

# Predation increases acoustic complexity in primate alarm calls

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**According to most accounts, alarm calling in non-human primates is a biologically hardwired behaviour with signallers having little control over the acoustic structure of their calls. In this study, we compared the alarm calling behaviour of two adjacent populations of Diana monkeys at Tai forest (Ivory Coast) and Tiwai Island (Sierra Leone), which differ significantly in predation pressure. At Tai, monkeys regularly interact with two major predators, crowned eagles and leopards, while at Tiwai, monkeys are only hunted by crowned eagles. We monitored the alarm call responses of adult male Diana monkeys to acoustic predator models. We found no site-specific differences in the types of calls given to eagles, leopards and general disturbances, but there were consistent differences in how callers assembled calls into sequences. At Tiwai, males responded to leopards and general disturbances in the same way, while at Tai, males discriminated by giving call sequences that differed in the number of component calls. Responses to eagles were identical at both sites. We concluded that Diana monkeys are predisposed to use their repertoire in context-specific ways, but that ontogenetic experience determines how individual calls are assembled into meaningful sequences.**

**Keywords:** predation; compositionality; vocal learning; evolution of language

## 1. INTRODUCTION

A long-standing hypothesis in animal communication states that, as signallers, non-human primates have little control over the acoustic structure of their call repertoire (Hammerschmidt & Fischer *in press*). Core evidence for the rigid nature of non-human primates' vocal behaviour comes from isolation and cross-fostering experiments, as well as some electrophysiological studies (Winter *et al.* 1973; Jürgens 1986; Owren *et al.* 1993; Hammerschmidt *et al.* 2001). Similarly, young vervet monkeys are predisposed to respond to aerial and terrestrial events with specific calls, while ontogenetic experience only affects the range of call-eliciting contexts (Seyfarth & Cheney 1986).

More recent work has shown that, in addition to the acoustic structure of calls, information can also be conveyed by the way individual calls are assembled into sequences. For example, white-handed gibbons (*Hylobates lar*) select from a limited set of song units

depending on whether they are singing to a terrestrial predator, a human observer or participating with their partner in a duet song (Clarke *et al.* 2006). King colobus (*Colobus polykomos*) and guereza monkeys (*Colobus guereza*) produce two basic alarm calls but assemble these meaningfully in predator-specific sequences (Schel *et al.* *in press*). Putty-nosed monkeys (*Cercopithecus nictitans*) produce two main alarm calls, which they assemble in predator-specific ways and also to signal forthcoming group travel (Arnold & Zuberbühler 2006, 2008).

In this study, we investigated the relationship between ontogenetic predator experience and alarm calling behaviour in West African Diana monkeys (*Cercopithecus diana*). We compared the vocal behaviour of free-ranging males at two sites, the Tai forest (Ivory Coast) and Tiwai Island (Sierra Leone). At Tai, the groups interacted regularly with leopards (*Panthera pardus*) and crowned eagles (*Stephanoaetus coronatus*; Zuberbühler & Jenny 2002; Shultz & Thomsett 2007), whereas at Tiwai, the groups only interacted with crowned eagles, as leopards have not been reported for at least 30 years (J. Oates 2007, personal communication).

## 2. MATERIAL AND METHODS

Data were collected in an approximately 100 km<sup>2</sup> area of Tai National Park (K.Z.: July 1994, June 1995, July to November 1996, January to June 1997, February 2000) and a 12 km<sup>2</sup> area of Tiwai Island (C.S.: February to May 2007) according to the same general protocol (Zuberbühler *et al.* 1997). Predator experiments were conducted in conjunction with continuous observations during which all vocal behaviours of adult male Diana monkeys, as well as their causes and consequences, were noted. Vocal responses of adult males were experimentally elicited by playing back predator vocalizations. For each trial, an unhabituated group was located, usually by acoustic cues, and approached as closely as possible. Recording distances were usually approximately 20 m. After positioning the speaker on a tree trunk or a fallen tree, the group was monitored for at least 20 min to ensure that the monkeys were unaware of the equipment and observer. Then, recording began with a 3–5 min pre-playback period, followed by a 10 min post-playback period. Playback stimuli consisted of a 15 s recording of crowned eagle shrieks or leopard growls, or a natural series of male Diana monkey alarm calls to a crowned eagle or a leopard.

On Tiwai Island, playback stimuli were broadcast using a CD player, connected to a Nagra DM speaker-amplifier. Vocal responses were recorded with a Sony WM-D6C recorder and Sennheiser ME80 microphone. At Tai, playback stimuli were broadcast with a Sony WM-D6C recorder, connected to a Nagra DSM speaker-amplifier. Responses were recorded with a Sony TCM-3000 recorder and Sennheiser ME80 microphone. After each trial, an area of a 500 m radius surrounding the location was not used for experiments with the same stimulus for at least two weeks. Trials were excluded from analysis if (i) the focal male was not with the group, (ii) two neighbouring groups responded to the playback stimuli, and (iii) local vegetation or technical problems prevented sufficient recording quality.

