously established changes in direct connections, here, we set out to test if age-based changes in social behaviour relate to measures of an individual's level of indirect connectedness [17,54], and if they scale up to influence network structure as a whole. Individual-level social network metrics, including measures of direct and indirect connectedness, relate to underlying, putatively simple, social differences or processes, such as individual-level variation in general sociability or reassociation tendency [51]. For example, an individual's general tendency to be sociable can be intuitively quantified using a direct network metric (e.g. strength, or the sum of the weights of an individual's ties to their partners), or using an indirect metric (e.g. weighted eigenvector centrality, which measures how well connected an individual is to its partners and how well connected those partners are to others [51]). Drawing on this idea that underlying processes can predict an individual's position in the network, and on the age-based changes in direct connections that we have previously documented in rhesus macaque females [6], we made

Table 1. Predictions for how indirect measures of connectedness are expected to change with age in female rhesus macaques from Cayo Santiago and how global network metrics are expected to change with increasing proportions of old individuals in the population.

network metric	definition	prediction	rationale
indirect network metrics			
eigenvector centrality	Measures how well connected an individual is in the network to individuals who are themselves well connected.	remains stable with increasing age	Females retain strength of connections to partners as they age.
betweenness	Calculates the number of shortest paths between others in a network that passes through an individual. Individuals with low betweenness have low capacity for linking discrete clusters in the network.	decreases with increasing age	Females increase their probability of interacting with strong, stable partners and kin. Therefore, females increase their tendency to reassociate with existing partners and have higher within-group association tendencies as they age.
closeness	Measures the inverse distance to all other individuals in the network. The lower an individual's closeness score, the more difficult it is for them to reach others or to be reached.	decreases with increasing age	
clustering	Measures cliquishness or sub-grouping. A high clustering coefficient indicates that an individual's partners are highly connected to each other.	increases with increasing age	
global network metrics			
mean degree	The average number of ties that each individual has to others in the network.	decreases with increasing proportion of old individuals	Older females have fewer social partners.
diameter	Measures the overall connectedness of the network. Networks with a larger diameter are less connected.	increases with increasing proportion of old individuals	
transitivity	The degree of clustering in the network.	increases with increasing proportion of old individuals	Older females cluster more with kin.

predictions for four common measures of indirect connectedness: eigenvector centrality, betweenness, closeness and clustering coefficient (table 1). Adult female rhesus macaques maintain their strength of grooming ties to others as they age, despite reducing their number of partners [6]. We therefore predicted that weighted eigenvector centrality (a measure of overall connectedness in the network) would also remain stable with age, as by retaining strong connections to some of their partners, females therefore (indirectly) retain connections to the partners of their partners. We also previously found that as female rhesus macaques age, they increase their likelihood of interacting with certain partners, especially their kin [6]. That is, ageing females increase both their tendency to reassociate with others and their tendency to interact with someone from their own (kin-based) sub-group. Individuals with higher within-group association tendencies mix less widely in their networks and those with greater reassociation tendency are less likely to associate with new individuals and connect disparate parts of the

network [51]. Therefore, we predicted that as females aged, they would have lower measures of betweenness (capacity for linking discrete clusters in a network) and closeness (capacity to reach others or be reached), but a higher clustering coefficient (cliquishness).

To determine if age-based changes at the individual level can result in changes in overall network structure, we used empirical data from 19 networks of female rhesus macaques to test how variable proportions of old individuals in a network were related to three common measures of global network structure: mean degree, diameter and transitivity (table 1). We predicted that networks with a greater proportion of old individuals would be more sparsely connected due to older animals having fewer social partners (i.e. have lower mean degree and network diameter). Given greater kin clustering with age, we expected that networks with more old individuals would be more clustered (i.e. have higher transitivity). Finally, to help inform our empirical findings and better understand the link between age

distribution and global structure, we built an agent-based model (parameterized using information from our empirical data) to simulate how different proportions of old individuals would be expected to affect network structure under simplified conditions where everything else is equal. Our results provide a first step to understanding how and when individual changes in social tendencies with age might scale up to detectable effects on global network structure, offering important insights into the consequences of demography for the structure and function of collectives.

2. Methods

(a) Study population and data

Data used in this study were collected from a well-studied population of rhesus macaques on the island of Cayo Santiago, off the southeastern coast of Puerto Rico. The current population is maintained by the Caribbean Primate Research Center (CPRC) and is descended from 409 macaques that were introduced to the island from India in 1938. The animals are food supplemented and provided with *ad libitum* access to water. There is no medical intervention, and so the major causes of death at this predator-free site are illness and injury [55,56]. The CPRC staff collect demographic data 5 days per week and thus track dates of births and deaths for all individuals with a high degree of accuracy.

Rhesus macaques are highly social cercopithecine primates that live in matrilineal kin groups and exhibit clearly differentiated social relationships with kin-biased affiliation [55,57,58]. At 6 years old, females are deemed adults [59] and previous research on the macaque population of Cayo Santiago has shown that, for females that survive to reproductive age, the median lifespan is 18 years with a maximum lifespan of about 30 years [53,60]. Female rhesus macaques have a strict dominance hierarchy, with maternal rank inheritance and youngest ascendency [61]. Patterns of social interactions and social attention vary between young and old adults [6,34,62]. Female macaques show clear evidence of within-individual declines in the number of grooming partners with age, although the amount of time spent giving and receiving grooming remains constant across adulthood [6].

For this study, subjects were mature adult females of 6 years and older [59] from six naturally formed mixed-sex social groups. We used data collected between 2010 and 2017, a time period for which we had detailed behavioural data from which to estimate social networks. We collected behavioural data between 07:30 and 14:00, which are the working hours of the field station, using 10 min focal animal samples and recording all behaviours continuously [63]. We balanced data collection to ensure equal sampling of individuals throughout the day and over the course of the year, resulting in approximately the same number of focal samples per individual per year. For these analyses, we used grooming interactions to build our networks, given the clear age-based changes in grooming associations previously demonstrated in this system [6]. Grooming behaviour was collected by recording the duration of a grooming bout along with the identities of the interactants and the direction of grooming. We focused only on interactions between adult females (aged 6 years and older) and did not include interactions with infants (less than 1 year old), juveniles (2–3 years old) or sub-adult females (4–5 years old). We also did not include interactions with males as we wanted to avoid capturing changes in socio-sexual behaviour with age. We established dominance ranks for all females in a given year using observed win–loss interactions (as per [64,65]). Rank was assigned as ‘high’ (80% or more of other females dominated),

‘medium’ (50%–79% of other females dominated) or ‘low’ (up to 49% of other females dominated).

