Explaining negative kin discrimination in a cooperative mammal society

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Kin selection theory predicts that, where kin discrimination is possible, animals should typically act more favourably towards closer genetic relatives, and direct aggression towards less closely related individuals. Contrary to this prediction, we present data from an 18-year study of wild banded mongooses, Mungos mungo, showing that females that are more closely related to dominant individuals are specifically targeted for forcible eviction from the group, often suffering severe injury, and sometimes death, as a result. This pattern cannot be explained by inbreeding avoidance or as a response to more intense local competition among kin. Instead, we use game theory to show that such negative kin discrimination can be explained by selection for unrelated targets to invest more effort in resisting eviction. Consistent with our model, negative kin discrimination is restricted to eviction attempts of older females capable of resistance; dominants exhibit no kin discrimination when attempting to evict younger females, nor do they discriminate between more closely or less closely related young when carrying out infanticidal attacks on vulnerable infants who cannot defend themselves. We suggest that in contexts where recipients of selfish acts are capable of resistance, the usual prediction of positive kin discrimination can be reversed. Kin selection theory, as an explanation for social behaviour, can benefit from much greater exploration of sequential social interactions.

kin selection | kin discrimination | conflict | cooperation | eviction

Kin selection theory aims to understand how selection acts on social traits, such as altruism and selfishness, that affect the fitness of social partners and local group members (1, 2). The theory predicts that where animals can discriminate between more closely and less closely related individuals within their social group, they will preferentially direct altruism towards closer genetic relatives, and aggression towards less closely related targets (1–3). Instances of such positive kin discrimination are taxonomically numerous and widespread (4), while reported examples of negative kin discrimination are rare (5, 6).

Contrary to the predictions of traditional kin selection theory, we show below that banded mongooses exercise negative kin discrimination during attempts at eviction; and that this result is not readily explained by inbreeding avoidance or local kin competition. However, a simple and very general sequential game model of selfish behaviour that takes into account the possibility of active resistance on the part of recipients does provide a potential explanation for negative kin discrimination. We first describe the model, and then test predictions of the model using our data.

Explaining negative kin discrimination: a model

Consider the interaction between two individuals, the first of whom (Player 1) may perform a selfish act at the other's expense, such as stealing a food item, killing offspring, or, in the case with which we are concerned, evicting the other from the territory or group. We suppose that this act entails some fitness cost to the actor, denoted c_1 , but that the cost is outweighed by the benefit to be gained, $b_1 > c_1$. The act, if carried out, also deprives the recipient, Player 2, of a benefit b_2 that it would otherwise enjoy. If the two individuals are related by a coefficient *r*, then Hamilton's

rule tells us that the act will be favoured by selection provided that $b_1 - c_1 - r b_2 > 0$. Consequently, selfishness will never be directed toward a closer relative where it would not also be directed toward a more distant one.

Suppose, however, that if Player 1 attempts the selfish act, Player 2 may then choose to resist. Resistance ensures that the act will fail; Player 1 will still suffer the cost c_1 of attempting the act, but will not enjoy the benefit b_1 of success, nor will Player 2 suffer the consequent loss of benefit b_2 . At the same time, resistance entails a fitness cost to Player 2 of c_2 . This situation may be modelled as a two-step, sequential game, as illustrated in Figure 1 (see Supplementary Information for a population genetic formulation of the model).

How does relatedness affect the outcome of this game? Applying Hamilton's rule once again, if $c_2 > b_2 - r b_1$, then the cost of resistance to Player 2 outweighs the benefit of preventing a selfish act. Under these circumstances, Player 2 will submit, and selection once again favours performance of the selfish act provided that $b_1 - c_1 - r b_2 > 0$. If, by contrast $c_2 < b_2 - r b_1$, then Player 2 will resist, and selection consequently does not favour the selfish act. Since a more closely related recipient of the selfish act may prefer to submit where a more distantly related recipient would resist, it follows that selfishness may be directed toward a closer relative where it would not be directed toward a more distant one. To be precise, if we consider two levels of relatedness, r_{high} and r_{low} ($r_{low} < r_{high}$), then provided that

$$\frac{b_1 - c_1}{b_2} > r_{\rm high} > \frac{b_2 - c_2}{b_1} > r_{\rm low}$$

Significance

Kin selection theory predicts that animals will direct altruism towards closer genetic relatives, and aggression towards more distantly related individuals. Our 18-year study of wild banded mongooses reveals that, unusually, dominant individuals target females who are more closely related to them for violent eviction from the group. This puzzling result can be explained by selection for unrelated individuals to resist eviction, and for related individuals to submit more easily. In support of this idea, we show that kin are targeted for aggression only when individuals are capable of resisting. Our results suggest that, where potential victims can oppose aggression, the usual predictions of kin selection theory can be reversed.

