Changes in concentration of serum prolactin during social and reproductive development of the spotted hyaena (Crocuta crocuta)

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Summary. A homologous radioimmunoassay system developed for humans was used to measure prolactin concentrations in spotted hyaenas. Concentrations of prolactin showed a significant ($P < 0.05$) decrease in lactating females, which is consistent with the infrequent suckling pattern of this species. This lack of hyperprolactinaemic conditions during lactation may explain the ability of females to resume reproductive activity soon after the loss of a litter, or even during lactation. Prolactin concentrations did not increase significantly during dispersion in male spotted hyaenas. This conforms to the pattern observed for cortisol, but differs from that for androgen, which fluctuates significantly with social suppression. Although comparative data from other species provide some circumstantial evidence for hyperprolactinaemic conditions during male dispersal, no obvious deductions regarding the recorded inverse relationship between prolactin and cortisol concentrations in mature males could be made.

Keywords: prolactin; spotted hyaena; lactation; dispersal

Introduction

The social unit, or clan, of the spotted hyaena (Crocuta crocuta) comprises a variable number of closely related matrilocal lineages (Mills, 1985; Frank, 1986; Henschel & Skinner, 1987) and a multimale group that consists of males in various stages of immigration or emigration (Frank, 1986; Henschel & Skinner, 1987). Females are generally philopatric and remain in their natal clans to breed; males tend to disperse at about puberty and join neighbouring clans, where they may have opportunities to mate. This emigration is not without risk, as the acceptance of a male into a new clan may take up to several months, during which time antagonistic encounters with members of the target clan can result in death (Kruuk, 1972; Henschel & Skinner, 1987). Females are generally the dominant social class, followed by their female cubs and then resident natal males that have not yet dispersed, central immigrant or mating males and peripheral immigrant males that are in the process of dispersing (Henschel & Skinner, 1987). This social structure results in the social suppression of concentrations of androgen in plasma of most males, with only the central immigrant males that have been accepted into a target clan having increased androgen titres (van Jaarsveld & Skinner, 1991a).

Spotted hyaenas are considered the only precocial fissiped carnivores (van Jaarsveld et al., 1988). They usually produce two young, which are dependent on milk for 4–6 months and are not weaned until 14–18 months old (East et al., 1989). Although this extensive lactation period suggests a low potential for population recruitment (Henschel, 1986), females can resume breeding soon after the loss of a litter, through either death or weaning (Grimpe, 1916; Golding, 1969; Henschel & Skinner, 1990) and even while still lactating (Schneider, 1926; Kruuk, 1972). Histological evidence as well as changes in concentrations of plasma progesterone and oestradiol suggest that some follicular development may occur during lactation in this species (van Jaarsveld et al., 1992). This
provides a mechanism whereby reproductive activity may be initiated during lactation or soon after the loss of a litter in this aseasonal breeder (Lindeque & Skinner, 1982).

This investigation evaluated the roles played by prolactin as a stress related hormone (Drago et al., 1989) during male dispersal and as a hormone involved in regulating ovarian activity during lactation in most mammals (McNeilly, 1987).

Materials and Methods

Animals

Twenty-eight spotted hyaenas (>1 year old) with established social histories were sampled between 1984 and 1987 from (a) the Mavumbye and Shingkelengane clans that are resident in the central district of the Kruger National Park, South Africa (24°20'S, 31°45'E) (Henschel & Skinner, 1987, 1989), (b) the Letaba clan (23°56'S, 31°40'E) resident around the Letaba Rest Camp in the Kruger National Park and (c) the Kousant clan that lives on the boundary between the Kalahari Gemsbok National Park (South Africa) and the Gemsbok National Park (Botswana) (25°15'S, 20°30'E). This clan is the subject of a long-term project on behavioural ecology (Mills, 1985).

