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The influence of stress hormones and aggression on cooperative behaviour in subordinate meerkats

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Abstract: In cooperative breeders, aggression from dominant breeders directed at subordinates may raise subordinate stress hormone (glucocorticoid) concentrations. This may benefit dominants by suppressing subordinate reproduction but it is uncertain whether aggression from dominants can elevate subordinate cooperative behaviour, or how resulting changes in subordinate glucocorticoid concentrations affect their cooperative behaviour. We show here that the effects of manipulating glucocorticoid concentrations in wild meerkats (*Suricata suricatta*) varied between cooperative activities as well as between the sexes. Subordinates of both sexes that were treated with a glucocorticoid receptor antagonist (mifepristone) exhibited significantly more pup protection behaviour ('babysitting') compared to those treated with glucocorticoids (cortisol) or controls. Females treated with mifepristone had a higher probability of exhibiting pup food provisioning ('pupfeeding') compared to those treated with cortisol. In males, there were no treatment effects on the probability of pupfeeding, but those treated with cortisol fed a higher proportion of the food they found to pups than those treated with mifepristone. We also used 19 years of behavioural data to show that dominant females did not increase the frequency with which they directed aggression at subordinates at times when their need for assistance was highest. Our results suggest that it is unlikely that dominant females manipulate the cooperative behavior of subordinates through the effects of aggression on their glucocorticoid levels and that the function of aggression directed at subordinates is probably to reduce the probability they will breed.

Keywords: Aggression, Behavioural plasticity, Cooperation, Glucocorticoids, Sociality, Stress

1. Introduction

In mammals that breed cooperatively, where group members raise young produced by dominant breeders, social rank can influence the stress-related hormone (glucocorticoid) concentrations of group members (1-3). In some species, dominant individuals direct aggression towards subordinates, raising their glucocorticoid concentrations (GCs: 4-7). The elevation of GCs in subordinates can suppress their reproductive activity (4-6,8) allowing dominants to monopolize reproductive opportunities (6). However, such changes in subordinate GCs could also carry additional benefits or costs for dominants by increasing or decreasing subordinate cooperative behaviour, respectively (9-16).

Studies of a diversity of taxa already suggest that changes in GCs can alter the expression of social behaviour. Studies in highly social wild animals are rare, but some studies have identified positive or negative associations between GCs and cooperative behaviour (12-15). In social mammals including humans, elevated GCs are associated with decreased expression of different social behaviours (17-19) and reductions in the formation of social attachments between parents and offspring (20-21) or two opposite-sex conspecifics (23). The lack of congruency in the direction of relationships between GCs and these different types of social behaviours is not surprising given the observational nature of these previous studies. Experimental studies are therefore needed to determine whether changes in GCs cause variation in cooperation or whether correlations between GCs and cooperation (12-15) occur because increased energetic expenditure on cooperative activities raises respective GCs.

Fluctuations in GCs can cause changes in behaviour by binding to central and peripheral glucocorticoid (GRs) or mineralocorticoid (MRs) receptors, subsequently altering the activity of those receptors that in turn regulate gene transcription, or through non-genomic effects that do

not involve GRs or MRs (23-25). However, there is at least some evidence that the effects of GCs on social behaviours are mediated by changes in the activity of GRs. For instance, mifepristone is a selective GR antagonist that can lower the expression of behaviours influenced by GCs such as reducing the expression of anti-social behaviours in humans (social withdrawal behaviour) that are associated with major depression, anxiety, or post-traumatic stress (26). How exactly changes in GCs or their genomic (binding to MRs or GRs) or non-genomic consequences affect behaviour is an active area of research (27). Previous studies suggest the possibility that increased GCs can reduce social behaviour by affecting the activity of neural circuits in the mesolimbic pathway such as promoting fear of conspecifics (28) or altering the reward value of social interactions (29-30). Regardless of the mechanism, a growing number of studies suggest that changes in GCs can alter the expression of a variety of social behaviours.

Although there is evidence that aggression directed at subordinates can increase their GCs (7) and reduce the probability that they will breed (6), how changes in subordinate GCs affect their cooperative behavior has rarely been investigated (9-16). Here, we describe the results of manipulations of circulating GC concentrations and GR activity in wild meerkats to determine if they cause changes in alloparental care behaviours (babysitting and pupfeeding) as well as analyses of the distribution of aggression directed by dominant females at subordinates. We provisioned subordinates of both sexes with either exogenous GCs (cortisol), a GR antagonist (mifepristone), or an oil vehicle (controls) for 10 consecutive days during either babysitting or pupfeeding periods. The goal of these treatments was to produce a range of variation in GR activity with those treated with cortisol predicted to have the highest, those treated with mifepristone having the lowest (because mifepristone antagonizes the GR), and the controls in between cortisol and mifepristone treatments. We confirmed that mifepristone was in

fact antagonizing the GR (see Fig. S1). To investigate whether variation in rates of aggression directed by dominant females at subordinates were related to their need for increased assistance, we used 19 years of long-term behavioural data.

Meerkats are obligate cooperative breeders that live in social groups containing 2-47 individuals (31), many of whom are close relatives (32). Each group contains a monogamous breeding pair who are socially dominant to all same-sex subordinates in their group (33-37) and are the parents of ~90% of all offspring produced within the group (33-34). Subordinate females but not males are known to be infanticidal especially if the former are pregnant (38, 39). Probably as a consequence, subordinate females, but not males, are usually evicted from their group when the dominant female is pregnant (6, 38). Aggression from dominant females can alter subordinate GCs. Subordinates that receive more aggression from the dominant female tend to have higher plasma GCs (7). Moreover, dominant females are more aggressive towards subordinate females when they are pregnant (6, 40) and, subordinate females (regardless of their pregnancy status), but not males, have higher GCs when the dominant female is pregnant (7).

Subordinates of both sexes provide alloparental care to offspring produced by dominants (35,36,41). During their first month of life, meerkat pups stay at their natal burrow with an older subordinate ('babysitter': 36) while the rest of the social group goes foraging away from the burrow for the entire day. From 1 to 3 months of age while the pups are foraging with the group, they are provisioned with food items that are found by older subordinates and dominant breeders in their social group (42). After around 3 months, meerkats obtain most of their food by foraging independently (43).

Previous correlative studies of the relationship between GCs and cooperative behaviour in subordinate male meerkats showed that elevated GCs were positively related to some forms of

cooperative behaviour and negatively with others. Specifically, they showed that increased GCs in males were positively associated with pupfeeding (12) but negatively associated with babysitting (15). Our initial expectation was consequently that subordinates of both sexes treated with cortisol (presumably those with the highest GR activity) would have the lowest levels of babysitting, but the highest levels of pupfeeding. Secondly, we predicted that subordinates of both sexes treated with mifepristone (presumably those with the lowest GR activity) would have the highest levels of babysitting (15), but lowest levels of pupfeeding (12).