(b) Social networks

We built 19 grooming networks including all adult females from the following group-years (group F 2010–2017; group HH 2014 & 2016; group KK 2013, 2015 & 2017; group R 2015–2016; group S 2011; group V 2015–2017), with data availability based on the focus of projects over time given limited person power. We used weighted network metrics, as these are more robust and provide higher resolution than binary measures [66]. In these weighted networks, edges represented the undirected rate of grooming between a pair of individuals (seconds of grooming/total number of hours that both individuals were observed in focal animal samples). We note that using undirected grooming rates may fail to capture some finer nuances of age-based variation in social behaviour if, for instance, older individuals were likely to give less grooming than they received [61]. However, our most recent within-individual analyses suggest that while the number of grooming partners does decline as females age, the amounts of both grooming given and received remain constant [6]. Therefore, the use of undirected grooming rates to build networks should provide a comprehensive picture of how individual connectedness to the wider network changes with age, without missing essential changes in the form that the connectedness takes. The average size of our networks was 50.7 (\pm s.e. = 3.9) adult females. All network metrics were calculated in R (v.4.2.0; [67]) using the *igraph* package (v.1.3.1; [68]).

(c) Empirical analyses

All empirical models were fitted in a Bayesian framework with different error structures and random effects dependent on the data analysed (see below). We conducted all analyses using R (v.4.2.0; [67]) and fitted all models in the Bayesian software STAN [69] using the *brms* package (v.2.17.0; [70]). All fixed effects were given weakly informative priors (see electronic supplementary material for more details). We ran all models for 10 000 iterations across two chains with a warm-up period of 2000 iterations. We assessed model convergence by examining trace plots to assess sampling mixing and by ensuring $R_{hat} = 1$. We considered estimates of fixed effects to be different from zero when the 95% credible intervals of the posterior distribution did not overlap zero.

(i) Investigating the relationship between age and indirect connectedness

For these analyses, we set out to test how individuals’ level of indirect connectedness to their social network changed from prime adulthood into later life. In our dataset, the median age of adult females was 10 years and previous studies in this system have shown that individuals aged 10 and beyond show clear evidence of physical [71–74], immunological [75], reproductive [71] and social ageing [6]. To capture later-life changes in indirect connectedness, we focused our analyses on individuals aged 10 years and older, in line with previous ageing studies in this system [6]. Females in this analysis therefore ranged between 10 and 28 years old, although, to be clear, all measures of indirect connectedness for these females were extracted from the networks including all adult females aged 6 years and older; see above). We had 563 macaque years of data over 204 unique females, with an average of 2.8 years of data per individual (range: 1–8 years; electronic supplementary material, figure S1). Because there was variation in the age-ranges over which individuals were sampled, we used a within-individual centring approach to capture changes in indirect connectedness across individuals’ lifespans [76]. Briefly, following the methodology of van de Pol & Wright

[76], we split our age term into a between-individual effect (calculated as the mean age of an individual across all observations; hereafter called average age) and a within-individual effect (calculated as the deviation of an individual's age from their mean age; hereafter called within-individual age). This within-individual age term was our primary variable of interest and reflects how an individual's deviation from its age affects its indirect connectedness in the network (see [6] for a more detailed description of these methods).

We fitted four models with our four response variables of interest: eigenvector centrality, betweenness, closeness and clustering coefficient. All statistical models included average age and within-individual age as continuous fixed effects. Given the strict dominance hierarchy exhibited by female rhesus macaques [61], we assessed whether social status affected the change in an individual's measures of indirect connectedness with age [6] by fitting an interaction between rank and within-individual age in all models. We removed the interaction when not significant. We included individual ID, group and year, as random effects to account for repeated observations and to capture any variation in indirect connectedness measures that might be due to differences between individuals, groups or years. We also fitted within-individual age as a random slope over individual ID to capture any among-individual variation in the change in indirect connectedness with age. We had no biological reason to expect nonlinearities in the relationship between age and measures of indirect connectedness given that we were looking at changes from prime adulthood to old age and we've previously found a linear relationship between direct measures of connectedness and age [6]. Nevertheless, we fitted a model with quadratic terms for within-individual age and average age and compared that model to the model with only linear age terms using leave-one-out cross-validation in the brms package (v.2.17.0) [70]. The quadratic terms never improved the model fit and so were not considered further. For the eigenvector centrality model, we log-transformed the response variable to improve model fit and fitted a model with a Gaussian error distribution. For betweenness, we fitted a model with a zero-inflated Poisson distribution. For closeness and clustering coefficient, we fitted models with a Beta error distribution and a zero-one-inflated Beta error distribution, respectively.

(ii) Investigating the relationship between age distribution and network structure

Our analyses quantifying changes in indirect connectedness with age focused on females aged 10 years and older to capture individual changes in social network position from prime age and beyond. Here we were interested in how the age distribution of a group was linked to network structure. For these analyses, we therefore used our complete networks including all adult females aged 6 years and older. We had three response variables of interest (three measures of global network structure) and fitted a separate model for each: mean degree, diameter and transitivity. To ask if age distribution relates to network structure, we included the proportion of old individuals as a continuous predictor in all our models. Since 18 years is the median age of death in this population and maximum lifespan is about 30 years, we considered individuals above 18 years to be 'old' [52,61]. The proportion of old females in our 19 networks ranged from 0.03 to 0.19 (figure 1; electronic supplementary material, figure S2). Given that rhesus macaques live in matrilineal groups with kin-biased behaviour [53,77], we included the average relatedness of the network as a continuous covariate to account for differences between groups in general relatedness. We also included network density (calculated as the number of existing ties in the network divided by the number of possible ties) as a continuous fixed effect in the models of diameter and

transitivity to account for the fact that density can drive variation in other global network metrics. We included year as a random effect to account for yearly variation that might affect network structure. For the diameter model, we log-transformed the response variable to improve model fit. We fitted all models with a Gaussian error distribution. To account for a potential nonlinear relationship between age distribution and network structure, we fitted all models with a quadratic term for proportion of old individuals and compared that to the model with only linear terms using leave-one-out cross-validation in the brms package (v.2.17.0; [70]). The quadratic terms never improved model fit and so were not considered further.