In addition, 12 cubs (<1 year old) were sampled from various localities in the Kruger National Park and from the National Zoological Gardens (Pretoria) in South Africa. This was an effective way of increasing the sample size for the younger age group, as they do not disperse before attaining reproductive maturity (Kruuk, 1972; Frank, 1983; Mills, 1985; Henschel & Skinner, 1987).

Animals were grouped into one of the following social categories, according to Henschel & Skinner (1987): female cubs (<1 year), resident females (>1 year), male cubs (<1 year), resident natal males, peripheral immigrant males or central immigrant males. Reproductive status was assessed through external examination according to the criteria specified by Matthews (1939) and animals were grouped as belonging to one of the following reproductive categories: nulliparous female, parous female, lactating female, immature male (<2 years) and mature males (>2 years). Ages of animals of unknown age were estimated from the height and surface area of the mandibular premolar (PM3) (van Jaarsveld et al., 1987).

Free-ranging and captive hyaenas were immobilized with Zoletil (about 4 mg kg \(^{-1}\) body weight, CI-744: Anchorpharm Pty Ltd, Bramley, S. Africa) following van Jaarsveld (1988). As all animals were the subjects of long-term field studies or in captivity, they were habituated to vehicles and human disturbance. The only stressors they were subjected to were therefore the mild response to the sting of the small air-powered dart (1 ml, Telinject S.A., Randburg, S. Africa) and the subsequent loss of consciousness owing to the anaesthetic effects of Zoletil. The behaviour of animals strongly suggested that the darting was not a traumatic experience, whereas the loss of consciousness was. Extended anaesthesia (90 min) for serial blood sampling was achieved by administering halothane (Fluothane:ICI Pharmaceuticals Ltd, Johannesburg, S. Africa) using a circle absorber machine. Induction was achieved with a 10% mixture of halothane in oxygen and the animals were maintained using 2-4% halothane in a closed circuit. This method was preferred as it provided a more stable plane of anaesthesia than other immobilizing chemicals. Clinical procedures followed van Jaarsveld et al. (1984).

Twenty-two adult males were subjected to serial blood sampling for the duration of anaesthesia. Blood samples (approximately 10 ml) were collected from the cephalic vein using multisample needles and venoject evacuated tubes. Blood was stored at 4°C until it was centrifuged for 10 min at 1000 g. Serum was stored at -20°C until assayed. Rectal temperature was monitored intermittently and, if temperature was falling, hot water bottles and solar blankets were provided. The other animals were only subjected to one-off sampling after immobilization.

Prolactin radioimmunoassay

Concentrations of prolactin were determined using a heterologous double-antibody radioimmunoassay kit (IM. 1061; Amersham Laboratories, Bucks, UK), as prolactin standards or pituitary glands of spotted hyaenas were unavailable. The assay uses ovine anti-human prolactin antiserum, human pituitary standards (0, 5, 15, 50, 100 and 200 ng ml \(^{-1}\)) in human serum, \(^{125}\)I-labelled human prolactin and donkey anti-sheep gamma globulin as second antibody. All freeze-dried controls were reconstituted to specified volumes using water distilled into glass. The incubation mixture consisted of reference preparations (100 μl) or serum samples (200 μl), antiserum (100 μl) diluted to 5-5 ml and \(^{125}\)I-labelled human prolactin (approximately 10 000 c.p.m. in 100 μl). The mixture was then incubated at room temperature (±20°C) for 24 h. Donkey anti-human gamma globulin (1 ml – second antibody) was then added and incubation continued at room temperature for 5 min before separation of bound and free hormone by centrifugation at room temperature (1500 g) for 15 min. The free iodinated prolactin in the supernatant was decanted into scintillation vials and 4 ml scintillation fluid (Scintillator 299RS; Packard Instrument Co., IL, USA) was added. Radioactivity was measured 4 h later for 2 min, using a Packard 1500 Tri-Carb scintillation counter (Packard Instrument Co., IL, USA). As no spotted hyaena standards were available, the prolactin concentrations were expressed as ng human prolactin equivalents ml \(^{-1}\) serum. Mathematical interpolation of recorded sample values against a standard curve was carried out using Securic® Plus RIA/QC software (Packard Instrument Co., Downers Grove, UK) over the range 5–200 ng ml \(^{-1}\).
Prolactin status. Two indices of prolactin status were evaluated, namely, the initial sample irrespective of the time since immobilization, and the total prolactin response, represented by the area beneath the prolactin response curve from the time of immobilization for 90 min, following the rationale of Brown et al. (1988).