2. Methods

(A) Documenting characteristics of social groups

We studied habituated meerkats at the Kuruman River Reserve (26° 58' S, 21° 49' E) in the Northern Cape, South Africa from 1998-2016. Individuals were uniquely and permanently marked with microchips (Identipet[®], Johannesburg, South Africa) and small dye marks so that they could be identified visually. Groups were visited for 4-8 hours per day once every two to three days throughout the year to collect *ad libitum* or focal behavioural observations during which we recorded the identity of all meerkats present in the group to quantify group size and the sex ratio of the group. We tracked the identity of the dominant female and male in each group (dominance is determined via behavioural observations, see supplementary material). Pregnancy status of all females was determined by noting steady mass gain and visible swelling of the abdomen and nipples. Parturition was identified by changes in the appearance of females, dramatic overnight mass loss, and the presence of subordinate individuals babysitting (44) at the sleeping burrow while the rest of the group went foraging. Presence of pups <90 days of age was determined either by direct observation or by the presence of babysitters at the burrow.

(b) Manipulating subordinate glucocorticoid (cortisol) concentrations

We provisioned subordinate meerkats with either cortisol (10 mg/kg, hydrocortisone, Sigma H4126), mifepristone (40 mg/kg, Sigma M8046), or oil vehicle (100% coconut oil). Dosages were chosen based upon previous studies and our own pilot experiments (see supplementary materials). We confirmed that our cortisol treatments significantly elevated plasma GCs (cortisol treatment: Fig. S2) and faecal glucocorticoid metabolites (Fig. S3) within a biologically relevant range and mifepristone altered the activity of the GR (Fig. S1, S4), and that these treatments would influence the behaviour of meerkats (Figs. S1, S4) without causing abnormal behavioural changes (Fig. S5). Treatments (cortisol, mifepristone, control) were randomly assigned to three adult subordinates (>12 months of age) within the same group that were the same-sex and of the same or similar age. When these conditions could not be met, we used as a subject another adult subordinate of the opposite sex but similar age. None of the subordinate females were pregnant during treatment (pregnancy status determined as described above). Subjects were first treated continuously for a 10-day period immediately following the birth of the litter (babysitting period) and then again for a second 10-day period (with a different treatment) during the peak pupfeeding period (42, mean age of pups during treatment = 45.3 d, range = 39-60 d). Subjects who consumed <50% of their treatments that they were provisioned with (n=4) were excluded from our analyses about the treatment effects on babysitting. However, in our analyses on pupfeeding, we included a covariate for whether the focal observations occurred on the same day of provisioning with the treatment or not (all individuals had consumed their treatment that day, see statistical analyses below).

(c) Body mass and foraging success

We measured subordinate body mass and foraging success to assess their effects on subordinate cooperative behaviour and to assess the influence of our treatments on foraging success (see supplementary material). Meerkats were weighed (to the nearest gram) in the morning immediately after emergence from their sleeping burrow, but before foraging had commenced, after 1-4 hours of foraging, and again in the evening after foraging was completed but before they entered their sleeping burrow (45, 46). Birth dates of meerkats were known so we calculated age-corrected body mass as a measure of body condition. We controlled for short-term fluctuations in body mass by averaging the morning body mass of individuals over the 30 days prior to capture or behavioural observations and then used the residuals from a general linear model (response variable was average morning body mass, predictor variable was age in days) to estimate age-corrected body mass. We estimated daily foraging success as the evening body mass minus the morning body mass within the same day and took the average daily foraging success over the previous 30 days prior to capture. We estimated mass-corrected foraging success as the residuals from a general linear model (response variable was daily foraging success, predictor variable was morning body mass).

(d) Quantifying aggression that subordinates received from dominants

We used *ad libitum* sampling (47) to quantify how much aggression subordinates received from dominants. During each visit to the group, we recorded all dominance assertions exhibited by dominant females towards subordinates. Dominance assertions that were collected include aggressive behaviours or signs of dominance exhibited by the dominant female towards subordinates such as charging, chasing, or hitting them (see 40, 48). The total number of dominance assertions was summed for each day of behavioural observations for each individual subordinate and the total amount of time of the session was recorded. In total, we examined the

distribution of variation in rates of aggression directed by dominant females at subordinate group members using >110,483 hours of behavioural data collected over 19 years that provided us with rates of aggression by 98 dominant females towards 1520 subordinates (713 females, 807 males) in 40 different groups. From this same dataset, we also had observations on all the babysitting and pupfeeding contributions of subordinates for each litter that survived to emergence from the natal burrow.

(e) Measuring babysitting behaviour

We measured the relative contributions of each subordinate male or female to babysitting, as we have done previously (36,41,49). When pups were born but not yet foraging with the group, we visited the natal burrow every day in the morning to record the identity of the babysitter (36, 49). Relative babysitting contributions of each individual for each litter was estimated by dividing the total number of days an individual babysat a litter over the total number of days that this specific litter had a babysitter. If a babysitter was replaced during morning observations, they were both assigned 0.5 d of babysitting. For the subordinates receiving treatments, we measured relative babysitting contributions of the three treated subordinates only during the treatment period. For the long-term correlative data analyses, the proportion of babysitting for a litter relative to all other individuals was used.

(f) Measuring pupfeeding behaviour

Pupfeeding in meerkats is a highly conspicuous event as pups emit distinctive begging vocalisations and subordinates engage in obvious behaviours where they bring prey items to pups (43). To assess how our experimental treatments affected pupfeeding behaviour in treated subordinate meerkats, we conducted 20 min continuous behavioural focal observation sessions on treated subordinates. Focal observations began in the morning after the group had left their

sleeping burrow and were foraging for at least 10-15 min. Focals were alternated among the treated subjects (first subject focalled was randomly selected) and focals on the same individual were not consecutive and were separated by 20-30 min. During each focal, we recorded all food items found, their size (tiny, small, medium, large, extra-large: 50), and the number and size of prey items that were found during the focal and fed to pups. Prey biomass was estimated as previously (50). This allowed us to assess the effects of our experimental treatments on 1) whether the subordinates exhibited pupfeeding at all during the focal session (probability of pupfeeding) and 2) the total proportion of prey biomass that was found during the focal session that was fed to pups that were foraging with the group (“generosity”). We analysed the effects of our treatments on both of these behaviours because they be different measures where the probability of pupfeeding reflects motivation to exhibit this type of alloparental care and generosity controls for its condition-dependence (35).

