Adult meerkats modify close call rate in the presence of pups

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Abstract

In animals, signaling behavior is often context-dependent, with variation in the probability of emitting certain signals dependent on fitness advantages. Senders may adjust signaling rate depending on receiver identity, presence of audiences, or noise masking the signal, all of which can affect the benefits and costs of signal production. In the cooperative breeding meerkat Suricata suricatta, group members emit soft contact calls, termed as “close calls”, while foraging in order to maintain group cohesion. Here, we investigated how the close calling rate during foraging was affected by the presence of pups, that produce continuous, noisy begging calls as they follow older group members. Adults decreased their overall close call rate substantially when pups were foraging with the group in comparison to periods when no pups were present. We suggest this decrease was likely due to a masking effect of the loud begging calls, which makes the close call function of maintaining group cohesion partly redundant as the centrally located begging calls can be used instead to maintain cohesion. There was some support that adults use close calls strategically to attract specific pups based on fitness advantages, that is, as the philopatric sex, females should call more than males and more to female pups than male pups. Dominant females called more than dominant males when a pup was in close proximity, while subordinates showed no sex-based differences. The sex of the nearest pup did not affect the calling rate of adults. The study shows that meerkats modify their close call production depending on benefits gained from calling and provides an example of the flexible use of one calling system in the presence of another, here contact calls versus begging calls, within the same species.

Key words: call rate, communication, contact calls, cooperation, flexibility, group cohesion.
depending on the function of the calls, and therefore differ between contexts.

One such context is seen in group foraging, where group members are spaced out but stay in contact with each other with vocalizations (Koda et al. 2008; Kondo and Watanabe 2009; Fichtel and Manser 2010). However, by emitting contact calls, a sender reveals its position not only to other group members, but also to predators (Deecke et al. 2002). Furthermore, in groups, where group members compete for resources rather than sharing them, they may try to avoid physical contact with others, or some specific individuals, and therefore not emit calls (Gros-Louis 2004). For example, in groups where dependent young follow adults around to be fed, such as in fledgling birds (Thompson et al. 2013) or pups in mongoose species (Kunc et al. 2007; Bell 2008), only highly successful foragers may be able to feed, while the others may try to avoid being followed by hungry offspring, and keep silent. Moreover, in species where relationships differ among group members, signals may be used strategically and senders will selectively emit calls to those individuals from which they will receive the most benefits.

Noise in the environment can mask vocal signals and reduce the efficacy of contact calls in maintaining spatial coordination between foraging group members. Most studies on the masking effects on vocal signals have dealt with the influence of anthropogenic or biotic noise in the environment (Derryberry et al. 2016). However, vocalizations produced by other members of the same group using a different call type may also have a disruptive influence or masking effect, if they are emitted at the same time. In the context of group coordination vocalizations, the loud begging calls from fledglings in birds (Thompson et al. 2013) or pups in mammals (Manser and Avey 2000) are likely to mask soft contact calls. The question arises: how do these 2 communication systems affect each other, whether the more obvious and louder begging calls make the low amplitude how do these 2 communication systems affect each other, whether the different call types are used for coordinating instead of the more quiet close calls. However, if the relative change in close calling rates differed between individuals, this could reflect differences in the direct and indirect benefits associated with cooperative care of pups. More successful foragers provide more of their food (Clutton-Brock et al. 2002), and potentially use close calls to attract pups to follow them. As the philopatric sex, females may benefit more from attracting and provisioning female pups since larger group sizes are associated with enhanced fitness (Clutton-Brock et al. 2002) because pup growth rate is positively related to the number of carers per pup, and pup survival between emergence from the burrow and foraging independence is higher in larger groups (Russell et al. 2002). Moreover, females feed pups at a higher rate than males, feed a higher proportion of found food than males (Brotherton et al. 2001; English et al. 2008), and feed female pups more than male pups (Brotherton et al. 2001). Female pups stay closer to adults than male pups (Hollein and Manser 2006). Therefore, females may emit close calls to attract or coordinate movements during foraging with female pups as this type of cooperative behavior could provide increased long-term benefits. Such, and if close calls are used strategically to make pups follow, we predict that females will produce more close calls in the presence of pups than males and they will produce more close calls when female pups are nearby compared with when male pups are nearby.