Reserved for Publication Footnotes

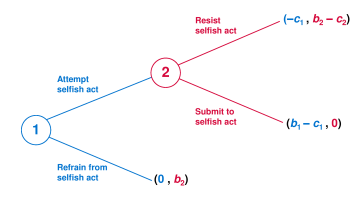


Fig. 1. A sequential model of selfishness and resistance. Player 1 (blue) first chooses whether or not to attempt a selfish act at the expense of Player 2 (red); in the event of such an attempt, Player 2 then chooses whether or not to resist. Direct fitness payoffs to both players are shown in their respective colours.

then the subgame-perfect equilibrium of the game features targeting of a selfish act towards a recipient related to the actor by r_{high} , but not towards a recipient related by r_{low} . To illustrate, suppose $b_1 = b_2 = 1$, $c_1 = 0.25$ and $c_2 = 0.75$; then the model predicts that a recipient of low relatedness ($r_{\text{low}} < 0.25$) would resist a selfish act, and hence should not be targeted, while a recipient of higher relatedness ($0.75 > r_{\text{high}} > 0.25$) will submit, and so should be targeted.

Unrelated recipients are more likely to resist a selfish act, favouring preferential targeting of more closely related victims, when c_2 (the cost of resistance) is small, and b_2 (the benefit to be retained) is large. These effects are illustrated in Figure 2. If c_2 is typically too large, or b_2 too small, even unrelated recipients are unlikely to resist, favouring indiscriminate selfishness; only for smaller values of c_2 or larger values of b_2 is negative kin discrimination predicted.

Negative kin discrimination in banded mongooses

We examine kin discrimination in the context of a conspicuous form of intragroup aggression in cooperatively breeding banded mongooses, Mungos mungo: the violent eviction of males and females from the group. Banded mongooses live in highly cooperative groups with limited dispersal and varying levels of relatedness between group members (7). In our study population in Uganda, groups consist of around 20 adults, plus offspring, and breed on average four times per year (8). Multiple females give birth synchronously to a communal litter that is cared for by members of both sexes (typically not the parents) (7). Each group contains a cohort of multiple dominant females (median = 4) that are older than the other females, breed more regularly and more successfully, and aggressively evict younger females (7, 9, 10). Older males monopolise mating with oestrus females by mate-guarding them and aggressively driving away younger, subordinate males (10, 11). Previous work shows that both sexes are capable of kin discrimination in the context of mating, supporting the assumption of our model that such discrimination is possible in this system (12). Evictions are relatively common, involve intense, targeted aggression, and result in the forcible mass exclusion of groups (median = 6 individuals) of females (female-only evictions) and, in around half of cases, groups (mean = 9 males) of males alongside them (mixed-sex evictions; 13). These mass evictions are triggered by high levels of intrasexual reproductive competition: females are evicted when there are many breeding females in the group, and males are evicted alongside females when there are many breeding males (13). Evictions are very violent, and evictees are often left with serious injuries as a result of the aggression they receive (see Video S1).

Among adult female banded mongooses, those more closely related to dominant individuals of both sexes (that is, males and females older than 3 years) in the group were more likely to be targeted for eviction (Generalised Linear Mixed Model (GLMM), $\beta \pm SE = 9.95 \pm 3.36$, $\chi^2_1 = 9.51$, p = 0.002; Figure 3A; Table S1). Younger females were also more likely to be subject to an eviction attempt (GLMM, $\beta \pm SE = -0.003 \pm 0.0005$, $\chi^2_1 = 39.98$, p < 0.0001; Table S1), but there was no effect of a female's pregnancy status or non-pregnant weight on her probability of being targeted for eviction (pregnancy status: GLMM, $\beta \pm SE = -0.61 \pm$ 0.49, χ^2_1 =1.63, p=0.20; weight: GLMM, $\beta \pm$ SE=0.002 \pm 0.001, χ^2_1 =2.59, p=0.11; Table S1). For a subset of eviction attempts we had data on the identity of 'primary aggressors' (individuals that were recorded as being notably more aggressive than other group members toward potential evictees). Females that were more closely related to primary aggressors were more likely to be targeted for eviction (GLMM, $\beta \pm SE = 5.47 \pm 2.87$, $\chi^2_1 = 3.85$, p = 0.0498; Table S2).