Pupfeeding contributions for each subordinate in our long-term data analyses were recorded using *ad libitum* sampling when there were pups (up to 90 d of age) foraging with the group. In these *ad libitum* observations, we did not record how much food each subordinate found, so we used the probability of pupfeeding as a response variable.

(h) Statistical analyses

We used an information-theoretic approach (Akaike information criterion corrected for small sample sizes, AIC_c: 51) to examine the degree of support among different models that contained a range of covariates (described in supplementary materials and results in Tables S1-S6). Our approach was to develop a list of candidate models, select the model containing biological predictor variables with the lowest AIC_c, and then evaluate the significance of each of the predictor variables using traditional null hypothesis significance testing. Further details of

model selection procedures, use of null models, and additional model averaging results are shown in supplementary materials.

We first used generalized linear mixed-effects model (GLMM) with binomial errors to examine whether our experimental treatments affected babysitting, the probability that pupfeeding occurred during our focal observations, and the proportion of biomass that was found by subordinates and subsequently fed to pups foraging with the group. We included random intercept terms for individual and litter identity as well as observer in both the pupfeeding models and a random intercept term for litter identity in the babysitting models because of repeated observations. Our model selection procedures revealed that the best model (with lowest AIC_c shown in Tables S1-S3) for babysitting and pupfeeding was the null model that did not contain any biological predictor variables. Because we were interested in the effects of the experimental treatments on babysitting and pupfeeding, we only report results from models (Tables S1-S3) containing biological predictor variables with the lowest AIC_c (model in bold face font in Tables S1-S3) in the main text and use these results for our interpretations. Because a second model containing biological predictor variables (i.e., not the null model) was within $\Delta AIC_c < 2$ in Tables S1-S3, we also include results from model averaging (53) from these top models in Tables S10-S12 (using the zero method: 90). However, our interpretations are focused on the one model with the lowest AIC_c containing biological predictor variables.

We next examined potential causes of variation in the amount of aggression by dominant females directed at subordinate meerkats (all models in Table S4, model with lowest AIC_c in Table S7). Aggression from the dominant female directed at subordinates was infrequently observed so we conducted a GLMM with a Poisson error structure that contained random intercept terms for individual, group, identity of the dominant female, and year. Because of

variation in the amount of time dominant females were observed, we included an offset for the total time each meerkat group was observed.

Finally, we examined associations between the amount of aggression subordinates received from dominant females and their 1) babysitting contributions (Table S5) and 2) the probability that pupfeeding occurred during our *ad libitum* behavioural observations (Table S6) using a correlative approach. Models assessing how aggression received from the dominant female affected subordinate babysitting contributions (Table S5) or the probability of pupfeeding (Table S6) are in the supplementary materials and we only present models with the lowest AIC_c (Tables S8-S9). The response variable in these models was either the relative proportion of babysitting contributions provided by that individual or a binary response variable to indicate whether pupfeeding did or did not occur in the behavioural observation sessions. All these models were GLMMs (binomial error structure) that contained random intercept terms for individual, identity of the litter and dominant female, and year because of repeated samples on the same individuals, litters, dominant females exhibiting the aggression, or within different years. The pupfeeding model included an offset for time we observed the group.

All analyses were conducted in R (version 3.21: 52). AIC_c was calculated using maximum likelihood (R package MuMin, version 1.15-1: 53). We used a graphical approach to confirm the normality and homoscedasticity of the residuals from the models. All continuous variables were standardized (mean of 0, SD of 0.5: 54). All GLMMs were run with the package lme4 (version 1.1-12: 55) in R. There was no or limited evidence of overdispersion in our GLMMs (dispersion parameters ranged from 0.7 to 1.08). P-values were generated using the R package lmerTest (version 2.0-3: 56). There was little evidence of strong collinearity among the covariates (variance inflation factors were consistently <2: 57) except for terms where we

included both a linear and second order term in the model or for covariates that were included in an interaction term with others. Unless otherwise indicated, below we present mean \pm SD.

3. Results

Effects of treatments on cooperative behaviour

Contrary to our prediction, treating subordinates with cortisol (n=9), which should increase GR activity, did not affect their contributions to babysitting relative to those of controls in either sex (n=8, $z=-0.2$, $P=0.84$, Fig. 1A). However, as predicted, mifepristone treatment (which should decrease GR activity), caused an increase in babysitting in both sexes: subordinate females and males treated with mifepristone (n=9) provided 64% more babysitting than did those treated with cortisol ($z=2.6$, $P=0.009$) and 55% more than the controls ($z= 2.34$, $P=0.019$, Fig. 1A). Model selection indicated that there was no evidence that the effects of the treatments on babysitting were sex-specific (Table S1).

Cortisol and mifepristone treatments both affected the frequency of pupfeeding (the probability that subordinates would feed pups during 20 min focal observations) though their effects differed between the sexes (Table S2). Contrary to our initial predictions, subordinate females treated with cortisol were significantly less likely to exhibit pupfeeding (1200 min observation on 10 females) compared to those treated with mifepristone (780 min observation on 8 females, $z=2.02$, $P=0.044$), but did not differ from control females (800 min observation on 7 females, $z=0.26$, $P=0.79$, Fig. 1B). Subordinate males treated with cortisol were not more likely to feed food items they found to pups (400 min observation on 4 males) compared to males treated with mifepristone (860 min observation on 7 males, $z=-1.59$, $P=0.11$) and also did not

differ from the controls (560 min observation on 7 males, $z=-1.01$, $P=0.31$, Fig. 1B). Females and males treated with mifepristone were equally likely to exhibit pupfeeding during the focals compared to the controls (females: $z=0.48$, $P=0.63$; males: $z=-1.63$, $P=0.1$, Fig. 1B)

The effects of cortisol and mifepristone on a subordinate's 'generosity' towards pups (the proportion of total food biomass found that they then fed to pups) also differed between the sexes (Table S3). Subordinate females treated with cortisol were not more generous compared to females treated with mifepristone ($z=-0.97$, $P=0.33$) or compared with the controls ($z=-0.83$, $P=0.41$, Fig. 1C). However, subordinate males treated with cortisol were more generous compared to those treated with mifepristone ($z=-2.41$, $P=0.016$), but did not differ from the controls ($z=-1.66$, $P=0.096$, Fig. 1C). Females and males treated with mifepristone and the controls did not differ in their generosity (females: $z=-0.13$, $P=0.89$; males: $z=0.92$, $P=0.36$, Fig. 1C).

