Recruitment calling to direct predator cues and its effect on

predator responses in meerkats

Markus Zöttl^{1,2,6}, Raphaela Lienert^{2,3}, Tim Clutton-Brock^{2,4,5}, Eva Millesi¹ & Marta B. Manser^{2,3,5}

¹Department of Behavioural Biology, University of Vienna, Althanstr. 14, 1090 Vienna, Austria

²Kalahari Meerkat Project, Kuruman River Reserve, Northern Cape, South Africa

³Institute of Evolutionary Biology and Environmental Studies, University of Zurich, Winterthurerstrasse 190 CH-8057, Zurich, Switzerland

⁴LARG, Department of Zoology, University of Cambridge, U.K.

⁵ Mammal Research Institute, University of Pretoria, Pretoria, South Africa

⁶Department of Behavioural Ecology, Institute of Ecology and Evolution, University of Bern, Switzerland

Corresponding Author: markus.zoettl@iee.unibe.ch

Summary

Behavioral responses of animals to direct predator cues (DPC; e. g. urine) are common and may improve their survival. We investigated wild meerkat (*Suricata suricatta*) responses to DPCs by taking an experimental approach. When meerkats encounter a DPC they often recruit group members by emitting a call type, which causes the group members to interrupt foraging and approach the caller. The aim of this study was to identify the qualities of olfactory predator cues, which affect the strength of response by meerkats, and determine the benefits of responses to such cues. Experimental exposure to dog (*Canis lupus*) urine as a DPC revealed that the recruited individuals increased vigilance to fresh urine in comparison to older urine, whereas a higher quantity of urine did not induce such an effect. Both freshness and higher quantities increased the proportion of group members recruited. These results indicate that recruitment might play a crucial role in correctly assessing the current level of danger and that recruiting might facilitate group decision-making. To test the prediction that the reaction to a DPC enhances early predator response, we presented a DPC of a predator and a control cue of a herbivore, and each time simultaneously moved a full-mounted caracal (*Caracal caracal*) in the vicinity of the group. Meerkats responded earlier to the caracal when the DPC was presented, indicating that the response to a DPC facilitates predator response and that they use information from the cue that reliably reflects the risk in the current moment.

Key words: direct predator cues, recruitment, olfactory cues, vigilance, meerkats, predator odor, predator detection

Introduction

Many animals face a trade-off between foraging and predator avoidance (Lima and Dill, 1990; Verdolin, 2006; Morrison, 2011). It is therefore highly beneficial for individuals to assess the actual predation risk and adjust their anti-predator investment according to the perceived danger. Evidence for this adjustment has been demonstrated in a number of species (Hilton *et al.*, 1999; Barta *et al.*, 2004; Devereux *et al.*, 2006). Theoretical models support the assumption that changes in foraging behavior help to minimize predator exposure and encounter rate (Lima and Dill, 1990; Lima, 1998) and experimental evidence suggests that vigilant individuals spot approaching predators at further distances than foraging ones (Lima and Bednekoff, 1999) and are probably less vulnerable to predation (Fitzgibbon, 1989; Krause and Godin, 1996; Hilton *et al.*, 1999).

Direct predator cues (DPC) are inadvertently left calling cards (e.g. urine, feces, hair heterospecific alarm calls) that can be used as indicators of nearby predators and magnified predation risk. Consequently, DPC recognition likely allows an individual to assess the current level of danger. Mammals respond to predator odors with changes in spatial activity, decreased feeding rate and increased vigilance (Berger *et al.*, 2001; Apfelbach *et al.*, 2005; Blumstein *et al.*, 2008) and similar responses to DPC are widespread in different taxonomic groups (birds: Roth *et al.*, 2008; Ridley *et al.*, 2010, fish: Wisenden, 2000; Brown, 2003; Ward and Mehner, 2010, reptiles: Ito and Mori, 2010, invertebrates: Foust *et al.*, 2001; Gherardi *et al.*, 2011, for a review see: Kats and Dill, 1998). Although behavioral changes are well documented and are frequently assumed to be adaptive, there currently exists little experimental evidence demonstrating that reactions to DPCs actually improve an animal's ability to avoid predator encounters – a response which would have direct fitness benefits to the prey.

Moreover, it is of advantage for prey species to be able to assess the reliability of DPCs as old cues are probably not associated with high risk situations. Such adjustment has been shown for the wolf spider (*Paradosa milvina*) and the brushtail possum (*Trichosurus vulpecula*), which react more strongly to a fresh than an old cue of their predator (Barnes *et al.*, 2002; Kirmani *et al.*, 2010). Other qualities of the cue, which are also highly variable, may be less reliable for assessing the current predation risk, such as the amount of predator urine deposited. Variation in the amount of urine deposited from carnivores can be caused by different ways of urinating that may serve different functions (e.g. fox; Jorgenson *et al.*, 1978, feral cat: Natoli, 1985, domestic dog: Hart, 1974). Carnivores mark their territory by spraying, a behavior where often small amounts of urine are excreted, or excrete larger amounts without dispersing. As expected, studies also found strong sex dependent variation in the use of these techniques in carnivores (Hart, 1974; Liberg, 1980). Hence, the quantity of a urine deposit might be an unreliable quality for risk assessment. Animals using DPC for risk assessment should therefore be sensitive to those qualities, which signal risk reliably.