Recordings were digitized using COOL EDIT 2000 (Syntrillium Software Corporation, Phoenix, USA) and submitted to acoustic analyses using PRAAT 4.4.33 (Boersma & Weenink 2003). Diana males produce two basic call types in response to a range of disturbances (Riede & Zuberbühler 2003). The two call types differ most strikingly with regard to the presence of formant transitions, an acoustic feature that can be discriminated by ear (figure 1). Males often produce acoustically identifiable inhalations between subsequent calls, which we also included in the analyses. Males rarely give calls singly, but usually produce them as part of longer series of varying numbers. For each response, we measured the (i) number of calls produced of each type, (ii) number of call series, and (iii) number of calls of each type per series. Non-parametric statistical comparisons (Mann–Whitney *U*-tests, two-tailed) were made using SPSS v. 13.0.

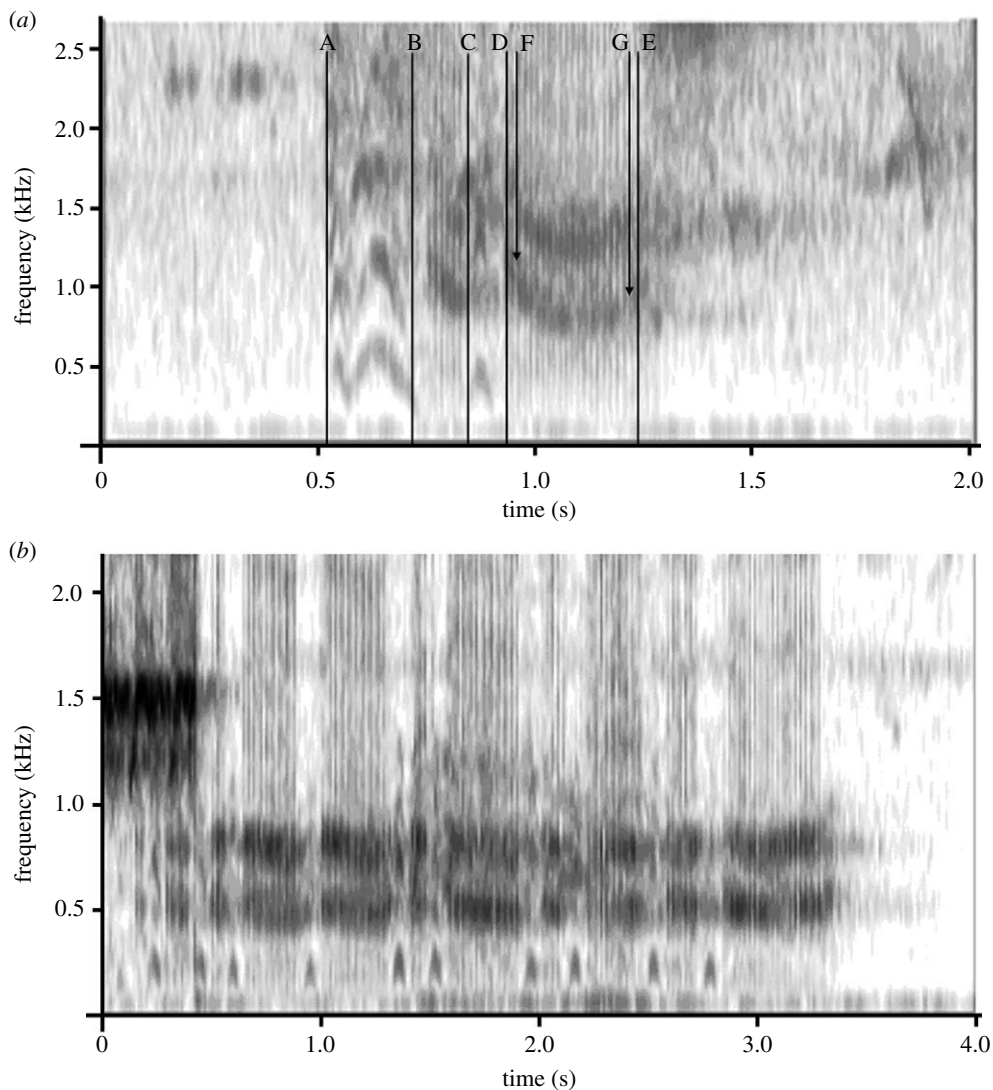


Figure 1. Spectrographic representation of a male Diana monkey alarm call sequence on Tiwai island in response to (a) leopard growls and (b) eagle shrieks. A–B and C–D, inhalations; B–C and D–E, exhalations; F–G, frequency transition.