The close calls, which meerkats only produce during foraging activities, are individually distinct and allow individuals to monitor group members spatially (Townsend et al. 2011; Reber et al. 2013). The rate of producing close calls is highly variable and is influenced by weather conditions, reproductive season, the social environment, and socio-spatial contexts (Mausbach et al. 2017). Meerkats can adjust their calling rate depending on the identity of the closest neighbor (Wolf 2014). Individuals at the leading edge of a group give more close calls than individuals at the back edge of the group, and the rate of close calling increases with decreasing proximity between group members (Engesser 2011). Preliminary observations further showed that meerkats decrease close calling rates when pups are foraging with the group (Manser 1998). Experimental studies have also demonstrated that close calls influence movement and cohesion during foraging by attracting receivers toward the caller (Gall and Manser 2017).

Here, we investigated different factors at the individual and group level to identify what explained the decrease in close calling rate of adult meerkats when pups were present. If all of the individuals decreased their calling rate, this may be related to the function of the close call to maintain group cohesion. When pups are foraging with the group, their loud and pervasive begging calls may be used by group members to guide coordinated movements and spatial positioning instead of the more quiet close calls. However, if the relative change in close calling rates differed between individuals, this could reflect differences in the direct and indirect benefits associated with cooperative care of pups. More successful foragers provide more of their food (Clutton-Brock et al. 2002), and potentially use close calls to attract pups to follow them. As the philopatric sex, females may benefit more from attracting and provisioning female pups since larger group sizes are associated with enhanced fitness (Clutton-Brock et al. 2002) because pup growth rate is positively related to the number of carers per pup, and pup survival between emergence from the burrow and foraging independence is higher in larger groups (Russell et al. 2002). Moreover, females feed pups at a higher rate than males, feed a higher proportion of found food than males (Brotherton et al. 2001; English et al. 2008), and feed female pups more than male pups (Brotherton et al. 2001). Female pups stay closer to adults than male pups (Hollen and Manser 2006). Therefore, females may emit close calls to attract or coordinate movements during foraging with female pups as this type of cooperative behavior could provide increased long-term benefits. Such, and if close calls are used strategically to make pups follow, we predict that females will produce more close calls in the presence of pups than males and they will produce more close calls when female pups are nearby compared with when male pups are nearby.

We used paired audio recordings of individuals taken before pups started foraging with the group and when pups were foraging with the group (1) to assess the influence the presence of pups has on close call rates and (2) to determine what group- or individual-level factors may predict individual differences in the relative change in call rate in these different contexts. The group-level traits we measured include group size, litter size, the ratio of pups to carers, and the perceived predation risk within the group. At the individual level, we investigated whether the relative change in overall calling rate was affected by the caller’s sex, social rank, and relative contribution to pup provisioning and vigilance behaviors within the group. Furthermore, we examined (3) flexibility of call rates within short time windows directly associated with the close presence of a pup to determine whether these rates were influenced by the sex or rank of the caller or the sex of the nearest pup.
Materials and Methods

Study area

Research was conducted on habituated wild meerkats at the Kalahari Meerkat Project, Kuruman River Reserve (26°59' S, 21°50' E) near Van Zylsrus in South Africa (Clutton-Brock et al. 1998). The arid landscape at the study site consists of wind-blown dunes, dry riverbeds, and flats containing a mixture of sparsely distributed and patchy vegetation including multiple grass, shrub, and tree species. The meerkats were habituated to observers walking with the foraging groups in order to collect behavioral data and sound recordings within 0.5–1 m (Manser and Avey 2000). Data were collected from April to June 2016 and January to March 2017 on 60 adult individuals in 8 different groups. This represented 9 different breeding events (i.e., unique litters). Group size when the pups began foraging with the group ranged from 9 to 20 individuals (mean ± SD = 12.44 ± 3.64) and litter size ranged from 1 to 5 pups (mean ± SD = 3.22 ± 1.30).