A common assumption regarding DPCs is that various behavioral adaptations to these cues, such as suppressed breeding (Fuelling and Halle, 2004), reduced activity, higher giving-up-densities (Apfelbach *et al.*, 2005) or increased vigilance (Monclus *et al.*, 2005; Monclus *et al.*, 2006), lead to increased prey survival. One mechanism that can increase survival chances of prey species is early predator detection because detected predators may abandon the hunt (Fitzgibbon, 1989; Lingle and Wilson, 2001) or because successful flight response of prey is facilitated (Krause and Godin, 1996).

We used a combined approach of observational data and experimental manipulation of DPC encounters in wild, free living meerkats (Suricata suricatta) to investigate which cue qualities affect the response to DPCs and how meerkats benefit from attending to DPCs. Meerkats are small, cooperatively breeding carnivores living in Southern Africa. They face high predation pressure by aerial and terrestrial predators (Clutton-Brock et al. 1999a) and have developed a coordinated sentinel (Clutton-Brock et al., 1999b; Manser 1999) and vigilance system (Townsend et al., 2011). The study population occupies farmland, where encounters with domestic animals including predators like domestic dogs (Canis lupus), and cats (Felis catus) have been documented (Kalahari Meerkat Project, long term data base, unpublished data). In response to predators they employ an elaborate spectrum of alarm calls encoding referential as well as motivational information (Manser et al., 2002). When a meerkat encounters a DPC such as cat, dog, caracal or bat-eared fox urine, feces or hair, it reacts immediately by emitting recruitment calls, whereas terrestrial alarm calls are emitted when a standing or moving predator on the ground is spotted (Manser, 2001). In response to recruitment calls, the rest of the group interrupts foraging and approaches the caller to inspect the cue (Manser et al., 2001). This gathering typically results in the group moving away from the cue to a different area before their onset of foraging again, or the group resuming foraging after several minutes in the vicinity of the cue.

We analyzed the natural frequency of recruitment events, and in the same population exposed meerkats to dog urine of different age and containing different quantities of urine. Finally, we tested experimentally if the reaction meerkats show towards a DPC leads to faster predator response. We predicted that meerkats should be sensitive to those qualities of DPC, which have the potential to reliably convey information to the receivers about risk, such as cue freshness, but not to qualities, which do not allow risk assessment, such as the amount of urine. Furthermore, we expected that DPC encounters facilitate the response of meerkats to alive predators by decreasing latency to emit an alarm call.

Material and Methods

The study animals

Observations and experiments were performed with wild meerkats at the Kalahari Meerkat Project (KMP), in the Kuruman River Reserve and on surrounding farm land in South Africa. Observations on recruitment behavior were analyzed for the period of January to December 2007. Experiments were conducted in August and September 2005, and from August 2006 to January 2007 (urine exposure experiment) and between June and August 2008 (predator detection experiment). The study site is located 30 km west of Van Zylsrus, in the southern part of the Kalahari desert (see Clutton-Brock *et al.*, 1999 a for detailed ecological description). The groups were habituated to human presence (closer than 1 m) and all group members were individually identified by unique dye marks (Jordan et al 2007). In total 38 DPC presentations in 12 groups (group size: median=20; range=5-32) were conducted for the urine exposure experiment and 14 cue presentations in 7 groups (group size: median=12; range=5-17) for the predator detection experiment.

The long-term observational data

To estimate the natural encounter rate of DPCs, we analyzed one year of adlib data from the long term data set of the KMP in eleven groups. Each group had been followed every week on at least three to four days for two to four hours during the morning foraging sessions and on two to four days during one to two hours in the afternoon foraging sessions, resulting in 6268 hours of observation (on average 570 per group; range 401 to 627 hours). According to the KMP protocol (Version 2006), every predator encounter and encounter of DPC or meerkat feces/scent that caused mobbing, inspection or recruitment was recorded. In most cases, we could not identify the actual cue but in other observations the meerkats behaved in similar manners regardless of whether the cue was secretions from glands, feces, carcasses, or hair hidden in the vegetation. From these observations we calculated the encounter rate resulting in recruitment to mob an alive predator, and inspect a scent (unidentified or identified) per observation hour per group.