The distribution and effects of aggression from dominant females

Our experimental results showed that mifepristone (which should decrease GR activity) elevated babysitting (Fig 1A) so, if dominant females adjust the frequency with which they direct aggression at subordinates to maximize the contributions of subordinates to babysitting, they would be expected to reduce the amount of aggression directed at subordinates during babysitting to lower their GCs and consequently their GR activity. Our long-term data analyses showed that dominant females directed significantly less aggression at subordinate females during babysitting compared to times when dominant females were pregnant and there were no pups in the group ($z=-3.06$, $P=0.0022$). However, rates of aggression directed at subordinate males did not vary in the same way ($z=0.18$, $P=0.86$, Table S7, Fig. 2), suggesting that that this difference was not the result of an attempt by dominant females to increase contributions to

subordinates to babysitting. In addition, if aggression is used by dominants to stimulate babysitting in subordinates, dominant females might also be expected to reduce rates of aggression directed at individuals that were infrequent babysitters to a greater extent than those directed at more frequent babysitters. However, the amount of aggression subordinates received from dominant females was also not associated with their contributions to babysitting in either sex (females: $z=-0.08$, $P=0.93$; males: $z=-0.23$, $P=0.82$, Table S8).

There was also no indication that dominant females adjusted their aggressive behaviour to stimulate pupfeeding by subordinates. Because our experimental results showed that mifepristone treatment increased pupfeeding frequency in females but decreased pupfeeding generosity in males compared to those treated with cortisol (Fig. 1C), dominant females that were using aggression to stimulate pupfeeding by subordinates should then have reduced the amount of aggression directed at subordinate females during pupfeeding but increased aggression directed at subordinate males. Although the amount of aggression dominant females directed at subordinate females was lower during pupfeeding compared to when the dominant female was pregnant with no pups in the group ($z=2.01$, $P=0.044$), it tended to be higher during pupfeeding compared to when subordinate females were babysitting (females: $z=-1.84$, $P=0.066$, Table S7, Fig. 2). Furthermore, during pupfeeding the amount of aggression dominant females directed at subordinate males was not higher compared to periods when they were in groups where the dominant female was pregnant with no pups in the group ($z=0.5$, $P=0.61$) or when they were babysitting ($z=0.11$, $P=0.91$, Table S7, Fig. 2). Finally, the amount of aggression received from dominant females by subordinates was not associated with the probability of pupfeeding in either sex (females: $z=-0.26$, $P=0.8$; males: $z=1.53$, $P=0.12$, Table S9).

An alternative interpretation of the distribution of aggression is that the frequency of aggression directed at subordinates by dominant females is adjusted principally to controlling the risk of infanticide and the frequency that subordinates will breed (6) and our analyses of the distribution of aggression is consistent with this. Subordinate females received significantly more aggression from the dominant female (females: $z=6.17$, $P<0.0001$, Table S7) when the dominant female was pregnant compared to when she was not pregnant with no pups in the group (Fig. 2). No similar changes in the amount of aggression received occurred in males ($z=1.52$, $P=0.13$, Table S7, Fig. 2) supporting the suggestion that the increase in aggression directed at subordinate females was associated with reproductive suppression. In addition, older and heavier subordinate females (age: $z=12.19$, $P<0.0001$; mass: $z=9.5$, $P<0.0001$, Table S7) and males (age: $z=6.71$, $P<0.0001$; mass: $z=5.03$, $P<0.0001$, Table S7) received more aggression from dominant females than those that were younger or lighter and the effect of age (sex \times age, $z=8.9$, $P<0.0001$) and body mass (sex \times body mass, $z=1.9$, $P=0.055$, Table S7) on aggression received was more pronounced for females than males.

4. Discussion

Our experimental results show that variation in GCs and likely GR activity can influence how much alloparental behaviour subordinates exhibit. Surprisingly, these treatment effects differed both between the sexes and between different forms of cooperative behaviour. Mifepristone (which should decrease GR activity) increased babysitting in subordinates of both sexes, but only elevated pupfeeding behaviour (probability of occurring) in females compared to those treated with cortisol (which should elevate GR activity). Cortisol decreased pupfeeding (probability of occurring) in females but it enhanced pupfeeding behaviour (generosity) in males.

Overall, this suggests that elevated GRs and perhaps increased GR activity reduces alloparental care in meerkats except in the case of pupfeeding by subordinate males.

Our results suggest that GCs and the activity of the GR may act as a general mechanism mediating behavioural plasticity in cooperative behaviour across cooperatively breeding species. Factors such as poor body condition (58), harsh environmental conditions (59), or low levels of relatedness (61, 62) are associated with decreases in the expression of cooperative behaviour presumably because under such conditions the ratio of costs to benefits of cooperation is increased. Similarly, GCs are highly responsive to fluctuations in the same intrinsic (body condition) and extrinsic (weather, group size) factors (35, 36, 49, 62). Increases in GCs from such intrinsic or extrinsic factors that affect GR activity (or MR activity or have non-genomic effects) could in part trigger these decreases in cooperative behaviour.

We found no evidence to support the hypothesis that dominant females use aggression to elevate the cooperative behaviour of subordinates. The amount of aggression subordinates of both sexes received from the dominant female was not associated with their contributions to babysitting or pupfeeding in our analyses of long-term behavioural data. This matches a previous study in meerkats showing that dominant females do not increase the amount of aggression they exhibit towards subordinates when the need for their help was experimentally increased (63).

Instead of using aggression to stimulate cooperative behaviour, our results are aligned with those of a previous study in meerkats (6) suggesting that dominant females use aggression to control reproduction in subordinate females. Subordinate females, but not males, are often infanticidal and pregnant subordinates are more likely to commit infanticide compared to when they are not pregnant (6,38,39). Subordinate females but not males received more aggression from the dominant female (this study) and had higher plasma GCs (7) when the dominant female

was pregnant. Increased GCs in subordinate females may reduce the probability of them committing infanticide by suppressing their own reproduction or causing abortions (6). Consequently, antagonistic interactions of social subordinates with dominant breeders and the consequent changes in their GCs likely play a role in the reproductive suppression of same-sex subordinates.

Our results suggest that the effects of GCs and perhaps GR activity on cooperative behaviour have different consequences in males and females. Previous studies of alloparental care in meerkats (11), African striped mice (*Rhabdomys pumilio*: 13) and the formation of pair-bonds in prairie voles (*Microtus ochrogaster*: 22) indicate that in males of these species, increased GCs are associated with increased cooperative behaviour (but see 14). We found that male meerkats treated with cortisol (presumably resulting in higher GR activity) were more generous when pupfeeding. This result is similar to a previous observational study in male meerkats (12) but differs from a previous short-term study in meerkats that also experimentally increased GCs (using injections of exogenous GCs) and observed no substantial changes in pupfeeding behaviour (64). This difference may be due to the increases in plasma GCs in the previous study that were higher than the increases in subordinate plasma GCs we induced in this study (Fig. S2) or because this previous study used manipulations that operated over a shorter timescale.