Data collection

To calculate close calling rate, we conducted focal animal samples on adult meerkats while they foraged. Two focal periods of 5–10 min (average of 8 min) were conducted on adults in each of 2 time periods: (1) from 10 days after litter birth until the pups left the burrow and began foraging with the group (no pups period) and (2) after the pups had been foraging with the group for at least a week but were not yet independent (with pups period). Each target period for recordings was approximately 2 weeks long. Recordings were made with a directional microphone (Sennheiser ME66 with K6 powering module) with windshield (Reinhartd, Germany) fixed to a tripod leg connected to a solid state recorder (Marantz PMD661, sampling frequency 48 kHz, 24 bit). A second microphone (Joseph E-280 Dynamic Microphone) was connected to the second channel on the recorder through which the observer annotated the focal individual’s behavior and the context in which each call was produced as well as the identity and distance of nearest neighboring adult and pup. Sound recordings were only conducted in a foraging context and the recording was paused when the focal individual was engaged in non-foraging behavior for more than 30 s.

All close calls produced by the focal individual and all nearest neighbor annotations in the .wav files were manually labeled in Cool Edit Pro v2.0 (Syntrillium) and Audacity v2.1.2 (http://audacity.sourceforge.net/). The overall rate of close calls was calculated for each individual in each period by first dividing the number of close calls by the duration of the recording period, and then averaging the call rates of the 2 sound focal topics together for each period. Forty-eight out of 60 individuals were recorded in both time periods and subsequent analysis of overall call rate was restricted to these individuals with paired recordings. Short-term calling rates occurring during the close presence of a single pup of known sex were calculated within 10 s windows surrounding the field annotation of pup presence between 1 and 5 m from the focal individual. To be selected for analysis, no other pups or adults could be present within this time window at a distance closer than 5 m. Instances of a pup present at less than 1 m distance were not included to avoid disruptions in calling rate due to aggressive interactions or pup feeds. Since meerkats occasionally move at a quick pace during foraging, the short window duration of 10 s was selected to increase the likelihood that calls were delivered while the only nearest neighbor was the specific pup described in the annotation.

We also attempted to record all pup provisioning events, bouts of anti-predator vigilance, and alarm events during foraging sessions for an average of 10 h per group per week. These observations were made on days adjacent to the day on which we recorded vocalizations. Thus, these data permitted us to make comparisons between periods of having the groups foraging without the pups and the period with pups, and calculate relative contributions by different individuals to these cooperative behaviors. We were careful to avoid any systematic bias in the data by constantly moving around among all different group members and covering the spatial range of the group equally. During these observation periods, observers noted the initiator of feeding events to pups as well as the recipient, the occurrence and duration of vigilance behaviors displayed by each individual, and the number of predator alarm events produced by the group. Each individual's relative contribution to pup provisioning within their group was calculated by dividing the number of times an individual provisioned a pup with food by the total number of pup provisioning events recorded for the group in the first 20 days after the pups started foraging with the group. Similarly, each individual's relative contribution to group vigilance behavior was calculated by dividing the total number of vigilance events per individual by the total number of vigilance events observed by all group members within their group during a 40-day period surrounding the date that the pups started foraging with the group (i.e., 20 days before and after this date). Group-wide alarm event rates for this same time period were calculated by dividing the total number of alarm events observed at each group by the total number of hours each group was observed during this 40 day period.

Statistical analysis

A Wilcoxon signed ranks test was used to compare the overall close call rate of each individual during sound focal samples when there were no pups to their overall close call rate when there were pups foraging with the group. Wilcoxon rank sum tests were used to test for significant differences in the overall call rate between the sexes and between dominants and subordinates within the 2 separate time periods. Non-parametric tests were used because the data were not normally distributed, as determined by both visual inspection of QQ plots and Shapiro–Wilk tests of normality. The relative change in call rate between the No Pup and With Pup periods was calculated as a percentage change for each individual as follows:

\[
\frac{\text{Call Rate}_{\text{With Pups}} - \text{Call Rate}_{\text{No Pups}}}{\text{Call Rate}_{\text{No Pups}}} \times 100.
\]