The effect of age and quantity of a direct predator cue

In our experiments, we used DPCs from terrestrial predators that had previously been shown to elicit a recruitment response by meerkats (Manser, 2001; Graw and Manser, 2007). We used dog urine to test whether meerkats show a different response towards fresh and old urine or to different quantities (1ml or 4ml) of DPC. The dog urine was either added to the sand five to ten minutes before the presentation (below referred to as fresh urine), or 24 hours before the presentation and left outside exposed to the sun and outdoor temperature (referred to as old urine). To test whether the quantity of dog urine had an influence we presented samples of sand with fresh dog urine of the according amount. These experiments were conducted in a randomized order. The dog urine was presented to one randomly chosen adult focal individual and the response was filmed with a camera (Sony Digital Camcorder HDR-HC5). In case the focal individual did not react, the dog urine was presented to other randomly chosen individuals until an individual sniffed on the cue. The samples of sand with

the dog urine were presented on cardboard (8cm x 12cm) allowing easy and consistent handling with the sample. All the experiments were conducted when the group was foraging. In case of any predator alarm or other disturbances the experiments were postponed until the group was foraging uninterrupted for at least fifteen minutes. To avoid habituation we left a minimum interval of one week between experimental presentations within the same group.

In total we conducted 38 cue presentations in twelve different meerkat groups and averaged measures from the same group if the same stimulus type was presented. This resulted in a sample size of 28 cue presentations. We modeled the proportion of group members recruited, average vigilance and the latency to resume foraging (time period between first recruitment call and foraging onset afterwards) with linear mixed effect models assuming normal error structure with identity link function. These analyses were carried out in R, Version 2.14.0 (R Development Core Team, 2011) using the package lme4 (Bates *et al.*, 2011). The response variables were log or square root transformed and subsequently did not

Table 3. Model selection process for: a) Vigilance of recruited individu	als, and b) Latency to resume
foraging. Final models with the lowest AIC (bold) are presented in Table 2	

a) Vigilance of recruited individuals	AIC
	82.
Age of urine + Number of animals recruited + Amount of urine	48
	80.
Age of urine + Number of animals recruited	82
	89.
Age of urine	95
	92.
Number of animals recruited	23
b) Latency to resume foraging	
	149
Age of urine + Number of animals recruited + Amount of urine	.4
	150
Age of urine + Number of animals recruited	.1
	164
Age of urine + Amount of urine	.8
	151
Number of animals recruited + Amount urine	.6

differ from a normal distribution. To analyze the influence of cue age and cue quantity on the proportion of group members recruited we included both as fixed factors with two levels each (fresh/old and 1ml/4ml) and the group identity as a random factor. Examination of the AIC (Table 3) suggested that all terms should be retained in the final model. When modeling vigilance and the latency to resume foraging, we included the same fixed and random effects as in the previous model and additionally included the number of individuals recruited as a covariate. We then simplified models and dropped terms if it decreased the AIC (Table 3) until no more terms could be dropped. Terms which were not included in the final model are displayed in Table 2 with the values before they were excluded in the model selection process. Preliminary data analysis suggested that group size is not correlated with the proportion of group members recruited (Spearman, N=28, p=0.85), with vigilance (Spearman, N=25, p=0.98) and that the reproductive state of groups (presence of dependent young) does not influence the response to predator cues (Lienert 2007). Hence, these factors were disregarded in the model selection process. All figures presented in this paper are based on untransformed raw data.

The caracal detection experiment

Each meerkat group was tested to determine how fast they responded to a terrestrial predator once with a DPC in the experimental treatment, and once with the herbivore cue in the control treatment. As predator cue we used cat hair or bat-eared fox fur, which had been stored at -20 °C and defrosted a few hours before presentation. Using the same type of standardized DPC would have been the preferred option, but this was not possible due to practical limitations in access to cues in the field. However, both indicate the presence of a terrestrial predator and previous experiments have shown that these cues elicit qualitatively the same response in meerkats (c.f. Manser, 2001). As a control cue we used antelope hair *(Oryx oryx)* stored and presented in the same way as the DPC. In the experimental and control

treatment we placed the cue in the centre of the foraging group. As soon as one of the group members inspected the cue, we started to move a full-mounted caracal parallel to the group in an average distance of 78 m (range: 49 to 142 m, see Table 1). Caracals are sympatric predators and prey on a range of mammals of various sizes from rodents to medium sized antelopes (Melville *et al.*, 2004). Prior to presentation the dummy predator was hidden behind a camouflage fabric and was therefore invisible to the group. We measured the latency of predator detection defined as the time when the first meerkat began to inspect the cue (sniffing) until the first terrestrial alarm call was given by any of the group members, in response to the mounted caracal being moved. To control for order effects, half of the groups

Table 1. Physical conditions for the presentations of direct predator cues (DPC) and control cues (Control) in each meerkat group. Visibility estimated by naive observers (modal values), distance to the predator in meters measured by a rangefinder, and landscape type in which the experiments were performed are displayed.