Validation of prolactin assay. Three spotted hyenas from the Kruger National Park were maintained on halothane for 4 h after immobilization with Zoletil. They were injected with $1 \mu g \, kg^{-1}$ luteinizing-hormone-releasing hormone (LHRH: Hoechst, Frankfurt, Germany) intravenously after 90 min. Plasma samples for androstenedione determination were collected at intervals of about 15 min for the total duration of anaesthesia. Three spotted hyenas from the National Zoological Gardens were immobilized with Zoletil and maintained on halothane for about 2 h on two occasions. During each of these extended immobilizations, the animals were injected intravenously with either bromocriptine ($1 \, mg \, kg^{-1}$ body weight; Sandoz Products (Pty) Ltd, Randburg, S. Africa) or metoclopramide ($1 \, mg \, kg^{-1}$ body weight; Beechams, Betchworth, Surrey, UK) after stabilizing for 30 min.

Specificity. The method proved to be acceptable for measuring prolactin concentration in spotted hyenas. The binding of $^{125}$I-labelled prolactin to the antiserum was displaced in a parallel manner by serial dilutions of serum from a mature male and a lactating female hyena (Fig. 1a). The addition of 200 µl serum from a male treated with bromocriptine to the standard curve did not affect the binding (Fig. 1a) and all samples were therefore assayed against the human standards. Measured concentrations of prolactin in serum were not affected by volume (100–200 µl). The injection with LHRH had no significant effect on concentrations of serum prolactin (Fig. 1b), whereas androstenedione, the principal steroid secreted from the ovary after stimulation with LHRH (van Jaarsveld & Skinner 1991b), showed a marked positive response (Fig. 1c).

![Graphs](image)

**Fig. 1.** (a) Standard curve for human prolactin prepared in distilled water (■) with and (●) without 200 µl serum from a male spotted hyena treated with bromocriptine and (▲) dose–response curves from dilutions of spotted hyena serum; (b) concentrations of serum prolactin in three spotted hyenas after an injection (arrow) of $1 \, \mu g$ luteinizing-hormone-releasing hormone (LHRH) kg$^{-1}$ body weight and (c) androstenedione responses from the ovary of three females after injection (arrow) with $1 \, \mu g$ LHRH kg$^{-1}$ body weight.

Accuracy. The recovery of known amounts of human prolactin (15–100 ng) added to the serum of a spotted hyena treated with bromocriptine did not differ significantly from expected values ($t_{31} = 1.81; P > 0.2$).

Sensitivity and precision. Sensitivity was $0.76 \pm 0.23 \, \text{ng prolactin ml}^{-1} \, \text{serum}$. The proportion of total radioactivity bound by the anti-serum in the absence of hormone ranged from 48.38 to 54.04%. The intra- and interassay coefficients of variation were 2.04 and 14.12%, respectively.