A linear mixed-effects model fit by maximum likelihood (with Satterthwaite approximations of degrees of freedom and P values) was used to determine what factors may partially explain the lower close call rate when pups were foraging with the group. Percentage change in close call rate between the 2 periods was the response variable and group nested within year was included as a random intercept term to account for repeated measurements within the same group and year. Focal ID was not included as a random effect because there was only one data point per individual. Predictor variables added to the model as main effects included sex, social rank (dominant or subordinate), relative contribution to pup feeding, relative contribution to vigilance behavior, group alarm rate, group size, litter size, and pup to carer ratio were included. The interaction term between sex and rank was also included. Age was not included in the model as a fixed effect because the age distribution was strongly associated with social rank (e.g., younger subordinates and older dominants). Prior to analysis of overall call rates or percentage
change in call rate, one extreme outlying data point was removed which was in the 99.9% percentile of the data distribution. Model assumptions were examined graphically with QQ plots and plots of fitted values versus standardized residuals.

A mixed model regression with Poisson distribution was used to assess what factors influence short-term call rates within narrow time windows directly associated with the close presence of a pup during the period when pups were foraging with the group. Here, we tested if the number of close calls produced within the 10 s windows of time surrounding each nearest neighbor annotation was influenced by the sex and rank of the caller and the sex of the nearest pup. Fixed effects within the full model included the sex and rank of the caller, the sex of the nearest pup present at a distance of 1–5 m, and the interactions between these factors. Focal ID nested within group was included as a random effect (intercept). Year was initially included as a random effect as well, with group and focal ID nested within it, but was subsequently removed as it displayed very little variation and likelihood ratio tests confirmed its inclusion did not improve model fit. To examine the influence of focal sex more closely, the dataset was subsequently split into dominant and subordi- nate subsets for further analysis. Model outcomes were examined for over- and under-dispersion by comparing the sum of squared Pearson residuals to the residual degrees of freedom. Zero-inflation was assessed by comparing the proportion of zero counts observed in the dataset to the proportion of zero counts predicted by the Poisson distribution. As there were no issues with dispersion or zero-inflation detected, the final models were run with the generalized log linear mixed model regression with Poisson distribution (fit by maximum likelihood, Laplace approximation).

Stepwise backward model selection procedures were used to determine the best fit models for both the linear and Poisson mixed effects models. Best fit models were compared with null models using likelihood ratio tests (within anova functions), with secondary comparisons conducted using AICc scores. Boxplots were used to visualize the data: the lower and upper hinges correspond to the first and third quartiles, the middle line represents the median, and the whiskers extend to the smallest and largest values within 1.5 times the interquartile range. All tests were run in R Studio v1.0.136 with R version 3.3.1 (R Core Team 2016), using the lme4 and glmer functions in the lme4 package (Bates et al. 2015).

**Results**

**Close calling rate of adults depending on pups’ presence**

There was a significant decrease in close call rate between the period before pups foraged with the group and the period when pups were foraging with the group (MedianNo Pups = 5.80 calls/min, MedianWith Pups = 2.04 calls/min; Wilcoxon signed rank test, \( V_{48} = 1,129, P < 0.0001 \), Figure 1).

**What factors predict change in overall call rate?**

Sex of the caller significantly influenced the percentage change of call rate between the 2 conditions (\( t_{19,74} = 2.05, P = 0.047 \), Table 1). While females exhibited a 65.6% reduction in call rate after pups started foraging with the group, males only reduced their call rate by 45.8%. This reduced model with focal sex as the only remaining fixed effect was a significantly better fit than the null model (\( \chi^2 = 4.032, P = 0.045 \)) and was less complex but similar in fit to the next best fit reduced model with focal sex, group alarm rate, and group count as fixed effects (\( \chi^2 = 4.066, P = 0.131 \), see Table 1 for full, reduced, and null models). Although females had a higher median call rate than males in the period before pups were foraging with the group (Medianfemales = 7.38 calls/min, Medianmales = 5.12 calls/min) and a slightly lower median call rate than males after pups started foraging with the group (Medianfemales = 1.38 calls/min, Mediansubordinates = 2.11 calls/min), there was no significant difference between the sexes in overall call rate within either period (no pups: \( W_{48} = 332, P = 0.177 \); with pups: \( W_{48} = 257, P = 0.712 \), Figure 2). Thus, although females showed a greater relative change in their call rate between the 2 conditions, there was no difference between the sexes in overall call rate within each condition. Similarly, there was no significant difference in overall call rate between dominant and subordinate individuals within each condition (no pups: \( W_{48} = 198, P = 0.163 \); with pups: \( W_{48} = 280, P = 0.733 \)).