	Visi	bility	Distan	Distance to predator (m) Landscape		
Group	DPC	Control	DP C	Control	DPC	Control
KU	good	good	142	87	Dunes Flats without	Dunes Flats without
F	good	good	51	49	bushes Flats without	bushes Flats without
D	medium	medium	94	95	bushes	bushes
AZ	poor	medium	93	78	Flats with bushes	Flats with bushes Flats without
W	poor	medium	80	64	Flats with bushes	bushes
L	poor	medium	90	56	Hills	Hills
CD	poor	poor	64	53	Flats with bushes	Flats with bushes

started with the experimental treatment, whereas the other half started with the control treatment.

We standardized, as much as possible, the distance between the group and the caracal, the visibility and the habitat structure in the experimental and control treatments. If impossible, we accepted a larger distance and poorer visibility in the experimental than in the control

treatment. This excluded the proximity or the visibility of the predator as an alternative explanation for faster predator detection (Table 1). Following an experiment, we took a photo with a digital camera (Konica Minolta Dimage X1) from the presentation spot towards the caracal. The camera was positioned at a standard height of 35cm, which corresponds to the height of the head of an adult meerkat standing on its hind legs. Later, these photos where shown to twelve naïve human observers, who were asked to score the visibility of the predator on a three-stage scale (good, medium, poor). At the time of the presentation, there were no meerkats acting as sentinels (look-out position at least 10 cm above ground) or meerkats emitting sentinel calls (Manser, 1999). We analyzed the latency to predator detection between the two treatments using an exact Wilcoxon test.

Results

Natural occurring recruitment events

In total, the eleven meerkat groups recruited group members 529 times in 6268 hours of observation during foraging (range: 36 to 77 recruitment events per group). This resulted in a recruitment frequency of one recruitment per 12.6 \pm 0.75 (range: 8.1-15.6) hours observation time (n=11 groups). On average 40.6 \pm 2.54 % were due to a predator that was encountered by one of the group members, who used the recruitment calls to initiate mobbing (c.f., Graw and Manser, 2007). In 53.2 \pm 2.82 % of the recruitment events, they were elicited by odors that could not be identified. Only 4 \pm 0.94% of all recruitment events were elicited by obvious identifiable feces of predators or conspecifics, and the rest (2 %) was caused by carcasses, body parts or artificial objects (e.g. bottles).

Effects of age and quantity of DPC on the recruitment

Presentations of dog urine elicited recruitment calls, which caused other individuals to approach the calling individual in 35 of 38 cases. Not all individuals inspected the cue, even

though a standardized distance and way of presentation to the different test animals was kept. However, the probability to inspect the cue was not dependent on the quality or quantity of the presented cue (fresh/1ml: 2.63 ± 0.53 times; fresh/4ml: 1.64 ± 0.48 times; old/4ml: 1.25 ± 0.57 times; GLMM, binomial distribution; age: estimate= 0.31 ± 0.54 , p=0.56; amount: estimate= 0.42 ± 0.50 , p=0.39). In three of the cases no individual approached the calling individual despite recruitment calls. On average 45 ± 4 (range: 0 to 100) % of the group members were recruited. Individuals which were recruited spent on average 22.06 ± 5.5 (range 0 to 127) seconds inspecting the cue and scanning the surroundings for predators until the groups after 122 ± 16.0 (range 0 to 399) seconds resumed normal foraging activity.

Table 2. Model parameters for a) proportion of the group recruited, b) vigilance of recruited individuals and c) latency to resume foraging. Final model parameters are displayed in bold. Terms which were not included in the final model are displayed with the p-value at which they were excluded from the model.

	Estimat			
	е	±SE	t-value	p-value
a) Proportion of group				
recruited				
Intercept	0.26	±0.06	4.47	
Age of urine	-0.23	±0.08	-2.91	0.004
Amount of urine	0.2	±0.08	2.59	0.01
b) Vigilance of recruited individua	als			
Intercept	1.13	±0.41	2.75	
Age of urine	-1.56	±0.37	-4.25	<0.001
Number of animals				
recruited	0.18	±0.03	5.25	<0.001
Amount of urine	0.21	±0.42	0.42	0.60
c) Latency to resume foraging				
Intercept	4.73	±1.18	4.01	
Age of urine	-1.75	±1.28	-1.36	0.19
Number of animals				
recruited	0.7	±0.11	6.18	<0.001
Amount of urine	0.38	±1.51	0.25	0.78



Figure 1: The proportion of the group recruited to dog urine (DPC) presentations (a) and (b), the mean vigilance per individuals in seconds (c) and (d), and the latency to resume foraging (e) and (f) plotted against cue age (a), (c), and (e) and cue quantity (b), (d), and (f). Asterisks indicate statistical significance (see Table 2 for model details).