(d) Agent-based model of the relationship between age distribution and network structure

We expected that age-based changes in patterns of direct association would scale up to affect overall network structure. However, our empirical data revealed no effect of the proportion of old individuals in the network on global network metrics (see §3 below). To better understand these results, we built an agent-based model where we could manipulate age-based differences in sociality in isolation of other variables to better explore how different age distributions would be expected to affect overall network structure, and why such scaling up of age-based differences in social behaviour may not be detectable in our data. Our general approach was to simulate artificial populations that exhibited the social age-dependencies observed in the rhesus macaque system on Cayo Santiago and to use these to investigate the relationship between age distribution and global network metrics. As mentioned above, adult female rhesus macaques change two aspects of their sociality with age: their number of social partners, and the proportion of partners that are kin [6]. These individual changes in sociality across adulthood lead to differences between young and old individuals in both their number of social connections and probability of connecting with kin [6].

The agent-based model therefore simulates social networks of varying age distributions, in which the probability that two nodes (i.e. individuals) have an edge (i.e. a grooming link) depends on the age of the individuals and their kinship to each other. Individuals belonged to two age categories (as in the empirical analysis for global structure; old adults and young adults, hereafter 'old' and 'young') and two kin categories (kin and non-kin). This gives six dyad types, which we denote by the age categories of the two individuals and their kinship status (e.g. old/old kin). We established the linking probability (i.e. the chance of having an edge in the social network) for each dyad type based on the mean proportion of dyads of that type that had an edge across the 19 empirical macaque grooming networks. The linking probability for dyads that were old individuals who were related to each other (i.e. old/old kin) was 0.33; for old/old non-kin dyads it was 0.02; for old/young kin dyads, it was 0.37; for old/young non-kin dyads, it was 0.05; for young/young kin dyads, it was 0.27; and for young/young non-kin dyads it was 0.08. We fixed group size at 50 individuals, which approximates the mean number of adult females in real groups on Cayo Santiago (mean \pm s.e. = 50.7 \pm 3.9). Each simulated network had 10 clusters of individuals who were related to each other (i.e. kin groups), with five individuals in each. The number of individuals within each kin group mirrors the mean number of close adult female kin (minimum relatedness coefficient 0.125) that adult female rhesus macaques have on Cayo Santiago (mean \pm s.e. = 5.2 \pm 0.87). In the model, kinship between pairs of individuals was determined by their kin group membership: individuals from the same kin group were classed as kin and individuals from different kin groups were classed as non-kin.

Each simulation round (i.e. construction and quantification of one network) proceeded as follows. We first randomly drew the