In contrast, studies of females of the same species as those described above suggest a negative association between GCs and perhaps GR activity and cooperative or social behaviour. Increases in GCs in female prairie voles (22) and African striped mice (13) are associated with reduced cooperative behaviour, suggesting that reduced GR activity elevates cooperation in females. We also found that subordinate female meerkats treated with mifepristone (presumably

with lower GR activity) exhibited more babysitting and pupfeeding behaviour. These studies indicate that, during the co-evolution of hormones and social or cooperative behaviours, selection has often favored sex-specific effects, raising important questions about the function and evolution of these differences that have yet to be investigated. One plausible explanation is that the fitness benefits of alloparental behaviour under stressful environments differ between the sexes. This can occur if there is sex-biased dispersal from the natal group such as in meerkats where there is male-biased dispersal (65) and the average relatedness between subordinate females and dominant females (and consequently the offspring produced in the group) is greater than in males who immigrate into the group (66). The indirect fitness benefits of staying and helping within the natal group for the sex that is more closely related to the dominant breeders may be greater than emigrating, especially when that sex also queued for the dominant breeding position. We predict that under these conditions, this sex should exhibit higher cooperative behaviour when GCs are elevated. However, our results reject this hypothesis because we did not that subordinate females exhibited higher cooperative behavior when their GCs were increased.

There are numerous possible mechanisms by which changes in GCs or GR activity can alter behaviour in a sex-specific fashion given the well-documented sex differences in the functioning of the vertebrate neuroendocrine stress axis (67,68). One hypothesis that could explain our results is that there are sex differences in the central distribution of GRs or receptors for other components involved the neuroendocrine stress axis (69) that in turn modify how GCs alter cooperative behaviour. For example, increased production of corticotropin-releasing factor (CRF) in the mesolimbic system due to elevated GCs could promote fear and anxiety of conspecifics thereby reducing social behaviour (27,28). CRF receptors in females may be hypersensitive compared to males (68) such that there is a lower threshold in females than males

at which GCs cause a reduction in social behaviour. There are many other possibilities to explain our results given the range of potential mechanisms that contribute to sex differences in the neuroendocrine stress axis (67,68). Clearly, the mechanisms underlying these sex differences is an area that deserves much more study. Future field based studies can contribute to this area by examining whether changes in social or environmental factors (group size, weather patterns, food availability) or individual-state (body mass) cause sex-specific changes in the expression of cooperative behaviour.

Our results have general implications for our understanding both of the causes of variation in the expression of cooperative behaviour in cooperatively breeding species and of the possible mechanisms underlying variation in social behaviour across taxa. First, our results reject the hypothesis that dominant breeders in cooperatively breeding species could manipulate the cooperative behaviour of subordinates by strategically increasing or decreasing their GCs and/or affect their GR activity. In both meerkats (63) and in other cooperatively breeding species (16), there is little evidence that dominant breeders use punishment or changes in GCs to alter the cooperative behaviour of subordinates.

Our research adds to the growing number of studies showing that changes in GCs or GR activity can alter the expression of different types of social behaviour ranging from social attachments formed between two opposite-sex individuals (22) or parents and offspring (20-21), antisocial behaviour in humans (17-18), or, in our study and others (12-15), alloparental behaviour. They provide new research avenues by showing that the effects of changes in GCs or GR activity on cooperative behaviour differs not only between the sexes but also between two different types of cooperative behaviour. Why they should vary in this way is still unknown, nor

is it clear whether similar differences in the effects of GCs and GR activity occur in other cooperative breeders.

Ethics: All protocols used in our experiments were approved by the Animal Ethics Committee at the University of Pretoria (Pretoria, South Africa) and the Northern Cape Conservation Authority (South Africa).

Date, code and materials: All data used in our analyses will be archived on Data Dryad.

Competing interests: We have no competing interests.

Authors' Contributions: B.D. designed experiments, T.H.C-B., M.B.M., D.G., B.D., and C.D. coordinated long-term data collection, B.D. T.H.C-B, M.B.M., D.G., I.BG., N.B., M.H., A.G., C.D., and H.C.S-J collected data, B.D. conducted analyses and produced figures, B.D. and T.H.C-B authored manuscript with contributions from M.B.M., D.G., I.BG., N.B., M.H., A.G, and C.D.

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References:

- 1) Creel S. 2001. Social dominance and stress hormones. *Trends Ecol. Evol.* **16**, 491-497.
(doi:10.1016/S0169-5347(01)02227-3)
- 2) Creel S, Dantzer B, Goymann W, Rubenstein DR. 2013. The ecology of stress: effects of the social environment. *Func. Ecol.* **27**, 66-80. (doi: 10.1111/j.1365-2435.2012.02029.x)
- 3) Goymann W, Wingfield JC. 2004. Allostatic load, social status and stress hormones: the costs of social status matter. *Anim. Behav.* **67**, 591-602. (doi:10.1016/j.anbehav.2003.08.007)
- 4) Abbott DH, Keverne EB, Bercovitch FB, Shively CA, Mendoza SP, Saltzman W, Snowdon, CT, Ziegler TE, Banjevic M, Garland T Jr. et al. 2003. Are subordinates always stressed? A comparative analysis of rank differences in cortisol levels among primates. *Horm. Behav.* **43**, 67-82. (doi: 10.1016/S0018-506X(02)00037-5)
- 5) Hackländer K, Möstl E, Arnold W. 2003. Reproductive suppression in female alpine marmots, *Marmota marmot*. *Anim. Behav.* **65**, 1133-1140. (doi: 10.1006/anbe.2003.2159)
- 6) Young, AJ, Carlson AA, Monfort SL, Russell AF, Bennett NC, Clutton-Brock T. 2006. Stress and the suppression of subordinate reproduction in cooperatively breeding meerkats. *Proc. Natl. Acad. Sci. USA* **103**, 12005-12010. (doi:10.1073/pnas.0510038103)
- 7) Dantzer B, Bennett NC, Clutton-Brock T. 2017. Social conflict and costs of cooperation in meerkats are reflected in measures of stress hormones. *Behav. Ecol.* **28**, 1131-1141.
(doi:10.1093/beheco/ax077)
- 8) Sanderson JL, Nichols HJ, Marshall HH, Vitikainen EIK, Thompson FJ, Walker SL, Cant