Inspection behavior of the meerkats to the presented dog urine differed depending on the age and the amount of the cue. The proportion of individuals recruited to the cue increased with cue freshness and with cue quantity presented (Table 2a; Figure 1a and 1b). However, the time individuals were vigilant after they were recruited and inspected the cue was only enhanced by fresh, but not by larger amounts of urine. Additionally, vigilance time increased when more individuals were recruited to the cue (Table 2b and 3a; Figure 1c and 1d). Finally, the latency to resume foraging was significantly increased by fresh urine and tended to be high when large quantities of urine were presented (GLMM; age: estimate= -5.56 ± 1.9 , p=0.014; amount: estimate= 3.66 ± 1.91 , p=0.05). However, when controlling for the number of animals which were recruited, neither of these factors explained a significant proportion of variation (Table 2, Figure 1e and f) and model selection by AIC suggested that the covariate "number of animals recruited" predicted how long a group would interrupt foraging after predator cue encounters and should be retained in the model.

Predator detection experiment

The detection time of the caracal was strongly influenced whether the groups were presented a DPC or a control cue. When the predator cues were presented, the animals inspecting the cue always gave a few or several recruitment calls, whereas when encountering the control cue, they never called (Binominal: p=0.016; N=7). This caused the rest of the group to approach the caller in all the cases of the DPC, but never to the control cue (Binominal: p=0.016; N=7). The latency to the first alarm call in response to the presented caracal was shorter when the meerkat groups were exposed to a DPC compared to a control cue (exact Wilcoxon test: p=0.02; N=7; Figure 2). The individual who first emitted the alarm call to the caracal in the DPC treatment was the individual that had encountered the cue first only in two groups. In three groups it was another individual. For the other experiments (N=2) we were unable to determine the first caller. Only in one out of seven cases the individual closest to the predator alarmed first. In the control treatment, the individual that inspected the cue first was not, in any trial, the individual to emit the alarm call to the caracal first.



Figure 2: Latency to predator detection (in seconds) by meerkat groups (N=7) after exposure to a direct predator cue (DPC) or a control cue (Control). Asterisks indicate statistical significance.

Discussion

Response by meerkats related to cue qualities

During foraging trips, meerkats regularly encountered direct predator cues (DPC), to which they typically recruited other group members and inspected it together. Often the whole group interrupted foraging to inspect the cue. In our experimental study, the intensity of the response by meerkats to a DPC depended on the age, but less on the quantity of the cue. Meerkats inspected the cue and scanned their surrounding longer when the cue was fresher, but not when the quantity of the cue was increased. This effect was still present when statistically controlling for the number of recruited individuals, indicating despite the fact that the number of recruited animals predicted vigilance of each individual, cue age also affected vigilance.

Meerkats appeared to assess the enhanced danger of predation indicated by the fresh urine, and adjusted their behavior accordingly. The freshness of the cue might indicate that the predator, which had left the cue, was still in the vicinity. Thus, a fresh cue might be a valuable indicator denoting an increased risk of predation in this area, similar to what has been demonstrated experimentally in the wolf spider (Barnes *et al.*, 2002), and brushtail possums (Kirmani *et al.*, 2010).

The presentation of a larger quantity of urine induced the recruitment of a higher proportion of group members, but did not increase individual vigilance or the latency to resume foraging in comparison to the lower quantity. This suggests that the meerkats initially discriminated between lower and higher quantity of the DPC, but then may not have perceived the situation more risky due to higher quantity as observed with fresh urine. By recruiting group members, the reaction of the recruited individuals may help to assess the situation more accurately, yielding benefits associated with group decisions (Conradt and Roper, 2007). Such an effect is known from fish, which make faster and more accurate choices due to quorum decisions (Ward *et al.*, 2008; Ward *et al.*, 2011). Quorum decisions also play a crucial role for meerkats when coordinating group movement during foraging (Bousquet *et al.*, 2011). The notion that recruitment partly facilitates collective decisions regarding how to react to the current threat of predation is supported by the results of our presentation experiments. However, additional experiments manipulating recruitment and information transfer are needed to generate firm evidence.

The amount of urine may be a potential indicator of predator size and affect prey response. However, the large variation within individuals and between sexes, for example due

15

to differences in marking behavior (Hart, 1974; Jorgenson *et al.*, 1978; Natoli, 1985) might make it more difficult for prey species to correctly assess predator size based on the quantity of urine used in deposits. Furthermore, enhanced predator size might not always be the best indicator of danger for meerkats as observations suggest that medium sized terrestrial predators, such as jackals, are one of the main terrestrial predators of meerkats (Clutton-Brock et al. 1999a). Meerkats, being small animals with a body mass of less than one kilogram, fit much better in the prey range of medium sized predators than larger predators. Thus, meerkats may not benefit from discriminating different quantities of dog urine, and hence did not increase vigilance when presented with magnified cue quantities.

Predator response

Meerkats responded earlier to the presented moving terrestrial predator in their close vicinity, when being exposed to a DPC that indicated the presence of a terrestrial predator in comparison to a control cue of a herbivore. The latency to emit the first alarm call to the predator was shorter when the meerkats encountered a DPC compared to a control cue. In our experiment, meerkats typically emitted medium to high urgency terrestrial alarm calls referring to terrestrial predators (Manser, 2001), when detecting the mounted caracal (Zöttl, personal observations), suggesting that the mounted caracal was identified as a terrestrial predator.