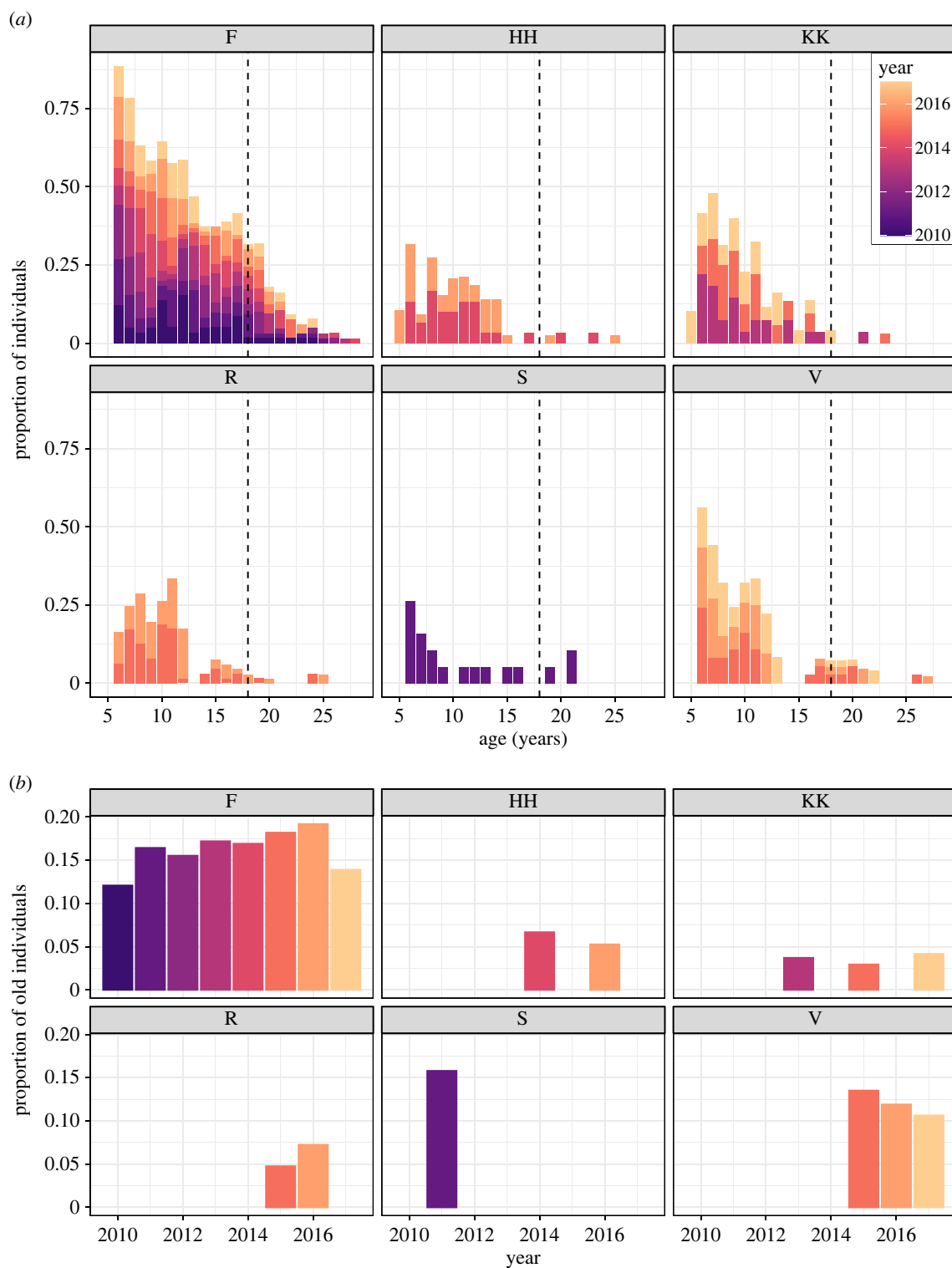


Figure 1. (a) Distribution of ages in each of the six empirical macaque groups for each year that those groups were observed (19 networks total observed between 2010 and 2017). The data are presented as a stacked bar chart, so to compare the proportion of x -year olds in a single group across years one should compare the height of coloured bars within a single age. The dotted black lines at age 18 indicate the median age of death in each population and the cut-off at which we considered individuals 'old' for the sake of calculating the proportion of old individuals in the group. (See also electronic supplementary material, figure S2.) (b) Proportion of old individuals in each group for each year that group was observed. Years with no bars indicate years in which the group was not observed.

number of old individuals in the group (n_{old}) from a uniform distribution with a set range [0, 50]. We then randomly assigned all 50 group members to age-groups (n_{old} old individuals and $n_{young} = 50 - n_{old}$ young individuals) and kin groups. This allowed the age structure of the groups to vary across the range from only young individuals (0% old) to only old individuals (100% old). We then constructed the social network by drawing links, where the chance of each dyad getting a link depended on their dyad type (linking probabilities given above). To determine whether a dyad was given a link, we extracted a random value from a binomial

distribution with a sample of 1 and probability equal to the linking probability of the type of dyad. If the extracted value was 1, the dyad was given a link, if it was 0, the dyad was not given a link. A schematic representation of the various steps of the model-building process can be found in the electronic supplementary material, figure S3.

To confirm that the model was working as intended, we ran 10 000 simulations (i.e. we generated 10 000 networks), from which we calculated the proportion of dyads of each type that had a link. We confirmed that the means of these proportions

(old/old kin = 0.33; old/old non-kin = 0.02; old/young kin = 0.37; old/young non-kin = 0.05; young/young kin = 0.27; young/young non-kin = 0.08) corresponded well to their respective empirical linking probability (see above, and electronic supplementary material, figure S4). We also confirmed that our model was not highly sensitive to our input linking probabilities (see electronic supplementary material, methods, table S1 and figure S5 for more information). Given that our input values are only estimates of the real values, robustness to these values is important for drawing general conclusions about the potential effects in the real system.

To investigate the general relationship between age distribution and network structure, we ran 100 000 replicates of the simulation (i.e. generated 100 000 networks) where we allowed the proportion of old individuals in the network to range from 0.00 to 1.00 and visualized the relationship between the proportion of old individuals and network mean degree, diameter, and transitivity. Additionally, we used the agent-based model to investigate if we should expect to find relationships between the proportion of old individuals and network metrics in the empirical analysis. To do this, we limited the proportion of old individuals in each network to between 0.04 and 0.20 (mirroring the empirical variation in age distributions), and we ran 19 simulations (mirroring the 19 empirical networks). We repeated this 50 times (i.e. 50 sets of 19 networks) to gauge consistency of the results.

3. Results

(a) Relationship between age and indirect measures of network connectedness

In line with our predictions, female macaques did not show any change in eigenvector centrality with age (within-individual age: $\beta = -0.05$; 95% CI = $-0.35, 0.24$; figure 2*a,b*, electronic supplementary material, table S2). That is, as females aged, the strength of their relationships to their partners, and to their partners' partners, was stable. We did find evidence of a within-individual decline in betweenness with age. However, this effect was rank dependent and seems to be driven primarily by a decline in betweenness as high-ranking individuals got older (within-individual age:rankH: $\beta = -0.35$; 95% CI = $-0.41, -0.29$; figure 2*c,d*, electronic supplementary material, table S3), although mid-ranking individuals also showed greater declines in betweenness with age than did low-ranking females (within-individual age:rankM: $\beta = -0.14$; 95% CI = $-0.18, -0.09$; figure 2*c,d*, electronic supplementary material, table S3). Therefore, some, but not all individuals became less effective at reaching disparate nodes in the network as they got older. Individuals showed a reduction in their closeness with age (within-individual age: $\beta = -0.18$; 95% CI = $-0.35, -0.02$; figure 2*e,f*, electronic supplementary material, table S4), meaning they were harder to reach and be reached in the network, as we predicted. Contrary to our predictions, there was no within-individual effect of age on clustering coefficient (within-individual age: $\beta = 0.04$; 95% CI = $-0.06, 0.14$; figure 2*g,h*, electronic supplementary material, table S5). That is, individual cliquishness was stable as individuals aged.

(b) Relationship between age distribution and network structure

(i) Empirical results

Contrary to our predictions, we found no evidence that groups with a greater number of old individuals were structured differently from groups with fewer old individuals. There

was no overall effect of the proportion of old individuals in the group on mean degree ($\beta = -2.76$; 95% CI = $-10.24, 5.25$; $n = 19$; figure 3*a*, electronic supplementary material, table S6) or network diameter ($\beta = -0.12$; 95% CI = $-3.93, 3.67$; $n = 19$; figure 3*b*, electronic supplementary material, table S7). Therefore, networks were not more sparsely connected as the proportion of older animals increased. Older networks were also not more clustered or cliquish, as measured by transitivity ($\beta = 0.20$; 95% CI = $-0.29, 0.70$; $n = 19$; figure 3*c*, electronic supplementary material, table S8).

(ii) Agent-based model results

Different group demographics in age appeared to have important consequences for network structure in our simulations (figure 4; electronic supplementary material, figure S6). Specifically, our 100 000 simulations where we allowed the proportion of old individuals in the network to range from 0.00 to 1.00 showed that, as the proportion of old individuals in the population increased, mean degree (i.e. the mean number of partners that each individual associated with) decreased (figure 4*d*), and the diameter of the network (i.e. the longest path length in the network) increased (figure 4*e*), both as expected. Network transitivity (i.e. the degree of clustering in the network; figure 4*f*) showed little variation in relation to age distribution.