- MA, Young AJ. 2015. Elevated glucocorticoid concentrations during gestation predict reduced reproductive success in subordinate female banded mongooses. *Biol. Lett.* **11**, 20150620. (doi:10.1098/rsbl.2015.0620)
- 9) Reeve HK, Sherman PW. Intra-colonial aggression and nepotism by the breeding female naked mole-rat. In *The Biology of the Naked Mole-rat* (eds PW Sherman, JUM Jarvis, RD Alexander), pp. 337-357. Princeton, NJ, USA: Princeton University Press.
- 10) Mulder RA, Langmore NE. 1993. Dominant males punish helpers for temporary defection in superb fairy-wrens. *Anim. Behav.* **45**, 830-833. (doi:10.1006/anbe.1993.1100)
- 11) Balshine-Earn S, Neat FC, Reid H, Taborsky M. 1998. Paying to stay or paying to breed? Field evidence for direct benefits of helping behavior in a cooperatively breeding fish. *Behav. Ecol.* **9**, 432-438. (doi: 10.1093/beheco/9.5.432).
- 12) Carlson AA, Manser MB, Young AJ, Russell AF, Jordan NR, McNeilly AS, Clutton-Brock T. 2006. Cortisol levels are positively associated with pup-feeding rates in male meerkats. *Proc. R. Soc. Lond. B* **273**, 571-577. (doi: 10.1098/rspb.2005.3087)
- 13) Raynaud J, Schradin C. 2015. Corticosterone levels correlate with alloparental care in a sex-dependent manner in African striped mice, *Rhabdomys pumilio*. *Ethology* **121**, 57-67. (doi:10.1111/eth.12317)
- 14) Sanderson JL, Young AJ, Hodge SJ, Kyabulima S, Walker SL, Cant MA. 2014. Hormonal mediation of a carry-over effect in a wild cooperative mammal. *Func. Ecol.* **28**, 1377-1386. (doi:10.1111/1365-2435.12307)
- 15) Carlson AA, Russell AF, Young AJ, Jordan NR, McNeilly AS, Parlow AF,

- Clutton-Brock T. 2006. Elevated prolactin levels immediately precede decisions to babysit by male meerkat helpers. *Horm. Behav.* **50**, 94-100.
(doi:10.1016/j.yhbeh.2006.01.009)
- 16) Riehl C, Frederickson ME. 2016. Cheating and punishment in cooperative animal societies. *Phil Trans. R. Soc. Lond. B* **371**, 20150090. (doi:10.1098/rstb.2015.0090)
- 17) Kathol RG, Anton R, Noyes R, Gehris T. 1989. Direct comparison of urinary free cortisol excretion in patients with depression and panic disorder. *Biol. Psych.* **25**, 873-878. (doi: doi.org/10.1016/0006-3223(89)90267-9)
- 18) Korte SM, Koolhaas JM, Wingfield JC, McEwen BS. 2005. The Darwinian concept of stress: benefits of allostasis and costs of allostatic load and the trade-offs in health and disease. *Neurosci. Biobehav. Rev.* **29**, 3-38. (doi: [10.1016/j.neubiorev.2004.08.009](https://doi.org/10.1016/j.neubiorev.2004.08.009))
- 19) van Goozen SH, Fairchild G. 2006. Neuroendocrine and neurotransmitter correlates in children with antisocial behaviour. *Horm. Behav.* **50**, 647-654. (doi: [10.1016/j.yhbeh.2006.06.021](https://doi.org/10.1016/j.yhbeh.2006.06.021))
- 20) Bardi M, French JA, Ramirez SM, Brent L. 2004. The role of the endocrine system in baboon maternal behaviour. *Biol. Psych.* **55**, 724-732.
(doi:/doi.org/10.1016/j.biopsycho.2004.01.002)
- 21) Saltzman W, Abbott DH. 2009. Effects of elevated circulating cortisol concentrations on maternal behaviour in common marmoset monkeys (*Callithrix jacchus*). *Psychoneuroendocrinol.* **34**, 1222-1234. (doi: [10.1016/j.psyneuen.2009.03.012](https://doi.org/10.1016/j.psyneuen.2009.03.012))
- 22) DeVries AC, DeVries MB, Taymans SE, Carter CS. 1996. The effects of stress on social

- preferences are sexually dimorphic in prairie voles. *Proc. Natl. Acad. Sci. USA* **93**, 11980-11984. (doi: 10.1073/pnas.93.21.11980)
- 23) Landys MM, Ramenofsky M, Wingfield, JC. (2006). Actions of glucocorticoids at a seasonal baseline as compared to stress-related levels in the regulation of periodic life processes. *Gen. Comp. Endocrinol.* **148**, 132-149. (doi: 10.1016/j.ygcen.2006.02.013)
- 24) Haller J, Mikics E, Makara GB (2008). The effects of non-genomic glucocorticoid mechanisms on bodily functions and the central nervous system. A critical evaluation of findings. *Front Neuroendocrinol.* **29**, 273-291. (doi: 10.1016/j.yfrne.2007.10.004)
- 25) Kumar R, Thompson EB. 2005. Gene regulation by the glucocorticoid receptor: structure: function relationship. *J. of Steroid Biochem. Mol. Biol.* **94**, 383-394. (doi:10.1016/j.jsbmb.2004.12.046)
- 26) DeBattista C, Belanoff J. 2006. The use of mifepristone in the treatment of neuropsychiatric disorders. *Trends Endocrinol. Metab.* **17**, 117-121. (doi:10.1016/j.tem.2006.02.006)
- 27) Raulo A, Dantzer B. Associations between glucocorticoids and sociality across a continuum of vertebrate social behaviour. *In review*
- 28) Schulkin J, Morgan MA, Rosen JB. 2005. A neuroendocrine mechanism for sustaining fear. *Trends Neurosci.*, **28**, 629-635. (doi: 10.1016/j.tins.2005.09.009)
- 29) Curtis JT, Liu Y, Aragona BJ, Wang Z. 2006. Dopamine and monogamy. *Brain Res.* **1126**, 76-90. (doi: 10.1016/j.brainres.2006.07.126)
- 30) Lemos JC, Wanat MJ, Smith JS, Reyes BA, Hollon NG, Van Bockstaele EJ, Chavkin C,