Enhanced predator response due to the decreased latency to alarm in response to the dummy predator may have been caused by i) an increased sensitivity to the terrestrial predator, or ii) a general increased perceived risk, resulting in a higher vigilance after the exposure to the DPC. With our experiment we cannot distinguish between these two non-mutually exclusive alternatives. Evidence for increased sensitivity due to perceived risk or vulnerability on anti-predator has been found in the brushtail possum which reacts stronger to

a DPC in the absence of shelter (Parsons and Blumstein, 2010). It seems likely that increased vigilance during and after recruitment is the key to faster predator detection in meerkats. From our results, we can infer that due to the encounter of odor cues a faster response to the predator by the whole group was achieved. Whether the reduction in latency to alarm to the simulated predator was potentially caused by specific information available to the receiver about the type of danger in the acoustic structure of the recruitment calls, or a general heightened perceived risk due to the DPC and the recruitment calls, needs further experiments. A broad body of literature documents behavioral changes in response to predator odors (Apfelbach *et al.*, 2005), and numerous authors assume that animals increase their survival rates by reacting to DPCs (Berger *et al.*, 2001; Brown *et al.*, 2004; Monclus *et al.*, 2005; Ferrari *et al.*, 2006; Templeton and Greene, 2007; Blumstein *et al.*, 2008; Roth *et al.*, 2009; Webb *et al.*, 2010). However, our study provides the first experimental evidence showing a faster response, likely due to faster detection of a predator, which might help to increase survival (Fitzgibbon, 1989; Lingle and Wilson, 2001; Krause and Godin, 1996) after recruitment to direct predator cues.

The individual that recruited the group to the DPC was not always the first to give the alarm call in response to the dummy predator, suggesting that individuals benefit from recruitment rather than from the DPC encounter per se. Likely, the individual encountering the cue may be at the greatest risk to be close to the predator, if it is still in the area. By recruiting others to the spot, it dilutes the risk of being predated (Foster and Treherne, 1981; Uetz and Hieber, 1994; Roberts, 1996), and several together, may be able to deter the predator (Lingle, 2001; Graw and Manser, 2007). This would directly benefit the caller, but may not fully explain why the other group members should interrupt foraging. Rather, the additional observation that it was rarely the individual closest to the predator that gave the initial alarm call indicates that recruiting group members enables information transfer and as a

consequence every individual is aware of the magnified predation risk. This may allow the costs of anti-predator behavior to be shared among group members, which likely benefits all of them including the recruiter.

Conclusions

We show that meerkats interrupted foraging and were more likely to spot a predator during this interruption, whereby they attended to specific qualities of the cue, i.e., more to the age than the quantity of the cue. Age is likely a more reliable indicator about the immediate risk of the situation than the quantity, which may be affected by many other factors. To fully understand costs and benefits involved in recruitment calling, further research is needed to disentangle whether recruitment is a selfish behavior from both sides of the recruiter and the recruited individual, which results in a coordinated action, or whether it is a cooperative behavior incurring net costs to one of the involved parties.

Funding

This work was supported by a "Förderungstipendium" and a "KWA" from the University of Vienna to MZ covering travel costs to him, the Zoological Institute of the University of Zurich to MBM for all research expenses in the field due to this study, and Cambridge and Zurich University for financing the long term field project.

Acknowledgements

We are most grateful to the Kalahari Research Trust and the neighboring farmers of the Kuruman River Reserve, SA, in particular Family Koetze, to be able to work on their land. Many thanks to Anne-Sophie Blanc and Christophe Bousquet for help with the fieldwork, to Simon Townsend for helpful comments on the manuscript and to Matt Bell and Tom Flower for inspiring discussions. Also, thanks to the project leaders Rob Sutcliffe and Dave Bell and

all the volunteers for maintaining the meerkat groups to work on. Sabrina Engesser was a great help in analyzing the data from the long term data set.

Reference List

- Apfelbach R, Blanchard CD, BlanchardRJ., Hayes RA, McGregor IS. 2005. The effects of predator odors in mammalian prey species: A review of field and laboratory studies. Neurosci Biobehav Rev. 29: 1123-1144.
- Barnes MC, Persons MH, Rypstra AL. 2002. The effect of predator chemical cue age on antipredator behavior in the wolf spider Pardosa milvina (Araneae : *Lycosidae*). J Ins Behav. 15: 269-281.
- Barta Z, Liker A, Monus F. 2004. The effects of predation risk on the use of social foraging tactics. Ani Behav. 67: 301-308.
- Bates D, Maechler M, Bolker B. 2011. lme4: Linear mixed-effects models using S4 classes. (R package).
- Berger J, Swenson JE, Persson IL. 2001. Recolonizing carnivores and naive prey: Conservation lessons from Pleistocene extinctions. Science 291: 1036-1039.
- Blumstein DT, Barrow L, Luterra M. 2008. Olfactory Predator Discrimination in Yellow-Bellied Marmots. Ethology 114: 1135-1143.
- Bousquet CA, Sumpter DJT, Manser MB. 2011. Moving calls: a vocal mechanism underlying quorum decisions in cohesive groups. Proc R Soc B Biol Sci. 278: 1482-1488.
- Brown GE. 2003. Learning about danger: chemical alarm cues and local risk assessment in prey fishes. Fish Fisheries 4: 227-234.
- Clutton-Brock TH, Gaynor D, McIlrath GM, Maccoll ADC, Kansky R, Chadwick P, Manser MB, Skinner JD, Brotherton PNM. 1999a. Predation, group size and mortality in a cooperative mongoose, Suricata suricatta. J Anim Ecol. 68: 672-683.