Importantly, these global metrics did not always exhibit linear relationships with network age (as might have initially been intuitively expected). The potential for nonlinearity is important for understanding when the effects of age on network structure might be the most pronounced and most detectable in the real world. In particular, the increase in network diameter with an increasing proportion of old individuals showed an accelerating trend (figure 4*e*). That is, the relationship between network diameter and proportion of old individuals was stronger (i.e. the slope was steeper) when the proportion of old individuals in the network was high, suggesting that effects of age on network structure might be most detectable when, for example, more than 50% of the individuals in the network are old. While we did not find any clear change in transitivity with the proportion of old individuals in the network, there was also some evidence of an accelerating trend at the higher end of the age distribution. These metrics (transitivity, diameter) are driven to a greater extent by modularity in networks and the nonlinear relationship could arise from the kin-biased nature of the relationships in our (modelled) study system. We also observed that transitivity and network diameter were more variable when there was a greater proportion of old individuals in the network. This suggests that changes in network structure with changing age distribution might be unpredictable. For example, networks might become more clustered or less clustered as a population gets older. Overall, these results suggest that compared to networks with more young individuals, networks with more old individuals are sparser and less cohesive, and can take on more varied structures.

When we limited the sample size and variation in age distribution to those of the empirical data (i.e. 19 networks, where the proportion of old individuals was only allowed to vary between 0.04 and 0.20), we observed no clear change in network structure as the proportion of old individuals in the network increased (figure 4*g-i*).

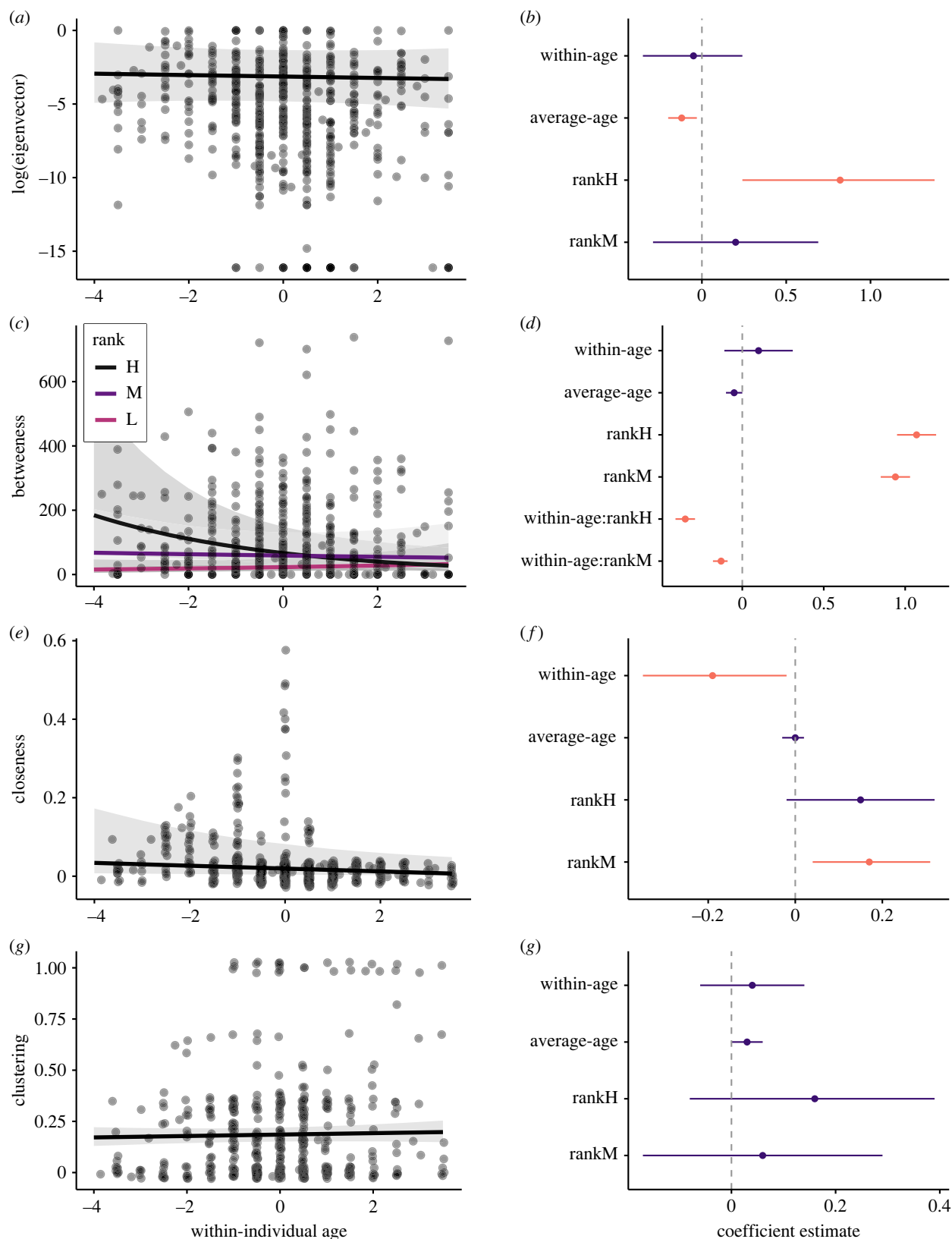


Figure 2. Relationship between within-individual changes in age and indirect measures of connectedness including (a,b) eigenvector centrality, (c,d) betweenness, (e,f) closeness and (g,h) clustering coefficient in female rhesus macaques. In (a,c,e,g), the points represent raw data. Shaded grey bars indicate 95% confidence intervals around the predicted values. In (b,d,f,h), effect sizes and 95% credible intervals (CI) for all fixed effects and interaction terms are shown. Instances where the 95% CI overlaps zero are coloured in purple.