### 3. RESULTS

#### (a) Acoustic features of alarm calls

Male Diana monkeys responded with the same basic alarm call types at both sites. Leopard growls and male alarm calls to leopards triggered call sequences with strong frequency transitions, and many voiced call inhalations, while eagle shrieks and male alarm calls to eagles triggered call sequences with only weak transitions (table 1).

#### (b) Organization of alarm call responses

There were no differences in the overall number of alarm calls produced at the two sites (leopard growls:  $N_{\text{Tai}}=12$  males,  $N_{\text{Tiwai}}=11$  males;  $N$  calls:  $U=46.0$ ,  $p=0.235$ ; leopard alarms:  $N_{\text{Tai}}=10$  males,  $N_{\text{Tiwai}}=7$  males;  $N$  calls:  $U=20.0$ ,  $p=0.601$ ; eagle shrieks:  $N_{\text{Tai}}=11$  males,  $N_{\text{Tiwai}}=6$  males;  $N$  calls:  $U=49.0$ ,  $p=0.660$ ; eagle alarms:  $N_{\text{Tai}}=11$  males,  $N_{\text{Tiwai}}=6$  males;  $N$  calls:  $U=73.0$ ,  $p=0.062$ ).

However, there were significant differences at the two sites with regard to how males assembled their calls into sequences. Although they had no prior experience with leopards, Tiwai males produced significantly more calls per call series in response to

leopard growls than Tai males ( $N_{\text{Tai}}=12$ ,  $N_{\text{Tiwai}}=11$ ,  $U=25.0$ ,  $p=0.011$ ). The same trend was found in these males' responses to other males' alarm calls to leopards, although the difference was not significant ( $N_{\text{Tai}}=10$ ,  $N_{\text{Tiwai}}=7$ ,  $U=19.0$ ,  $p=0.133$ ). In response to eagle-related stimuli, we found no differences in terms of how calls were assembled into sequences (eagle shrieks:  $N_{\text{Tai}}=11$ ,  $N_{\text{Tiwai}}=6$ ,  $U=25.0$ ,  $p=0.591$ ; eagle alarms:  $N_{\text{Tai}}=11$ ,  $N_{\text{Tiwai}}=6$ ,  $U=53.0$ ,  $p=0.961$ ).

At Tai, males discriminate acoustically between their responses to leopards and general disturbances, such as falling trees or fleeing duikers (Zuberbühler *et al.* 1997), whereas at Tiwai, males also responded regularly to such general disturbances, but these call sequences were not different from the ones given to leopard-related stimuli ( $N_{\text{leopard}}=11$ ,  $N_{\text{unspecific}}=6$ ,  $U=21.0$ ,  $p=0.256$ ;  $N_{\text{leopardalarm}}=7$ ,  $N_{\text{unspecific}}=6$ ,  $U=15.5$ ,  $p=0.445$ ; figure 2).

### 4. DISCUSSION

Our study provides evidence that differences in predation pressure affect the vocal behaviour of non-human primates. Although we found no differences in the

Table 1. Median percentages of different alarm call types given to leopard- and eagle-related stimuli at Tiwai Island and Tai Forest. (Statistical comparisons between study sites (Mann–Whitney  $U$ -tests, two-tailed) are as follows. Leopard growls—exhalations, transitions:  $U=27$ ,  $p=0.437$ ; exhalations, no transitions:  $U=36$ ,  $p=1$ ; inhalations:  $U=33$ ,  $p=0.82$ . Leopard alarms—exhalations, transitions:  $U=9$ ,  $p=0.022$ ; exhalations, no transitions:  $U=30$ ,  $p=1$ ; inhalations:  $U=0$ ;  $p=0.000$ . Eagle shrieks—exhalations, transitions:  $U=32.5$ ,  $p=0.961$ ; exhalations, no transitions:  $U=26.5$ ,  $p=0.525$ ; inhalations:  $U=0$ ,  $p=0.000$ . Eagle alarms—exhalations, transitions:  $U=33$ ,  $p=1$ ; exhalations, no transitions:  $U=16$ ,  $p=0.098$ ; inhalations:  $U=9$ ,  $p=0.015$ .)

	playback type			
	leopard growls	leopard alarms	eagle shrieks	eagle alarms
Tai Forest	$N=12$ males	$N=10$ males	$N=11$ males	$N=11$ males
call inhalations	56.4	57.1	50.0	52.0
call exhalations, transitions	43.7	42.9	4.2	6.3
call exhalations, no transitions	0	0	45.8	41.7
Tiwai Island	$N=11$ males	$N=7$ males	$N=6$ males	$N=6$ males
call inhalations	55.3	54.2	47.2	44
call exhalations, transitions	44.7	45.8	3.8	4.3
call exhalations, no transitions	0	0	49.0	51.7

acoustic structure of individual alarm calls, Diana monkey males differed consistently and significantly in how they assembled individual calls into sequences.