- Clutton-Brock TH, O'Riain MJ, Brotherton PNM, Gaynor D, Kansky R, Griffin AS, Manser M. 1999b. Selfish sentinels in cooperative mammals. Science 284: 1640-1644.
- Conradt L, Roper T. 2007. Democracy in animals: the evolution of shared group decisions. Proc R Soc B Biol Sci. 274: 2317-2326.
- Devereux CL, Whittingham MJ, Fernandez-Juricic E, Vickery JA, Krebs JR. 2006. Predator detection and avoidance by starlings under differing scenarios of predation risk. Behav Ecol. 17: 303-309.
- Ferrari MC, Messier F, Chivers DP. 2006. The nose knows: minnows determine predator proximity and density through detection of predator odours. Anim Behav. 72: 927-932.
- Fitzgibbon CD. 1989. A Cost to Individuals with Reduced Vigilance in Groups of Thomsons Gazelles Hunted by Cheetahs. Anim Behav. 37: 508-510.
- Foster WA, Treherne JE. 1981. Evidence for the Dilution Effect in the Selfish Herd from Fish Predation on A Marine Insect. Nature 293: 466-467.
- Foust SJ, Thompson SA, Griswold SL, Gurrola EC, Kats LB. 2001. Response of leaf-cutting ants to predator chemical cues. Am Zool. 41: 1445-1446.
- Fuelling O, Halle S. 2004. Breeding suppression in free-ranging grey-sided voles under the influence of predator odour. Oecologia 138: 151-159.
- Gherardi F, Mavuti KM, Pacini N, Tricarico E, Harper DM. 2011. The smell of danger: chemical recognition of fish predators by the invasive crayfish Procambarus clarkii. Freshw Biol. 56: 1567-1578.
- Graw B, Manser MB. 2007. The function of mobbing in cooperative meerkats. Anim Behav. 74: 507-517.
- Hart BL. 1974. Environmental and Hormonal Influences on Urine Marking Behavior in Adult Male Dog. Behav Biol. 11: 167-176.

- Hilton GM, Cresswell W, Ruxton GD. 1999. Intraflock variation in the speed of escape-flight response on attack by an avian predator. Behav Ecol. 10: 391-395.
- Ito R, Mori A. 2010. Vigilance against predators induced by eavesdropping on heterospecific alarm calls in a non-vocal lizard Oplurus cuvieri cuvieri (Reptilia: Iguania). Proc R Soc B Biol Sci. 277: 1275-1280.

Jordan N.R., Cherry M.I. & Manser M.B. (2007) 'The spatial and temporal distribution of

meerkat latrines reflects intruder diversity and suggests a role of mate defence', Animal

Behaviour, 73, 613-622

- Jorgenson JW, Novotny M, Carmack M, Copland GB, Wilson SR, Katona S, Whitten WK. 1978. Chemical Scent Constituents in Urine of Red Fox (Vulpes-Vulpes L) During Winter Season. Science 199: 796-798.
- Kats LB, Dill LM. 1998. The scent of death: Chemosensory assessment of predation risk by prey animals. Ecoscience 5: 361-394.
- Kirmani SN, Banks PB, McArthur C. 2010. Integrating the costs of plant toxins and predation risk in foraging decisions of a mammalian herbivore. Oecologia 164: 349-356.
- Krause J, Godin JGJ. 1996. Influence of prey foraging posture on flight behavior and predation risk: Predators take advantage of unwary prey. Behav Ecol. 7: 264-271.
- Liberg O. 1980. Spacing patterns in a population of rural free roaming domestic cats. Oikos 35: 336-349.
- Lienert R. 2007. Secondary Predator Cue Inspection in Meerkats (Suricata Suricatta). MSc Thesis, University Zürich.
- Lima SL. 1998. Stress and decision making under the risk of predation: Recent developments from behavioral, reproductive, and ecological perspectives. Stress Behav. 27, 215-290.