4. Discussion

Age has begun to emerge as an important attribute shaping the social decisions of individuals [6,7,14]. This implies that age is a potentially significant feature underlying the behaviour of animals in collectives. Here we have shown that

age-based changes in social behaviour are likely to have important consequences for an individual's position within their wider social network and may scale up to influence network topology. As female rhesus macaques aged, they showed declines in some, but not all of their indirect measures of connectedness. But despite age-based changes

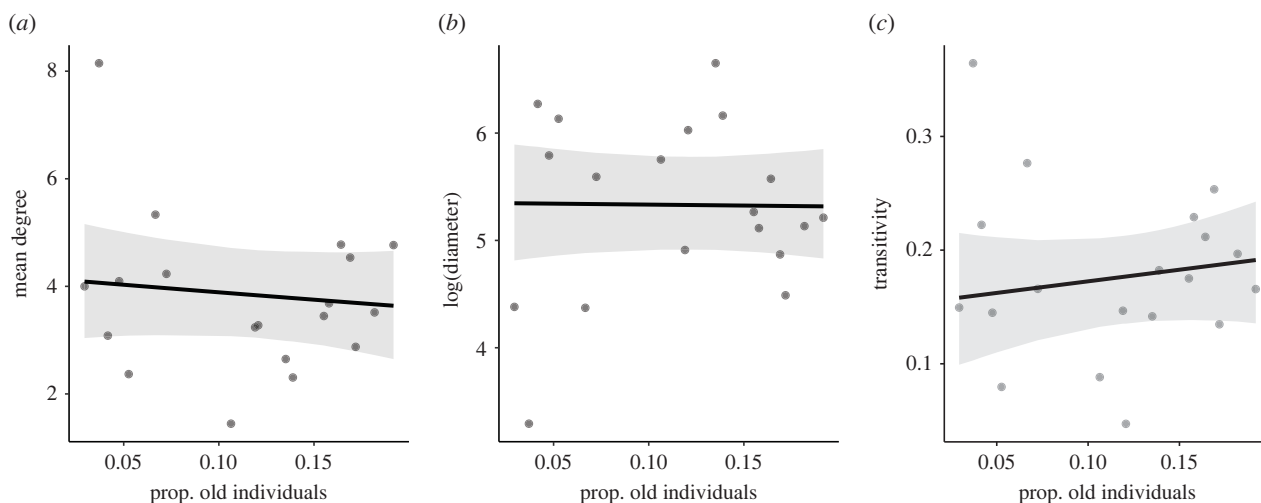


Figure 3. The relationship between the proportion of old individuals in female rhesus macaque networks and (a) mean degree, (b) network diameter and (c) network transitivity. Points represent raw data and shaded grey bars indicate 95% confidence intervals around the predicted values.

in both direct connectedness [6] and indirect connectedness (this paper), we detected no effect of age distribution on the overall structure of rhesus macaque networks. The agent-based model gives insight into this surprising result, as it implies that age-based differences in social behaviour do not necessarily, as expected, scale up to affect network structure, but that these effects may be nonlinear and so may not always be easily detectable.

In line with our predictions, we found no evidence for changes in eigenvector centrality with age. This was expected because, despite reducing their number of social partners, female rhesus macaques maintain the amount of time they engage in social interactions as they get older [6], allowing them to continue to have strong connections to some of their partners, and (indirectly) to the partners of their partners. A similar pattern has been observed in a population of wild rhesus macaques in China whereby older individuals had fewer partners but exhibited similar weighted eigenvector centrality to those of younger individuals [34]. Unexpectedly, we did not find any evidence that clustering coefficient increased as individuals aged in this population. Females increase their preference for grooming kin as they age [6], suggesting that within-group (e.g. homophilic) tendencies increase with age, which should lead to greater clustering as females from the same matriline would cluster together in the network. However, although kin-biases increase with age, they exist for females of all ages [58]. It is therefore possible that ties with kin are the main driver of clustering coefficient scores, regardless of age. The loss or removal of a few non-kin relationships as an individual ages could thus have a relatively minor impact on their clustering coefficient, leading to stable values for this metric across the adult lifespan. In general, the stability of weighted eigenvector centrality and clustering coefficient across the adult lifespan suggests that older animals can remain socially central and well integrated in some respects, despite other aspects of their social life changing (e.g. fewer partners overall, and the patterns observed below). Old individuals may therefore continue to reap some of the advantages of social relationships [5].

As predicted, we found declines in betweenness and closeness with age, although some of these results were rank-dependent. As individuals got older, particularly those

that were higher-ranking had less influence on and became less well connected to the wider network. Betweenness and closeness are associated with an individual's level of reassociation tendency and within-group association [51]. In other words, the more likely individuals are to reassociate with the same partners, and to select partners of similar characteristics to themselves, the less they mix with the wider network and the less likely they are to connect distinct sub-groups (i.e. have high betweenness) or be easily reached by all others in the network (i.e., to have high closeness). We've previously shown that ageing rhesus macaque females have fewer partners, but engage with those partners more often, and increase their preference for close relatives. In other words, while mean dyadic association strength to associates increases with age, mixing with the wider network should decline [6]. Our betweenness and closeness results seem to reflect these underlying changes in behavioural patterns. The declines in betweenness with age were rank-dependent, with mid-ranking and high-ranking rhesus macaque females showing a greater decline in betweenness with age than low-ranking individuals. This may simply be due to a floor effect—because low-ranking individuals have such low betweenness to begin with, there is little capacity for further decline with age.

Generally, our findings demonstrate that how indirectly connected an individual is to their social world can change across their lifespan. How an individual is positioned in the wider social environment can modulate their exposure to information [26,31], parasites [18,77] and pathogens [20–22]. Individuals with high betweenness and closeness occupy a critical position in the acquisition and transfer of 'goods' within a network [17,54,78]. Decreases in both of these measures of indirect connectedness with age may benefit aged individuals who may experience greater susceptibility to disease or illness as a result of immunosenescence [79]. Such benefits need not imply that individuals actively change complex network positions with age. By changing simple behavioural rules or processes with age, changes in polyadic ties with age are likely to emerge [51]. Measures of indirect connectedness have been linked to fitness proxies including future social status [80,81], survival [82,83] and reproductive success [32,64,81]. Indirect network metrics can be an even stronger predictor of fitness proxies [17,32] than measures of direct