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We did not find evidence that this pattern of negative kin discrimination was an attempt to reduce inbreeding between targeted females and dominant males. We found that it was relatedness of females to female dominants (not male dominants) that predicted whether they were targeted for eviction. Females were more likely to be targeted if they were more closely related to female dominants (females older than 3 years; GLMM, $\beta \pm$ SE=5.10 ± 2.65, χ^2_1 =4.07, p=0.044; Table S3), but not if they were more closely related to male dominants (males older than 3 years; GLMM, $\beta \pm SE = 3.69 \pm 2.55$, $\chi^2_1 = 2.09$, p = 0.15; Table S3). Behavioural data indicate that females are almost entirely responsible for the attempts at eviction of other females: males rarely attacked females in female-only eviction attempts (GLMM, β \pm SE=1.80 \pm 0.56, χ^2_1 =12.48, *p*<0.001; Figure 3B; Table S4). Previous work indicates that eviction attempts are triggered when the level of reproductive competition in the group is high, not when the potential for inbreeding is high (13); and that both sexes use kin discrimination during mating to avoid inbreeding (12).

We also did not find evidence for the alternative hypothesis that closer relatives are targeted for eviction because they inflict higher costs on the reproductive success of dominants. Dominant females did not suffer greater reproductive costs when co-breeding with more closely related females. There was no difference in the number of emergent pups to which dominant females gave birth when co-breeding with more closely related versus less closely related females (GLMM, $\beta \pm SE=-0.06 \pm 0.12$, $\chi^2_1=0.23$, p=0.63; Table S5); nor was there a difference in the proportion of the emergent litter that were assigned maternity to dominant individuals (GLMM, $\beta \pm SE=-0.08 \pm 0.14$, $\chi^2_1=0.27$, p=0.60; Table S5).

Among males, by contrast with females, there was no effect of an individual's mean pairwise relatedness to dominant group members (males and females older than 3 years) on the probability of being subject to an eviction attempt (mixed-sex eviction attempts: GLMM, $\beta \pm SE=0.84 \pm 3.08$, $\chi^{2}_{1}=0.07$, p=0.79; Table S1). We found no evidence of any discrimination as to which males were targeted for eviction (Table S1). There was also no discrimination of any kind when we restricted our analysis to cases where the identity of primary aggressors was known (Table S2), or when we tested the effect of mean pairwise relatedness to same-sex and opposite-sex dominants (Table S3). Unlike the case for female-only eviction attempts, both males and females were primary aggressors in eviction attempts directed at both sexes (GLMM, $\beta \pm SE=1.80 \pm 0.56$, $\chi^{2}_{1}=12.48$, p<0.001; Figure 3B; Table S4).

Testing model predictions

Two specific predictions of the model are: (1) that selfish acts will be directed preferentially towards closer relatives only when recipients can resist; and (2) that resistance to selfish acts offered by recipients should decrease as their relatedness to the actor

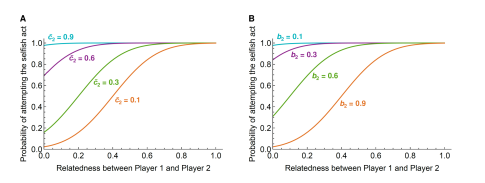


Fig. 2. Impact of model parameters on patterns of kin discrimination. Here we focus on a region of parameter space in which Player 1 should carry out the selfish act when unopposed, but in which Player 2 may do best to resist (i.e. $b_1 - c_1 - r b_2 > 0$). We show the probability that, at equilibrium, the actor attempts a selfish act, as a function of relatedness, when c_2 is drawn from a normal distribution with specified mean and standard deviation equal to 0.2. (A) Probability that a selfish act is attempted at equilibrium, assuming that $b_1 = 1$, $c_1 = 0.1$, $b_2 = 0.5$ and mean \bar{c}_2 as specified for the plotted curves. (B) Probability that a selfish act is attempted at equilibrium, assuming that $b_1 = 1$, $c_1 = 0.1$, $\bar{c}_2 = 0.5$, and b_2 is as shown in the plot.