Biological validation. All three treated spotted hyenas showed a significant increase ($t_{31} = -37.39; P < 0.01$) in serum prolactin concentrations after the injection of metoclopramide, the effect being significantly greater in the immature male than in a nulliparous female and a mature male. A significant decrease ($t_{31} = -39.65; P < 0.01$) was seen after the injection of bromocriptine (Fig. 2).
Androstenedione radioimmunoassay

Plasma androstenedione was assayed using the same chemical reagents, and following procedures described by van Jaarsveld & Skinner (1991b). Plasma aliquots of 0.1 ml were assayed using antisera raised in rabbits against androstenedione-7-hemisuccinate bovine serum albumin (Miles Yeda, Kiryat Weizman, Rehovot, Israel). Cross-reactions with other steroids were 32% for 5-androstane-3,17-dione, 3% for testosterone and dehydroepiandrosterone, 0.6% for 11-deoxycorticosterone and progesterone, 0.2% for oestrone and <0.01% for oestradiol. Sensitivity of the assays ranged from 3.92 to 16.70 pg ml\(^{-1}\) (\(x = 5.98 \pm 3.63\) sd; \(n = 7\)). Intra- and interassay coefficients of variation were 7.31 and 5.28%, respectively.

Statistical analyses

All statistical procedures used were in accordance with the statistical principles described by Sokal & Rohlf (1980). Statistical manipulations were carried out using SAS Institute Inc. (Illinois, USA) software. The relevant procedures used are specified with the results. The frequency distributions of all data sets were tested for normality, and variances for homoscedasticity. When raw data failed to comply with the required criteria for parametric analyses, data transformations were used to obtain normality and/or homoscedasticity. When data sets or subsets still failed to comply with parametric criteria, nonparametric statistical procedures were used.

Results

Correlation analysis showed that there was a strong positive rank correlation (Spearman - \(r_s = 0.78; P < 0.001; n = 22\)) between the initial sample and the total prolactin response (Fig. 3). In 12 (55%) of the animals subjected to serial sampling, the highest prolactin concentrations recorded were the initial samples. Of the ten animals that showed subsequent increases, six (27%) were represented by the lactating females in the sample, all of which tended to show a general increase after starting from low initial values (Fig. 3b). Other animals that showed subsequent increases in serum prolactin concentrations after the initial sample included two resident natal males (Fig. 3d), one peripheral immigrant male (Fig. 3e) and a parous female (Fig. 3a). This tendency to increase and decrease after the first sample showed no relationship to the time of first sampling, as individuals that showed an increase were, on average, first sampled after those that showed a decline (\(x = 18.90\) min compared with \(x = 3.82\) min). Visual inspection of the profiles also showed that males generally displayed a larger variance than females in prolactin concentrations after immobilization stress.

The initial values were used as an index of prolactin status and a parametric ANOVA (PROC GLM) comparing \(\log_{10}(\text{prolactin})\) showed no significant difference (\(F_{0.05[1.55]} = 4.60\)) between
sexes or with respect to the month of the year that samples were collected ($F_{0.05[1, 55]} = 4.52$). This lack of a seasonal pattern is further supported by the presence of the highest and lowest recorded concentrations of serum prolactin in the same month (1.34 versus 50.80 ng ml$^{-1}$ in November). A similar analysis for the various reproductive ($F_{0.05[4, 55]} = 15.79$) and social ($F_{0.05[5, 53]} = 3.65$) categories showed significant variation. A comparison of means revealed that lactating females had significantly lower (HSD$_{51} = 3.99, P < 0.05$) prolactin concentrations than animals in all the other reproductive categories. Parous females, however, showed a lower mean prolactin concentration than the remaining categories but the difference was not statistically significant (Fig. 4a). Among the social categories, the peripheral immigrant males, the male cubs and the female cubs had high mean prolactin concentrations, but these differences did not reach statistical significance, and only the male cubs had significantly higher (HSD$_{48} = 4.197, P < 0.05$) mean titres than the resident females (Fig. 4b).