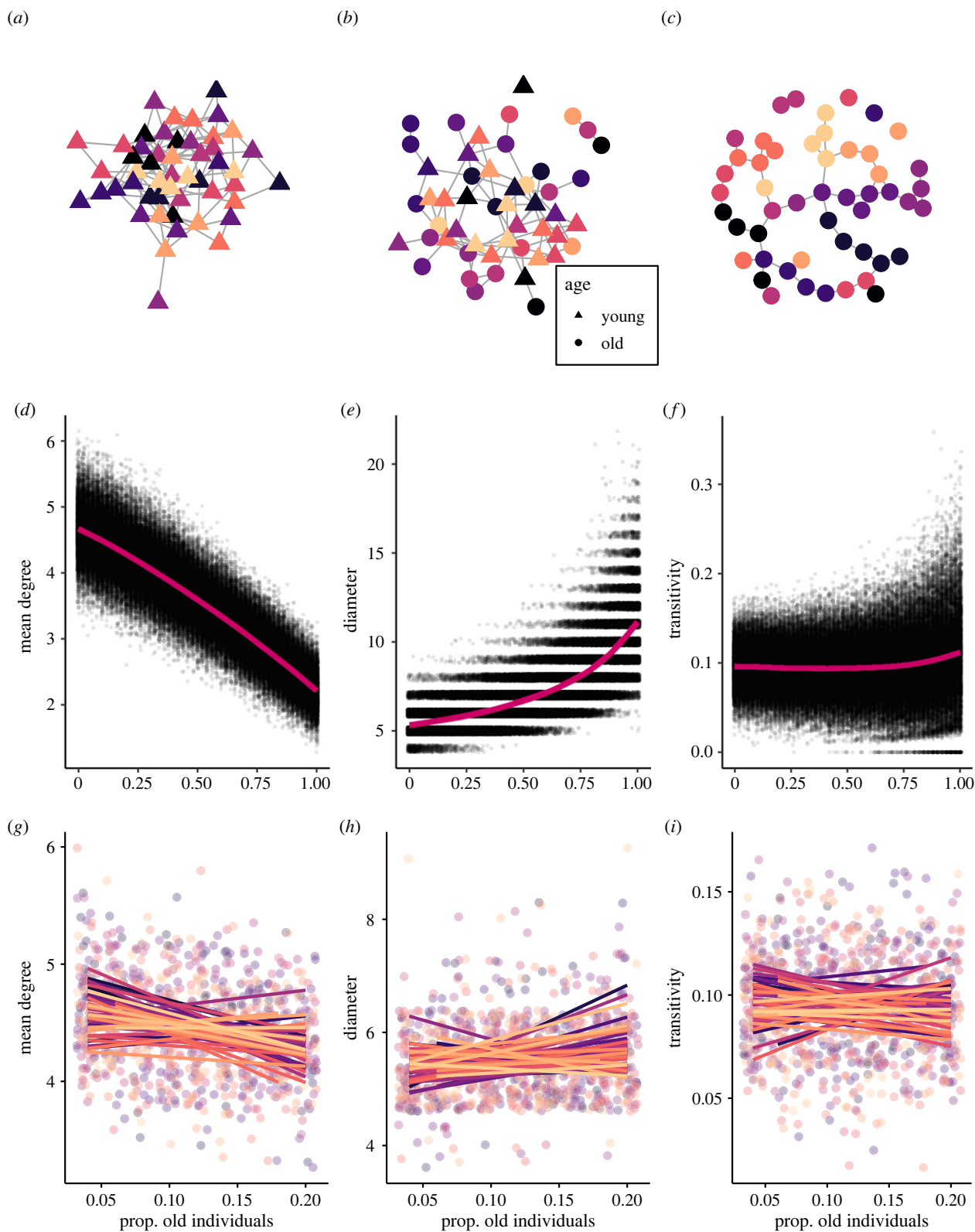


Figure 4. (a–c) Example networks from the agent-based model illustrating differences in network structure with the proportions of old individuals in the network equal to 0 (a), 0.5 (b) and 1 (c). Node colour represents kin groups, where nodes of the same colour belong to the same kin group. (d–f) Results from the agent-based model showing relationship between the proportion of old individuals and (d) mean degree, (e) diameter and (f) transitivity. Data points from the 100 K simulations are shown with a small amount of jitter introduced to show overlapping points and are fitted with a smoothing term. (g–i) Relationship between the proportion of old individuals and (g) mean degree, (h) diameter and (i) transitivity based on 19 simulations from the agent-based model where the proportion of old individuals in the population was restricted between 0.04 and 0.20. We re-simulated these 19 networks with limited age variation 50 times to help ensure that our results were robust. Each colour represents a different ‘bout’ of 19 networks. Data points are shown fitted with a linear smoothing term. (Online version in colour.)

connections and may more strongly reflect the underlying behavioural rules that give rise to individual differences in sociality [51]. As such, documenting how ageing shapes the

polyadic social world may be particularly relevant for understanding how changes in sociality across the lifespan influence patterns of senescence and fitness in later life.

While age is clearly associated with changes in the behaviour and social connectedness of animals living in groups [10,13,14], including in this population of rhesus macaques [6,60], we did not find any empirical evidence that age impacts the overall structure of those groups. We found no relationship between age distribution and network mean degree, diameter or transitivity in the observed macaque data. This was surprising to us given that we expected that age-based changes at the individual level would scale up to the network level. That is, it should be self-evident that a network with more old individuals who each interact with fewer partners would, for example, have a lower mean degree. Given that we did not find empirical evidence of this, we used an agent-based modelling approach to try to understand why these results might emerge and whether under a scenario where everything else is equal, we could recreate these expected effects on network structure.

By modelling age-based differences in two interaction patterns (number of social partners and the tendency to link with kin) based on the age-based changes we observed in the female macaques [6], we found that the age composition of a group can have important consequences for its cohesiveness and connectedness. As predicted, mean degree declined and diameter increased as the proportion of old individuals increased, while transitivity exhibited no strong relationship to age distribution. Interestingly, these effects did not necessarily scale in the linear manner that might be expected in response to a linear increase in proportion of old individuals in the network. Network diameter showed a steeper increase as networks became older, suggesting that the strength of the effect, and thereby the ability to statistically detect an effect in a real-world case, might depend on where along the continuum of age distributions one's data lie. The combination of this and a limited sample size could potentially have led to the null result in our empirical analysis. This interpretation is supported by the fact that when we limited the age variation and sample size (number of networks) of our model to match that of the empirical data (i.e. we only ran the model for 19 simulations and allowed the proportion of old individuals in the network to range between 0.4 and 0.20) there was no clear relationship between the proportion of old individuals and network structure.