**Factors influencing short-term call rate when pups are nearest neighbors**

The short-term call rate of adults when pups were in close proximity was not significantly affected by focal sex, focal rank, pup sex, or their interactions within the full model. Additionally, there was no significant difference between the null model and the full model (\( \chi^2 = 5.707, P = 0.574 \)) or between the null model and any reduced model. However, when examining call rates of only dominant individuals, the sex of the caller significantly influenced the call rate during periods when a pup was the nearest neighbor at a distance of 1–5 m. Dominant females called at a significantly higher rate near close pups than dominant males (2.205 calls/min vs. 0.727 calls/min; \( Z_{diff} = 2.327, P = 0.020 \), Figure 3). The sex of the pup did not affect call rate and was removed from the final model. The best fit reduced model, with focal sex as the main effect, tended toward being significantly different from the null model (\( \chi^2 = 3.325, P = 0.068 \)) and possessed the lowest AICc score.

Neither focal sex nor sex of the pup predicted call rate by subordi- nate individuals, although there was a non-significant trend for subordinates to call more in the presence of male pups than female pups (\( Z_{diff} = 1.715, P = 0.086 \), Figure 3). The best fit reduced model for subordinates included focal sex and pup sex as the main
Table 1. Results of linear mixed effects model selection for factors predicting percentage change of close call rate between the no pups and with pups conditions

| Fixed effects | Value  | Standard error | DF  | t value | P value | AICc   | logLik  |
|---------------|--------|----------------|-----|---------|---------|--------|---------|
| Null model   | -55.043| 7.527          | -   | -7.313  | <0.0001 | 479.673| -235.360|
| Best fit model | -65.532| 8.813          | 15.66 | -7.438  | <0.0001 | 478.152| -233.344|
| Sex M         | 19.706 | 9.604          | 39.74 | 2.052   | 0.0468  |        |         |
| Next best fit model | -96.678| 32.194         | 6.49  | -3.003  | 0.022   | 479.494| -231.311|
| Sex M         | 19.748 | 9.628          | 41.08 | 2.051   | 0.047   |        |         |
| Group alarm rate | 32.872| 15.545         | 7.53  | 2.115   | 0.070   |        |         |
| Group count   | -4.264 | 1.960          | 7.30  | -2.176  | 0.065   |        |         |
| Full model    | -114.981| 41.606        | 47.00 | -2.764  | 0.008   | 495.577| -229.273|
| Sex M         | 23.571 | 17.521         | 47.00 | 1.345   | 0.185   |        |         |
| Rank SUB      | -4.390 | 14.605         | 47.00 | -0.301  | 0.765   |        |         |
| Pup carer ratio | 110.154| 78.285         | 47.00 | 1.407   | 0.166   |        |         |
| Pup feed ratio | -56.515| 57.204         | 47.00 | -0.988  | 0.328   |        |         |
| Vigilance ratio | -21.577| 50.699         | 47.00 | -0.426  | 0.672   |        |         |
| Group alarm rate | 30.652| 17.837         | 47.00 | 1.718   | 0.092   |        |         |
| Group count   | -1.244 | 3.867          | 47.00 | -0.322  | 0.749   |        |         |
| Pup count     | -14.415| 13.519         | 47.00 | -1.066  | 0.292   |        |         |
| SexM:RankSUB  | 0.884  | 20.812         | 47.00 | 0.042   | 0.966   |        |         |

Discussion

In meerkats, group members produced fewer close calls when they had pups foraging with them in comparison to periods when no pups were foraging with the group. There was a significant difference in the relative change in close call rate between the 2 periods as a function of the caller’s sex: females reduced calling rate more than males. This may have been because females appeared to call more than males when pups were not foraging with the group, while when pups were foraging with the group, both females and males called at about the same rate (Figure 2). Within each period, however, there was no significant difference in overall call rate between the sexes. The relative change in calling rate between these 2 periods was not significantly affected by other individual-level factors such as social rank, relative pup provisioning, or relative vigilance behavior, or other group-level factors such as group size, litter size, pup to carer ratio, or group alarm rates. However, within the period when pups were foraging in close proximity, dominant females had higher short-term close calling rates than dominant males.