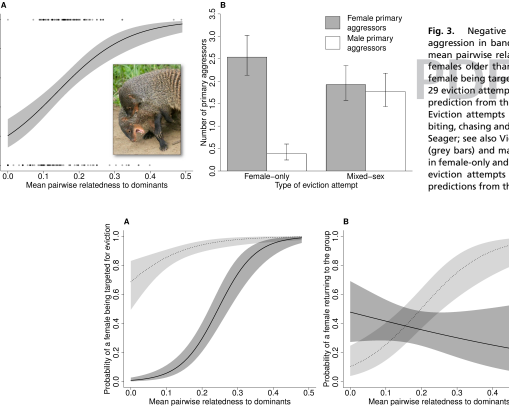


Fig. 3. Negative kin discrimination and sex-specific aggression in banded mongooses. **(A)** The effect of mean pairwise relatedness to dominants (males and females older than 3 years) on the probability of a female being targeted for eviction (*N*=207 females in 29 eviction attempts in 5 groups). The line shows the prediction from the GLMM (± standard error). **(Inset)** Eviction attempts are highly aggressive and involve biting, chasing and wrestling (photo courtesy of Dave Seager; see also Video S1). **(B)** The number of female (grey bars) and male (open bars) primary aggressors in female-only and mixed-sex eviction attempts in 6 groups). The bars show the predictions from the GLMM (± standard error).

Fig. 4. Patterns of kin discrimination towards recipients capable of offering varying levels of resistance. **(A)** In eviction attempts, dominants (males and females older than 3 years) exhibit negative kin discrimination only towards older females (*N*=207 females in 29 eviction attempts in 5 groups). The lines show the predictions from the GLMM (± standard error) for younger females (dotted line and light grey shaded area, 25th percentile of age=522 days) and older females (solid line and dark grey shaded area, 75th percentile of age=1636 days). **(B)**Following an attempt at eviction, older females who were more related to dominants (males and females older than 3 years) were less successful in regaining entry to the group (*N*=76 females in 14 eviction attempts in 4 groups); this pattern was reversed in younger females. The lines show the predictions from the GLMM (± standard error) for younger females (solid line and dark grey shaded area, 25th percentile of age=446 days) and older females (solid line and dark grey shaded area, 75th percentile of age=922 days).

increases. After we had developed the model, we tested these predictions. To test the first prediction, we examined how the effect of relatedness varied with the age and weight of potential evictees. Our reasoning was that younger or lighter females should be less able to resist eviction attempts, and that the pattern of negative kin discrimination should therefore be more pronounced when eviction is targeted at older or heavier individuals. We found that older females were indeed more likely to be targeted for eviction when more closely related to dominants, but that no such effect of relatedness was apparent for younger females (interaction

between relatedness and age: GLMM, $\beta \pm$ SE=0.008 \pm 0.004, χ^2_1 =5.98, p=0.014; Figure 4A; Table S6). The strong overall positive relationship between the probability of being targeted for eviction and relatedness was thus driven almost entirely by the pattern in older females (see Supplementary Information; Figure S2; Figure 4A). A similar interaction was also found between relatedness and weight (GLMM, $\beta \pm SE = 0.03 \pm 0.02$, $\chi^2_1 = 5.63$, p=0.018; Table S6). Consistent with prediction (1), therefore, negative kin discrimination was restricted to cases in which the

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eviction 1.0

Probability of a female being targeted for 0.0 0.2 0.4 0.6 0.8

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targets of eviction were older or heavier, and potentially more capable of offering resistance. As predicted by our model, we found no discrimination on the basis of relatedness for cases where the targets of eviction were younger or lighter, and therefore likely to suffer high costs of resistance (a high value of \bar{c}_2 ; Figure 2A).

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We found no evidence for the alternative hypothesis that the pattern of negative kin discrimination among older females arises because older, more closely related females inflict higher reproductive costs on dominants (number of emergent pups assigned to dominant female: GLMM, interaction between female cobreeder age and relatedness to dominant female, $\beta \pm SE=0.00002 \pm 0.0002$, $\chi^2_1=-0.07$, p=1.00; proportion of emergent litter assigned to dominant female: GLMM, interaction between female cobreeder age and relatedness to dominant female, $\beta \pm SE=0.00002 \pm 0.0002$, $\chi^2_1=-0.07$, p=1.00; proportion of emergent litter assigned to dominant female: GLMM, interaction between female cobreeder age and relatedness to dominant female, $\beta \pm SE=-0.00005 \pm 0.0002$, $\chi^2_1=0.08$, p=0.78; Table S5).