- Lima SL, Bednekoff PA. 1999a. Temporal variation in danger drives antipredator behavior: The predation risk allocation hypothesis. Am Nat. 153: 649-659.
- Lima SL, Bednekoff PA. 1999b. Back to the basics of antipredatory vigilance: can nonvigilant animals detect attack? Anim Behav. 58: 537-543.
- Lima SL, Dill LM. 1990. Behavioral Decisions Made Under the Risk of Predation A Review and Prospectus. Canad J Zool. 68: 619-640.
- Lingle S. 2001. Anti-predator strategies and grouping patterns in white-tailed deer and mule deer. Ethology 107: 295-314.
- Lingle S, Wilson WF. 2001. Detection and avoidance of predators in white-tailed deer (Odocoileus virginianus) and mule deer (O-hemionus). Ethology 107: 125-147.
- Lohrey AK, Clark DL, Gordon SD, Uetz GW. 2009. Antipredator responses of wolf spiders (Araneae: Lycosidae) to sensory cues representing an avian predator. Anim Behav. 77: 813-821.
- Manser MB. 1999. Response of foraging group members to sentinel calls in suricates, Suricata suricatta. Proc R Soc B Biol Sci. 266: 1013-1019.
- Manser MB. 2001. The acoustic structure of suricates' alarm calls varies with predator type and the level of response urgency. Proc R Soc B Biol Sci. 268: 2315-2324.
- Manser MB, Bell MB, Fletcher LB. 2001. The information that receivers extract from alarm calls in suricates. Proc R Soc B Biol. Sci. 268: 2485-2491.
- Manser MB, Seyfarth RM, Cheney DL. 2002. Suricate alarm calls signal predator class and urgency. T Cog Sci. 6: 55-57.
- Melville HIAS, Bothma JD, Mills MGL. 2004. Prey selection by caracal in the Kgalagadi Transfrontier Park. South African J Wildlife Res. 34: 67-75.

- Monclus R, Rodel HG, Von Holst D, De Miguel J. 2005. Behavioural and physiological responses of naive European rabbits to predator odour. Anim Behav. 70: 753-761.
- Monclus R, Roedel HG, von Holst D. 2006. Fox odour increases vigilance in european rabbits: A study under semi-natural conditions. Ethology 112: 1186-1193.
- Morrison EB. 2011. Vigilance behavior of a tropical bird in response to indirect and direct cues of predation risk. Behaviour 148: 1067-1085.
- Natoli E. 1985. Behavioural Responses of Urban Feral Cats to Different Types of Urine Marks. Behaviour 94: 234-243.
- Parsons MH, Blumstein DT. 2010. Feeling Vulnerable? Indirect Risk Cues Differently Influence How Two Marsupials Respond to Novel Dingo Urine. Ethology 116: 972-980.
- Pulliam HR, Pyke GH, Caraco T. 1982. The Scanning Behavior of Juncos A Game-Theoretical Approach. J Theor Biol. 95: 89-103.
- R Development Core Team. 2011. R: A language and environment for statistical computing. R Foundation for Statistical Computing. Vienna, Austria., R Development Core Team.
- Ridley A, Raihani N, Bell M. 2010. Experimental evidence that sentinel behaviour is affected by risk. Biol Lett. 6: 445-448.
- Roberts G. 1996. Why individual vigilance declines as group size increases. Anim Behav. 51: 1077-1086.
- Roth TC, Cox JG, Lima SL. 2008. Can foraging birds assess predation risk by scent? Anim. Behav. 76: 2021-2027.
- Templeton CN, Greene E. 2007. Nuthatches eavesdrop on variations in heterospecific chickadee mobbing alarm calls. Proc Nat A Sci. 104: 5479-5482.
- Townsend SW, Zoettl M, Manser MB. 2011. All clear? Meerkats attend to contextual information in close calls to coordinate vigilance. Behav Ecol Sociobiol. 65: 1927-1934.

- Uetz GW, Hieber CS. 1994. Group-Size and Predation Risk in Colonial Web-Building Spiders -Analysis of Attack Abatement Mechanisms. Behav Ecol. 5: 326-333.
- Verdolin JL. 2006. Meta-analysis of foraging and predation risk trade-offs in terrestrial systems. Behav Ecol Sociobiol. 60: 457-464.
- Ward AJ, Herbert-Read JE, Sumpter DJ, Krause J. 2011. Fast and accurate decisions through collective vigilance in fish shoals. Proc Nat A Sci. 108: 2312-2315.
- Ward AJ, Mehner T. 2010. Multimodal mixed messages: the use of multiple cues allows greater accuracy in social recognition and predator detection decisions in the mosquitofish, Gambusia holbrooki. Behav Ecol. 21: 1315-1320.
- Ward AJ, Sumpter DJ, Couzin LD, Hart PJ, Krause J. 2008. Quorum decision-making facilitates information transfer in fish shoals. Proc Nat A Sci. 105: 6948-6953.
- Webb JK, Pike DA, Shine R. 2010. Olfactory recognition of predators by nocturnal lizards: safety outweighs thermal benefits. Behav Ecol. 21: 72-77.
- Wisenden BD. 2000. Olfactory assessment of predation risk in the aquatic environment. Phil. Trans R Soc. 355: 1205-1208.