- Phillips PE. 2012. Severe stress switches CRF action in the nucleus accumbens from appetitive to aversive. *Nature* **490**, 402-406. (doi: 10.1038/nature11436)
- 31) Bateman AW, Ozgul A, Nielsen JF, Coulson T, Clutton-Brock TH. 2013. Social structure mediates environmental effects on group size in an obligate cooperative breeder, *Suricata suricatta*. *Ecology* **94**, 587-597. (doi: 10.1890/11-2122.1)
- 32) Nielsen JF, English S, Goodall-Copestake WP, Wang J, Walling CA, Bateman AW, Flower TP, Sutcliffe RL, Samson J, Thavarajah NK, Kruuk LEB, Clutton-Brock TH, Pemberton JM. 2012. Inbreeding and inbreeding depression of early life traits in a cooperative mammal. *Molec. Ecol.* **21**, 2788-2804. (doi:10.1111/j.1365-294X.2012.05565.x)
- 33) Hodge SJ, Manica A, Flower TP, Clutton-Brock TH. 2008. Determinants of reproductive success in dominant female meerkats. *J. Anim. Ecol.* **77**, 92-102. (doi:10.1111/j.1365-2656.2007.01318.x)
- 34) Spong GF, Hodge SJ, Young AJ, Clutton-Brock TH. 2008. Factors affecting the reproductive success of dominant male meerkats. *Molec. Ecol.* **17**, 2287-2299. (doi:10.1111/j.1365-294X.2008.03734.x)
- 35) Clutton-Brock TH, Russell AF, Sharpe LL, Young AJ, Balmforth Z, McIlrath GM. 2002. Evolution and development of sex differences in cooperative behavior in meerkats. *Science* **297**, 253-256. (doi:10.1126/science.1071412)
- 36) Clutton-Brock TH, Brotherton PNM, O'Riain MJ, Griffin AS, Gaynor D, Sharpe L,

- Kansky R, Manser MB, McIlrath GM. 2000. Individual contributions to babysitting in a cooperative mongoose, *Suricata suricatta*. *Proc. R. Soc. Lond. B* **267**, 301-305.
(doi:10.1098/rspb.2000.1000)
- 37) Clutton-Brock TH, Hodge SJ, Spong G, Russell AF, Jordan NR, Bennett NC, Sharpe LL, Manser MB. 2006. Intrasexual competition and sexual selection in cooperative mammals. *Nature* **444**, 1065-1068. (doi:10.1038/nature05386)
- 38) Clutton-Brock TH, Brotherton PNM, Smith R, McIlrath GM, Kansky R, Gaynor D, O'Rian MJ, Skinner JD. 1998. Infanticide and expulsion of females in a cooperative mammal. *Proc. R. Soc. Lond. B* **265**, 2291-2295. (doi:10.1098/rspb.1998.0573)
- 39) Young AJ, Clutton-Brock T. 2006. Infanticide by subordinates influences reproductive sharing in cooperatively breeding meerkats. *Biol. Lett.* **22**, 385-387.
(doi:10.1098/rsbl.2006.0463)
- 40) Kutsukake N, Clutton-Brock TH. 2006. Aggression and submission reflect reproductive conflict between females in cooperatively breeding meerkats *Suricata suricatta*. *Behav. Ecol. Sociobiol.* **59**, 541-548. (doi: 10.1007/s00265-005-0079-7)
- 41) English S, Nakagawa S, Clutton-Brock TH. 2010. Consistent individual differences in cooperative behaviour in meerkats (*Suricata suricatta*). *J. Evol. Biol.* **23**, 1597-1604.
(doi:10.1111/j.1420-9101.2010.02025.x)
- 42) Brotherton PNM, Clutton-Brock TH, O'Riain MJ, Gaynor D, Sharpe L, Kansky R, McIlrath GM. 2001. Offspring food allocation by parents and helpers in a cooperative mammal. *Behav. Ecol.* **12**, 590-599. (doi:10.1093/beheco/12.5.590)

- 43) Kunc HP, Madden JR, Manser MB. 2007. Begging signals in a mobile feeding system: the evolution of different call types. *Am. Nat.* **170**, 617-624. (doi: 10.1086/521233)
- 44) Clutton-Brock TH, Gaynor D, Kansky R, MacColl ADC, McIlrath G, Chadwick P, Brotherton PNM, O’Riain JM, Manser M, Skinner JD. 1998. Costs of cooperative behaviour in suricates (*Suricata suricatta*). *Proc. R. Soc. Lond. B* **265**, 185-190. (doi: 10.1098/rspb.1998.0281)
- 45) English S, Bateman AW, Clutton-Brock TH. 2011. Lifetime growth in wild meerkats: incorporating life history and environmental factors into a standard growth model. *Oecologia* **169**, 143-153. (doi: 10.1007/s00442-011-2192-9)
- 46) English S, Bateman AW, Mares R, Ozgul A, Clutton-Brock TH. 2014. Maternal, social and abiotic environment effects on growth vary across life stages in a cooperative mammal. *J. Anim. Ecol.* **83**, 332-342. (doi:10.1111/1365-2656.12149)
- 47) Altmann J. 1974. Observational study of behaviour: sampling methods. *Behaviour* **49**, 227-265. (doi: [10.1163/156853974X00534](https://doi.org/10.1163/156853974X00534))
- 48) Thavarajah NK, Fenkes M, Clutton-Brock TH. 2014. The determinants of dominance relationships among subordinate females in the cooperatively breeding meerkat. *Behaviour* **151**, 89-102. (doi:[10.1163/1568539X-00003124](https://doi.org/10.1163/1568539X-00003124))
- 49) Russell AF, Sharpe LL, Brotherton PNM, Clutton-Brock TH. 2003. Cost minimization by helpers in cooperative vertebrates. *Proc. Natl. Acad. Sci. USA* **100**, 3333-3338. (doi: 10.1073/pnas.0636503100)
- 50) Thornton A. 2008. Early body condition, time budgets and the acquisition of foraging skills in meerkats. *Anim. Behav.* **75**, 951-962. (doi:10.1016/j.anbehav.2007.08.004)

- 51) Burnham KP, Anderson DR. 2002. *Model Selection and Multimodel Inference*. (Springer-Verlag, New York, NY).
- 52) R Development Core Team. 2013. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria; available at www.r-project.org/.
- 53) Barton K. 2015. MuMIn: Multi-model inference. R package version 1.15.1; available at <http://CRAN.R-project.org/package=MuMIn>.
- 54) Gelman A, Hill J. 2007. *Data Analysis using Regression and Multilevel/Hierarchical Models* (Cambridge University Press, New York, NY).
- 55) Bates DM, Maechler M, Bolker B, Walker S. 2015. Fitting linear mixed-effects models using lme4. *J. Stat. Soft.* **67**, 1-48. (doi:10.18637/jss.v067.i01)
- 56) Kuznetsova A, Brockhoff PB, Christensen RHB. 2016. lmerTest: Tests in Linear Mixed Effects Models. R package version 2.0-30; available at <http://CRAN.R-project.org/package=lmerTest>
- 57) Zuur AF, Ieno EN, Elphick CS. 2010. A protocol for data exploration to avoid common statistical problems. *Meth. Ecol. Evol.* **1**, 3-14. (doi: 10.1111/j.2041-210X.2009.00001.x)
- 58) Clutton-Brock TH, O’Riain MJ, Brotherton PNM, Gaynor D, Kansky R, Griffin AS, Manser M. 1999. Selfish sentinels in cooperative mammals. *Science* **284**, 1640-1644. (doi:10.1126/science.284.5420.1640)
- 59) Nichols HJ, Amos W, Bell MBV, Mwanguhya F, Kyabulima, Cant MA. 2012. Food availability shapes patterns of helping effort in a cooperative mongoose. *Anim. Behav.* **83**, 1377-1385 (doi:10.1016/j.anbehav.2012.03.005).