To test further whether negative kin discrimination depends on the capacity of recipients to resist, we examined kin discrimination in cases of infanticide of new born pups. Between one and 12 females reproduce in each breeding attempt (14) and birth is highly synchronised, with pregnant females giving birth on exactly the same morning in 63% of cases (15). Experimental and observational evidence suggest that asynchronous litters are often killed by dominant females (14, 15), and that in asynchronous litters the death of a litter in the first week after birth can be used as a proxy for infanticide (15, 16). In contrast to the pattern of negative kin discrimination in the eviction of adult females, there was no evidence of kin discrimination in cases of presumed infanticide (GLMM, $\beta \pm SE = -0.64 \pm 2.00$, $\chi^2_1 = 0.10$, p = 0.75; Table S7). This is again consistent with our model, which predicts zero or positive kin discrimination where resistance is impossible or prohibitively costly (Figure 1; Figure 2A).

440 To test the second prediction, that resistance to the selfish 441 act should decrease as the recipient's relatedness to the actor 442 increases, we examined the effects of relatedness to dominants 443 (males and females older than 3 years) and age on the probability 444 of females targeted for eviction overcoming efforts to perma-445 nently exclude them from the group. After being targeted for 446 eviction, some individuals leave the group, splitting into single-447 sex cohorts and dispersing separately in cases where males and 448 females are evicted together. Other targeted individuals, how-449 ever, actively resist eviction: they persist in following the rest of 450 the group, despite being aggressively driven away, until efforts 451 to expel them eventually cease (9). Some attempts at eviction 452 may thus be said to 'fail', largely due to sheer persistence on the 453 part of the potential evictees. We predicted, therefore, that the 454 probability of resisting eviction and regaining entry to the group 455 should decline with relatedness to dominant individuals, partic-456 ularly among older females who are potentially more capable 457 of resisting eviction. There was a significant interaction between 458 the effect of relatedness and the age of targeted females on the 459 probability of regaining entry (interaction between relatedness 460 and age: GLMM, $\beta \pm SE = -0.03 \pm 0.01$, $\chi^2_1 = 5.96$, p = 0.015; Figure 461 4B; Table S8). Among older females, individuals that were more 462 closely related to dominants were less likely to regain entry to 463 the group, although this slope was not significantly different from 464 zero (see Supplementary Information; Figure S3; Figure 4B). By 465 contrast, among younger females, more closely related targets 466 were significantly more likely to regain entry (see Supplementary 467 Information; Figure S3; Figure 4B). 468

Discussion

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We have shown in very general terms that where recipients can offer resistance, individuals can gain from targeting selfish acts at closer, rather than more distant, relatives. In the particular case of violent eviction in banded mongooses, this model may therefore explain why dominant females target closer female relatives, and why such discrimination is restricted to cases where recipients can offer most resistance. Patterns of resistance, particularly among477younger females, deviated from the model predictions, as we478discuss in more detail below. Further tests of the model could479adopt an experimental approach to manipulate resistance or the480costs and benefits of selfishness, which was not possible in our481long-term field study.482While our findings offer qualified support for the predictions483

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While our findings offer qualified support for the predictions of our model, it is important to consider alternative explanations for our results. For example, there has been much recent theoretical interest in the possibility that local competition among kin can erode selection for local helping and instead favour indiscriminate harming behaviour (16, 17). A prediction of these models is that, across groups or species, rates of aggression may be independent of relatedness (17, 18). However, these models cannot explain the targeting of closer kin for aggression when less closely related, but otherwise equivalent targets are available. A second possibility is that relatedness is correlated with some other factor influencing aggression, such as resource holding potential (RHP) or the level of reproductive competition. For example, in sea anemones, higher aggression among closer relatives has been attributed to their greater similarity in RHP (5). In banded mongooses, there is no evidence that related females are of higher RHP, or represent more of a reproductive threat. In fact, younger females (with lower RHP, and who reproduce less often) are more likely to be targeted for eviction overall, regardless of relatedness (Figure 4A). Moreover, our data show that dominant females do not suffer greater reproductive costs when they cobreed with more closely related females (Table S5). Our findings also cannot be explained as a non-adaptive side effect of selection to discriminate between species of heterospecific competitor, which has been suggested to explain negative kin discrimination in polyembryonic wasps (6).