The results of our model suggest that populations made up of very large numbers of young, or old, individuals may have detectable levels of divergence in network structure, whereas populations within a smaller range of age distributions may not. Sample size also appears to have played an important role in our ability to detect an effect of network age on network structure. Each 'bout' of 19 networks (simulations) produced a slightly different result. Sometimes slopes were positive, sometimes negative, sometimes flat, suggesting there is a fair amount of stochasticity in our ability to detect the true effect when the sample size is small, even in the model world where there are no other confounding factors. While 19 rhesus macaque networks from six groups collected across 8 years represent a considerable investment of research time and effort, it may not have been enough to detect a clear pattern. However, it is also possible that our empirical results reflect a true null result if there are processes occurring that our model has not accounted for. Our model is a simplistic version of the real world and does not fully replicate the complete suite of changes that might occur as individuals age. For example, our model does not allow for

potential rewiring between nodes that might occur if younger individuals choose to build new ties in response to the loss of ties with older individuals. Such rewiring could explain why some measures of network structure do not change despite individual declines in sociality with age.

While the general result from the agent-based model was expected, these simulations help inform our understanding of the empirical data and offer us a glimpse into how age-based differences in sociality may affect network structure and when these effects might be detected. The fact that, even in our simplified model where there is nothing to obscure individual linking probabilities from scaling up to affect global network metrics, changes in some measures of network structure appear most likely to be detected in very old populations raises the question of when such effects are likely to be detectable in wild systems. The macaques of Cayo Santiago are a managed population that are food supplemented and have no predators. As such, the proportion of old individuals seems likely to be higher than in many wild populations and aligns with other natural but non-predated systems (e.g. red deer [84], where the proportion of old individuals in the population appears to reach a maximum of about 20%). Our model suggests that the effects of age on network structure are also present at this lower end of the age distribution, but are less strong, or negligible, for some network measures. Hence, detecting an effect in real systems could necessitate large amounts of data. We note, however, that the effect on the network measures (the shapes of the curves) will depend both on how sociality changes with age and the social organization of the system (such as relatedness patterns). While other systems may be similar in these regards to the one investigated here, making predictions for detectability of social ageing effects in general would require more comprehensive modelling.

Future work in other systems, conditions and contexts could reveal consequences of age-based changes in sociality for network structure that are even more stark. For example, greater differences between age classes in the overall probability of forming a social tie may be expected in societies that experience selective disappearance of the most sociable individuals or where declines in sociality with age are exacerbated by partners dying and not being replaced (which we do not observe in this system [6]). The kin structure of populations is also likely to be important if the probability of connecting decreases with age but there is no compensatory increase in the probability of forming ties with kin. This could lead to stronger absolute effects on global network metrics, including substantial changes in transitivity, and thus would likely have further consequences for network connectivity and spreading processes. While our analyses bin individuals into age categories of 'old' and 'young', there may be much more nuanced effects of age on network structure if the full age distribution of a population or group is taken into account.

It is also important to recognize that the results presented here only reflect how changes in female sociality across adulthood are expected to influence adult female social networks. Understanding how social networks of entire groups are affected by age-based changes in behaviour more broadly will necessitate incorporating an understanding of the patterns of social ageing in both sexes, as well as an understanding of how interactions with juveniles shift as adults get older. While much less is known about the social

ageing trajectories of males, some studies suggest that they show lower levels of connectedness at older ages, similar to those observed in females [13,33]. But even in cases where social ageing trajectories are similar across the sexes, changes in socio-sexual interactions with age, in addition to shifting interactions between age classes, are likely to produce much more complex network dynamics than observed here, necessitating further research.

Despite these limitations, our study provides a first glimpse into the effects that age-based changes in behaviour might have on network structure, offering a deeper understanding of the potential importance of demographic changes for the structure and function of collectives. Much attention has been given to the potential for the social structure of populations to be affected by the removal of older individuals, for example through practices like trophy hunting or fishing [85,86]. This is relevant given that older individuals can play a critical role in their social groups through leadership [4,39], inter-generational transfers [40] and the stabilization of social relationships [87]. Despite the importance of the loss of older individuals from networks, our results suggest that asking what happens when populations contain large numbers of older individuals may be an equally salient question.

In humans, population ageing is poised to be one of the most significant social transformations of the twenty-first century, with the global number of older persons projected to more than double over the next three decades [88], facilitated by falling fertility rates and increasing longevity [89,90]. Meanwhile, in other group-living species, anthropogenic challenges, including climate change, poaching, disease outbreaks and pollution, among others, are likely to cause major population declines and demographic shifts [91,92]. The consequences of declining fertility and/or increasing mortality rates may be further exacerbated by the concomitant shifts in age structure and resulting implications for network structure and function. For instance, declines in cohesiveness and structural connectedness in older networks may limit or slow the transfer of information [24,93], restrict the potential for cooperation [94] or reduce the stability of populations [30], feeding back to further limit population growth. Alternatively, such changes

could reduce the vulnerability of populations to infectious diseases or parasites [30,95] or enhance cooperation if network modularity is increased [96]. Consequently, through its effects on network structure, shifting age demography may have broader implications than previously appreciated for group dynamics and persistence and warrants further research.

Data accessibility. All data are available from the Figshare Repository: <https://doi.org/10.6084/m9.figshare.21936426> [97].

The data are provided in the electronic supplementary material [98].

Authors' contributions. E.R.S.: conceptualization, formal analysis, investigation, methodology, writing—original draft and writing—review and editing; A.S.P.: formal analysis, visualization, writing—original draft and writing—review and editing; J.B.B.: formal analysis, visualization, writing—original draft and writing—review and editing; J.N.V.: data curation; D.P.: data curation; CBRU.: resources; M.L.P.: funding acquisition and resources; J.P.H.: conceptualization, funding acquisition, project administration, supervision and writing—review and editing; N.S.-M.: conceptualization, funding acquisition, project administration, supervision and writing—review and editing; L.J.N.B.: conceptualization, data curation, funding acquisition, investigation, methodology, project administration, resources, supervision, writing—original draft and writing—review and editing.

All authors gave final approval for publication and agreed to be held accountable for the work performed therein.

Conflict of interest declaration. We declare we have no competing interests.

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