- 60) Russell AF, Hatchwell BJ. 2001. Experimental evidence for kin-based helping in a cooperatively breeding vertebrate. *Proc. R. Soc. Lond. B* **268**, 2169-2174 (doi:10.1098/rspb.2001.1790).
- 61) Griffin AS, West SA. 2003. Kin discrimination and the benefit of helping in cooperatively breeding vertebrates. *Science* **302**, 634-636 (doi:10.1126/science.1089402).
- 62) Clutton-Brock TH, Russell AF, Sharpe LL. 2003. Meerkat helpers do not specialize in particular activities. *Animal Behaviour* **66**, 531-540. (doi: 10.1006/anbe.2003.2209)
- 63) Santema P, Clutton-Brock T. 2012. Dominant female meerkats do not use aggression to elevate work rates of helpers in response to increased brood demand. *Anim. Behav.* **83**, 827-832. (doi:10.1016/j.anbehav.2011.12.032)
- 64) Santema P, Teitel Z, Manser M, Bennett N, Clutton-Brock T. 2013. Effects of cortisol administration on cooperative behavior in meerkat helpers. *Behav. Ecol.* **24**, 1122-1127. (doi:10.1093/beheco/art039)
- 65) Doolan SP, Macdonald DW. 1996. Dispersal and extra-territorial prospecting by slender-tailed meerkats (*Suricata suricatta*) in the south-western Kalahari. *J. Zool.* **240**, 59-73. (doi: 10.1111/j.1469-7998.1996.tb05486.x)
- 66) Griffin AS, Pemberton JM, Brotherton PNM, McIlrath G, Gaynor D, Kansky R, O'Riain J, Clutton-Brock TH. 2003. A genetic analysis of breeding success in the cooperative meerkat (*Suricata suricatta*). *Behav. Ecol.* **14**, 472-480. (doi: [10.1093/beheco/arg040](https://doi.org/10.1093/beheco/arg040))
- 67) Handa RJ, Burgess LH, Kerr JE, O'Keefe JA. 1994. Gonadal steroid hormone receptors and sex differences in the hypothalmo-pituitary-adrenal axis. *Horm. Behav.* **28**, 464-476. (doi: [10.1006/hbeh.1994.1044](https://doi.org/10.1006/hbeh.1994.1044))

- 68) Bourke CH, Harrell CS, Neigh GN. 2012. Stress-induced sex differences: Adaptations mediated by the glucocorticoid receptor. *Horm. Behav.* **62**, 210-218. (doi: [10.1016/j.yhbeh.2012.02.024](https://doi.org/10.1016/j.yhbeh.2012.02.024))
- 69) Owen D, Matthews SG. 2003. Glucocorticoids and sex-dependent development of brain glucocorticoid and mineralocorticoid receptors. *Endocrinol.* **144**, 2775-2784. (doi: [10.1210/en.2002-0145](https://doi.org/10.1210/en.2002-0145))

Fig. 1. (A) Subordinate female and male meerkats fed mifepristone (n=9) exhibited significantly more babysitting (proportion of total babysitting provided by subordinate) than did those fed cortisol (n=9) or the controls (n=8). (B and C) Unlike babysitting, the effects of the treatments on pupfeeding were sex-specific (Tables S2-S3). Subordinate females treated with mifepristone (n=8) were significantly more likely to feed pups compared to those treated with cortisol (n=10) or the controls (n=7) but the amount of prey biomass found by subordinate females and fed to pups did not differ among the treatment groups. The probability of pupfeeding by subordinate males was not affected by the treatments but males treated with cortisol (n=4) fed significantly more of the prey biomass they found to pups compared to those fed mifepristone (n=7) but not controls (n=7).

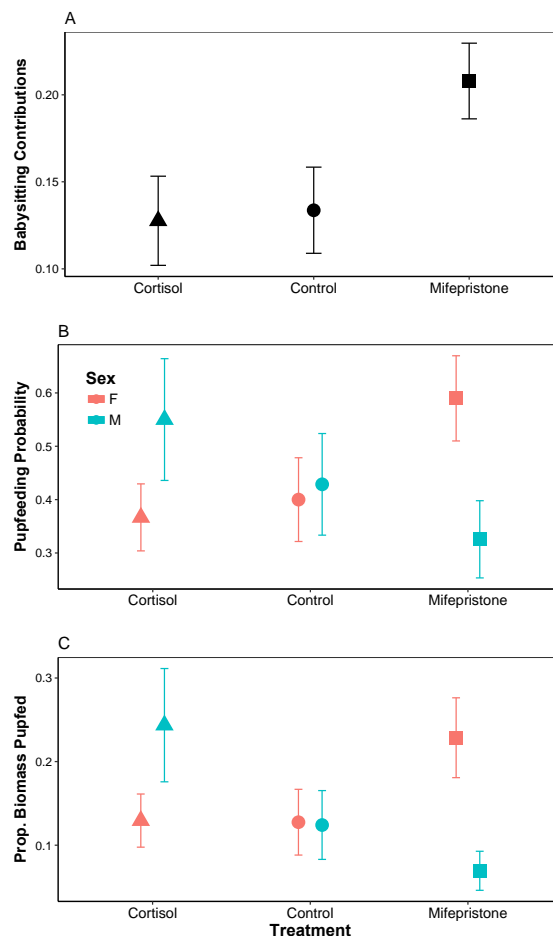


Fig. 2. Behavioral data collected over 19 years (>110,483 hours) indicated that the amount of aggression subordinate female meerkats (n=713) received from the dominant female (n=98 females from 40 groups) varied according to whether the dominant female was pregnant and whether there were offspring in the group that subordinates were taking care of (babysitting and pupfeeding: Table S7). Subordinate females received the highest levels of aggression from the dominant female when she was pregnant and there were no pups in the group. The amount of aggression directed at subordinate males (n=807) was not significantly influenced by the pregnancy status of the dominant female or whether there were offspring in the group being babysat or fed.