Lastly, it has been suggested that targeting relatives for eviction could be part of an adaptive forced dispersal strategy by breeders to maximise metapopulation fitness in a structured population (13). In a previous study we did not find support for this hypothesis as a predictor of eviction at a group level (13). Moreover, it is difficult to reconcile this idea with our observations of negative kin discrimination within a given eviction attempt. In classic models such as that of Hamilton & May (19), dispersal entails direct costs for individuals who leave their natal patch, but is nevertheless favoured because it reduces local competition among kin. Because offspring value their own survival more than that of their siblings, while parents value all their offspring equally, offspring favour a lower dispersal rate than do their parents, and selection can therefore favour forced eviction of young (20-22). However, eviction by an adult of unrelated young offers equal benefits, in terms of reduced local competition, to eviction of related young, without inflicting the direct costs of dispersal on a relative. Hence, where adults can choose whom to evict, local kin competition alone cannot explain why they should preferentially target more related over less related young for expulsion. There may be other asymmetries associated with the forced dispersal of kin versus non-kin, deriving, for example, from variation in local competitive ability, or variation in the bet-hedging benefits of dispersal (23), but the effect of such variation on forced dispersal in heterogeneous groups has been little explored theoretically or empirically (13, 24). In banded mongooses, there is no evidence that closer kin compete more intensely (Table S5), or that forced dispersal of kin yields bethedging benefits.

We predicted that resistance to eviction should decrease with increasing relatedness to dominants. Contrary to this prediction, among younger females, closer relatives were more likely to regain entry to the group than older or less closely related females (Figure 4B). This result runs directly counter to our assumption that these females are weaker and less able to force 545 their way back into the group, and suggests that factors other 546 than strength or the costs of resistance may underlie the pattern 547 by which younger females return to the group. For example, 548 following eviction, dominants may voluntarily readmit closely 549 related, younger females, that would otherwise fare very badly 550 outside the group. The negotiation process by which females 551 regain entry to the group may thus be more complex than the 552 simple two-step sequence of eviction and resistance assumed by 553 our model. While our analysis shows that even a two-step game 554 can yield results that diverge from classical predictions, it is likely 555 that many negotiations in family groups may better be modelled 556 as a sequence of three or more steps, something we have not 557 attempted here. 558

Negative kin discrimination was evident only in eviction attempts of females: in males we found no relationship between relatedness and the probability of being targeted for eviction. This difference between the sexes could reflect differences in the direct fitness incentive for males and females to retain group membership, which in our model is represented by the parameter b_2 (Figure 1). Females gain greater direct fitness from group membership (i.e. higher b_2) than males because there is little or no reproductive suppression, and most females breed from the age of 10 months (11, 25). In males, by contrast, most individuals are excluded from mating by the two or three oldest males within the group (10). In our model, low values of b_2 favour little or no kin discrimination (Figure 2B). A relatively low value of b_2 in males compared to females may explain why males sometimes disperse voluntarily as a group, whereas females are invariably forced to leave after being subject to violent attack (7, 13).

574 More generally, our model shows that incorporating even 575 very simple forms of behavioural anticipation can radically 576 change the predictions of kin selection theory. Where such an-577 ticipation is possible, higher relatedness can lead to outcomes 578 that are less favourable on average for all those involved, because 579 the threat of resistance or punishment of selfish behaviour is less 580 credible between relatives than between unrelated individuals. 581 Many empirical studies have shown that animals are in fact 582 capable of adjusting their behaviour according to the anticipated 583 responses of their social partners in a range of contexts. Examples 584 include signalling systems (26), negotiation over care of offspring 585 (27), restraint in competitive growth (28), and audience effects 586 (29). However, few have considered the possibility that this kind 587 of anticipation might lead to less cooperative outcomes among 588 closer kin. An example comes from economic studies of human 589 behaviour in the context of joint-liability group lending, which 590 have found higher rates of loan default when there are more 591 relatives within a group (30, 31), leading to barring of remaining 592 group members from future borrowing. This pattern has been 593 attributed to the difficulty of group members imposing penalties 594 on relatives to enforce repayment (see 32, 33). Our results suggest 595 that similar patterns might also occur in the behaviour of other 596 species, and that the influence of kinship on aggression and co-597 operation within animal groups may be considerably more subtle 598 and variable than predicted by classical kin selection theory.