There are several reasons that may explain the reduction in the rate of emitting close calls when pups are foraging with the group. Pups are typically located in the center of the group and are constantly emitting loud begging calls. Thus, the soft close calls are likely to be masked by these begging calls. The function of the close calls to maintain group cohesion becomes at least partly redundant, because group members can direct themselves toward the center of the group using the loud begging calls rather than the highest rate of close calls (Gall and Manser 2017). There was also no effect of group size, litter size, or pup–carer ratio, suggesting, as soon as meerkats had a begging pup with them they reduced the rate of emitting close calls. These results support the hypothesis that the decreased calling rate was because of the redundant function of close calls to maintain group cohesion when begging pups emitting the loud calls were present.

The other explanation, that older group members may selectively use close calls to attract the attention of specific pups where they gain the largest fitness benefit, was partially supported. We predicted that as the philopatric sex, females would produce more close calls in the presence of pups than males and would produce more close calls when female pups were close compared with when male pups were close. The sex of the caller significantly predicted call rate when a close pup was present: dominant females called more in the presence of a close pup than dominant males. These results suggest that dominant females call more than dominant males when pups are near because they gain a greater fitness benefit by doing so, potentially from attracting pups for provisioning. This is supported by the observation that female parents provision pups more than male parents (Brotherton et al. 2001). However, subordinate females and males showed no difference in their calling behaviors in the presence of a close pup, despite the fact that female helpers provision pups more than male helpers (Clutton-Brock et al. 2001) and female...
helpers provision a similar proportion of food to pups as female parents (Brotherton et al. 2001). Although females provide more food to female pups than male pups (Brotherton et al. 2001), the sex of the nearest close pup did not affect the calling rate of adult group members. Furthermore, when foragers were foraging with the group, the overall call rate was similarly low for all different individuals, independent of sex or rank. Since older group members not only have to feed pups, but also forage for themselves, they may try to avoid having pups following them as long as they are not satiated to a certain level themselves. This is supported by the fact that the least successful foragers provided the least amount of prey items to the pups (Clutton-Brock et al. 2001) but not supported by the lack of a correlation between the relative change in close call rate and generosity of carers. Experiments are needed to identify whether close calls attract pups to follow the vocalizing carer.

This adjustment of close call rate in meerkats to the pups’ presence indicates the flexibility and control of senders on the production of this signal type. Furthermore, it strongly suggests that the reduction in close call rate is due to a masking effect of the obvious loud begging calls, which make the close call function of maintaining group cohesion partly redundant. The few close calls emitted when pups were present may still have the function to attract pups, with dominant females potentially attracting more pups than dominant males, although with the current sample size we have no clear evidence that these calls are used to attract specific pups by the different social categories of adult group members. This study suggests that meerkats adjust their close call production to the masking effect due to pup begging calls, and therefore according to the benefit of emitting these calls in the specific situation. It provides an example of the flexible use of one calling system in the presence of another, here contact calls versus begging calls, within the same species.

Ethical Note

All data collection adhered to ASAB guidelines. All research was conducted under the permission of the ethical committee of Pretoria University ( Permit number: EC031-13) and the Northern Cape Conservation Service (FAUNA 192/2014), South Africa.

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Figure 3. The number of close calls produced by dominant individuals (top) and subordinate individuals (bottom) within the 10 s time window when a pup was the only neighbor between 1 and 5 m. Boxplot color represents the sex of the nearest neighbor (gray = female pup; white = male pup). The numbers under each boxplot represent the number of time windows included in the analysis for each combination (e.g., there were 14 time windows where a female pup was the nearest neighbor to a subordinate female).
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