We found no site-specific differences in the males' calling behaviour to crowned eagles, both in terms of acoustic structure and call sequencing. Crowned eagles are present at both sites and their impact as primate predators is well documented (Shultz & Thomsett 2007). By contrast, we observed consistent differences in how the monkeys responded to leopard-related stimuli. At Tai, monkeys interact regularly with leopards, and males typically produce call sequences consisting of a small number of calls, usually introduced by voiced inhalations. Males also produce the same call types to a variety of general disturbances, such as falling trees, fleeing ungulates and other sudden loud noises, but then they consistently assemble them into significantly longer sequences (Zuberbühler *et al.* 1997). At Tiwai, leopards have not been seen for at least 30 years, but crowned eagles are regularly present, and the males' calling behaviour reflected this fact. By contrast, although Tiwai males reliably responded to leopard-related stimuli, they produced calling sequences that were indiscriminate from the ones given to general disturbances, such as falling trees.

We concluded that these monkeys were biologically predisposed to produce acoustically distinct alarm calls to aerial and terrestrial disturbances, confirming earlier empirical work and current theory (e.g. Seyfarth & Cheney 1986; Hammerschmidt & Fischer *in press*). However, Diana monkeys did not produce their alarm calls singly but instead assembled them into larger sequences, which added an additional layer of complexity. At both sites, males produced long sequences to eagles and general disturbances, but only in Tai did males show evidence of discriminating vocally between leopard-related stimuli and general disturbances. By contrast, if Tiwai males heard a playback of leopard growls, or short-sequenced leopard alarm calls, they responded as if they had perceived a general disturbance (figure 2).

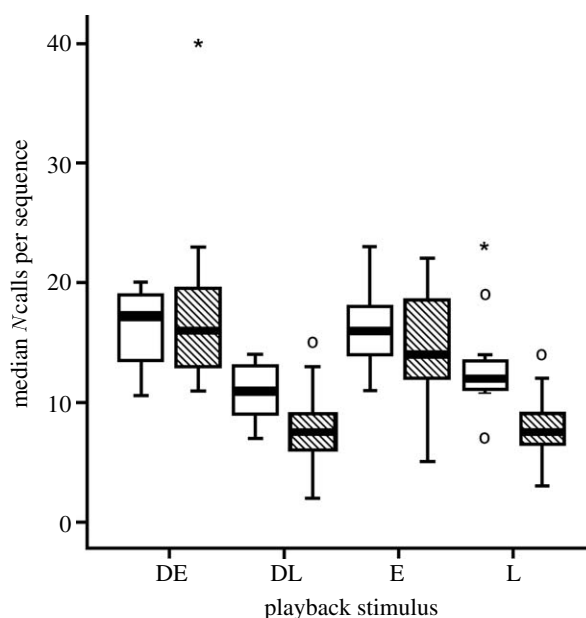


Figure 2. Median numbers of calls per sequence of Tai (hatched boxes) and Tiwai (open boxes) Diana monkeys to playbacks of eagle shrieks (E), leopard growls (L), male Diana alarm calls to eagles (DE) and male Diana alarm calls to leopards (DL). Box plots show median values, quartiles, range and outliers. Circles and asterisks show extreme values, asterisks being more extreme.

These differences in vocal behaviour are unlikely to be the result of genetic differences between the two populations; the time of isolation has been much too short. More likely, these are reflections of differences in ontogenetic history, particularly of growing up without a key predator, suggesting that Tiwai males would learn to discriminate between leopards and general disturbances, and mark these differences acoustically if the habitat changed accordingly (Berger *et al.* 2001).

Ontogenetic studies on vervet monkeys and meerkats have shown that antipredator behaviour becomes increasingly more complex as individuals mature and gain experience (Seyfarth & Cheney 1980; Hollen & Manser 2006). How exactly experience with predators influences this process is largely unknown, but social learning is likely to play an important role (Curio



*et al.* 1978). Most accounts of primate communication accept the notion that individuals can modify the timing and duration of calls (Hammerschmidt & Fischer in press), but how individuals use this flexibility offered in the temporal domain to encode the meaning has not been explored systematically (e.g. Arnold & Zuberbühler 2006, 2008).

In our study, we found no flexibility in the acoustic fine structure of individual calls, but significant flexibility in the way calls were organized into context-specific sequences. The two populations were identical in all relevant ecological and social parameters, apart from the presence of leopards, a key predator of monkeys. The most reasonable explanation for the differences in calling behaviour is, therefore, that leopard predation has led to increased complexity in the calling behaviour of Tai but not Tiwai monkeys.

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