Methods

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Study population and data collection

601 Data were collected from 15 groups of banded mongooses living on the 602 Mweya Peninsula, Queen Elizabeth National Park, Uganda (0°12'S, 27°54'E), 603 between September 1997 and October 2015. For further details of habitat and climate, see (8). Groups were visited every one to three days to record group composition, life history and behavioural data. Individuals were easily 604 605 identifiable by unique shave markings on their back, and were regularly 606 trapped to maintain these markings (see (34)). On first capture a 2 mm skin sample was collected from the end of the tail for genetic analyses. For details 607 of genetic analyses and calculating relatedness see Supplementary Infor-608 mation. Individuals were trained to step onto portable electronic scales to 609 obtain weight measurements. The research was conducted with permission 610 from Uganda Wildlife Authority and Uganda National Council for Science 611 and Technology, and all methods approved by the Ethical Review Committee of the University of Exeter. 612

613 We observed the attempted eviction of 405 individuals from 8 groups in 44 eviction attempts. Eviction attempts were conspicuous, violent events and 614 easy to recognise. We defined an eviction attempt to have occurred if one 615 or more individuals left their group for at least one day following a period 616 of intense aggression towards themselves or other group members (13, 35, 617 36). In 21 out of 44 eviction attempts all targeted individuals re-joined their group; in 14 attempts some targeted individuals returned while others did 618 not; and in 9 eviction attempts all targeted individuals dispersed. 619 Statistical analyses 620

Statistical analyses were performed in R 3.3.0 (37). We used generalised linear mixed effect models (GLMM) with a binomial error structure using a logit link function, or a Poisson error structure using a log link function, in the 'Ime4' package (38). Poisson models were checked for overdispersion (39). In all analyses, we assessed the significance of each fixed effect by comparing the likelihood ratio of the maximal model to that of the model without the fixed effect (38). We present parameter estimates and standard errors from maximal models, rather than removing non-significant fixed effects due to problems associated with stepwise model reduction (40). We did, however, remove non-significant interactions to allow the significance of the main effects to be tested (41).

(i) Negative kin discrimination in banded mongooses

Models were fitted to male and female data separately because not all eviction attempts involved males. We considered adult females over 10 months old, since females younger than 10 months are unlikely to be regular breeders and are rarely evicted (7, 11, 25). We fitted whether or not a female was targeted for eviction during an eviction attempt as the binomial response variable. We included mean pairwise relatedness to dominants in the group, which we defined as males and females over 3 years of age, as a fixed effect. We use this age criterion for social dominance throughout 636 because, in both males and females, individuals over 3 years of age are more 637 likely to breed, have higher fertility, and appear to be socially dominant (7, 638 10, 14, 42). In males there is a clear age based dominance hierarchy, evident during oestrus (10, 42). In females, experimentally suppressing reproduction 639 in older females (> 3 years) results in failure of the communal litter, whereas 640 suppressing reproduction in younger females (< 3 years) does not (14). 641 Older females also breed more frequently (7, 11), and produce larger litters 642 (43). Age (days), pregnancy status (pregnant or not pregnant), mean nonpregnant weight (g) in the 60 days before the eviction attempt, mean rainfall 643 (mm) in the 30 days preceding the eviction attempt, and group size (number of individuals over 6 months) were included as additional fixed effects. 644 645 We controlled for repeated measures of individuals, eviction attempts and groups by including these terms as random intercepts, and fitted the model 646 to data on 207 females in 29 eviction attempts in 5 groups. To examine which 647 males were targeted for eviction we considered adult males over 1 year old, 648 since males under 1 year do not sire young (7, 42) and fitted whether or not 649 a male was targeted for eviction during a mixed-sex eviction attempt as the binomial response variable. We included the same fixed and random effects 650 as the female model but, instead of pregnancy status, we included the male's 651 breeding status (whether or not the male had been observed mate-guarding 652 or had sired pups in the current breeding attempt), and fitted the model to 653 data on 177 males in 15 eviction attempts in 5 groups. 654

Aggressive interactions between targeted individuals and other group members during eviction attempts were numerous, but systematic data on the identity of aggressors was difficult to record. However, since 2000 we have noted ad libitum the identity of individuals that were notably more aggressive than other group members toward potential evictees. Data on 90 of these 'primary aggressors' were available for 26 eviction attempts. To investigate the relationship between the sex of primary aggressors and the sex of individuals targeted for eviction we fitted the number of primary aggressors in each eviction attempt as the Poisson response variable and included the sex of the primary aggressors, the type of eviction attempt (female-only or mixed-sex), and the interaction between these two variables as fixed effects. We included group identity and eviction attempt as random intercepts and fitted the model to 26 eviction attempts in 6 groups. See Supplementary Information for details of analyses of negative kin discrimination by primary aggressors, to avoid inbreeding, or as a response to reproductive competition.

(ii) Testing model prediction (1): the selfish act will be directed preferentially towards closer relatives only when recipients can resist

We repeated the original analysis investigating negative kin discrimination among females in eviction attempts, but included two-way interactions between female relatedness to dominants and age (days), and female relatedness to dominants and weight (g). Other fixed and random effects were as in the original analysis. We fitted this model to data on 207 females in 29 eviction attempts in 5 groups.

673 To test further model prediction (1), we examined kin discrimination in 674 cases of infanticide of newborn pups. Infanticide is known to occur in this system, but is difficult to observe directly as it typically occurs in the den. 675 In asynchronous litters, infanticide appears to be common because early life 676 pup mortality is strongly dependent on the pregnancy status of other females 677 in the group (14, 15): pups that are born early in asynchronous litters almost 678 always die in the first few days after birth, whereas pups born last almost 679 always survive (15). Between November 1997 and October 2015, we recorded the order in which each female gave birth in an asynchronous litter (i.e. if 680

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681 she gave birth first, middle or last) and whether any of her pups survived the first week after birth. Following (15), we used the presence or absence 682 of 'babysitters' (adults left at the den to guard newly born pups) to measure 683 patterns of early life litter mortality. Specifically, to determine the survival 684 of an individual female's pups, we only considered females who gave birth early relative to the rest of the breeding females, and for which the failure of 685 a specific female's litter could be detected using cessation of babysitting (i.e. 686 we did not consider females who gave birth when babysitting of other early 687 birthing females' pups was still ongoing). A female's litter was determined to have survived the first week after birth if there were still babysitters being 688 689 left 7 days after birth, or if she was retrospectively assigned maternity to at least one emergent pup from the communal litter following genetic analysis. 690 We observed 166 females that fitted the criteria outlined above, who gave 691 birth to 120 asynchronous litters in 15 groups. Where none of the female's pups survived the first week after birth, they were assumed to have been 692 693 subject to an infanticidal attack. To test whether pups are more likely to be targeted for infanticide when mothers are less related to female dominants 694 in the group, we fitted whether or not each female's pups survived the first 695 week after birth as the binomial response variable. We included the mother's 696 mean pairwise relatedness to female dominants older than 3 years of age, mother's age (days), group size and rainfall (mm) in the 30 days before birth as fixed effects. We controlled for repeated measures of mothers, litters and 697 698 groups by including these terms as random intercepts, and fitted the model 699 to data on 59 females giving birth to 52 communal litters in 12 groups. 700

(iii) Testing model prediction (2): resistance to the selfish act offered by recipients should decrease as their relatedness to the actor increases

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We fitted whether or not a female over 10 months old that was targeted for eviction overcame efforts to permanently exclude them from the group as

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the binomial response variable. We only considered eviction attempts where some evictees were allowed to return and others were not due to problems with fitting a binomial model to outcomes that are exclusively successes or failures. We included mean pairwise relatedness to dominants in the group, age (days), and the interaction between these two variables as fixed effects. We were unable to include additional fixed effects in the model due to problems with model convergence. We controlled for repeated measures of individuals, eviction attempts and groups by including these terms as random intercepts, and fitted the model to data on 46 females in 14 eviction attempts in 4 groups.

Author contributions

F.J.T., M.A.C. and R.A.J. conceived the study. R.A.J. wrote the theoretical model. F.J.T., M.A.C. and R.A.J. designed the analyses. F.J.T., H.H.M., J.L.S., E.I.K.V., J.S.G., M.B.V.B. and S.J.H. collected data. F.J.T. analysed data. J.L.S. and H.J.N. carried out genetic analyses. F.J.T., M.A.C. and R.A.J. drafted the paper. All authors contributed to the final version of the manuscript.

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