No significant difference was found between male and female cubs in mean serum prolactin concentrations (Fig. 4b) or in the observed trends during the first year of development (Fig. 5a, b). More important, however, is that there is an indication of reduced prolactin titres during this developmental phase, as was the case for cortisol (van Jaarsveld, 1990). Correlation analysis revealed a negative relationship between age and initial prolactin concentrations (Pearsons $r = -0.28, P < 0.05; n = 56$), reflecting the high prolactin concentrations observed in cubs (Fig. 4b).
Fig. 4. Serum human prolactin equivalents (mean ± SEM) for (a) reproductive categories: mature males (MM; > 2 years), immature males (IM; ≤ 2 years), lactating females (LF), nulliparous females (NF), parous females (PF) and (b) social categories: central immigrant males (CIM), peripheral immigrant males (PIM), resident natal males (RNM), male cubs (MC), female cubs (FC), and resident females (RF). ANOVA was run on log10[prolactin]. Statistical significance was measured at $P < 0.05$. Numbers above columns indicate sample size.

Fig. 5. Serum prolactin profiles for (a) female ($n = 4$) and (b) male ($n = 7$) spotted hyaena cubs. Different animals are represented by different symbols; serial samples from the same animals are connected.

**Discussion**

Although it is generally accepted that prolactin concentration increases during acute stress, this is not always the case; a decrease in circulating prolactin concentrations often occurs when concentrations have already been raised (Krieg et al., 1984; Gala & Haisenlander, 1986), or during chronic stress (Collu et al., 1979; López-Calderon et al., 1989). Thus, the general trend of declining prolactin concentrations observed in animals subjected to serial sampling in this study suggests that some prolactin stress response to immobilization had already occurred by the time the first sample was collected. The mean time of first sampling for these serial samples was 20-23 min and rats are known to show increased concentrations of prolactin 20-45 min after immobilization stress (López-Calderon et al., 1989). The initial samples used in the present study therefore probably represent stress-induced concentrations rather than basal concentrations.

The lack of any significant increase in the initial prolactin concentrations in the dispersing peripheral immigrant males (Fig. 4b), which are usually at the bottom of the social hierarchy (Henschel & Skinner, 1987), is similar to the pattern found for cortisol concentrations in plasma (van Jaarsveld, 1990). These data conform with earlier work on captive talapoin monkeys (*Miopithecus talapoin*)
in which no correlation between male social rank and prolactin concentrations could be found (Yodingyuad et al., 1982). Although social status was not measured in this study, serum prolactin concentrations did not differ in dominance-related social categories, clearly reflecting the effects of social suppression on circulating concentrations of androgen in males (van Jaarsveld & Skinner, 1991a).

A negative correlation between plasma cortisol and serum prolactin in mature male spotted hyaenas (van Jaarsveld, 1990) suggests that these hormones may act in a stress-related manner in this species. This relationship could indicate a negative feedback between adrenocortical activity and prolactin stress responsiveness, a process documented in rats (López-Calderon et al., 1989). On the other hand, it could be the result of the different control mechanisms of these stress-related hormones (Naylor et al., 1990), or that these hormones respond in different ways to social modifications and that they do not simply reflect a single intervening variable, such as stress (Eberhardt et al., 1985). Some support for the view that these hormones are regulated independently comes from the observation that the apparent cortisol stress nonresponsive period observed in juvenile spotted hyaenas (van Jaarsveld, 1990) is not accompanied by a change in prolactin-stress responsiveness. However, if prolactin primarily functions as a protective factor in stress-induced biological changes (Drago et al., 1989), high prolactin concentrations would complement the unperturbed concentrations of cortisol found during the cortisol stress nonresponsive period and ensure normal central nervous system and behavioural development of juvenile animals (Sapolsky & Meaney, 1986). This line of reasoning and the observed negative relationship between these two hormones in adult males does not rule out a negative feedback relationship between cortisol and prolactin.

The similarity between the concentrations of prolactin in peripheral immigrant males and male and female cubs (Fig. 5) indicates a form of hyperprolactinaemia in dispersing males. These concentrations are similar to those found in prepubertal pigs (Sus scrofa) and blue foxes (Alopex lagopus) in seasonal anoestrus (Mondain-Monval et al., 1985; Smith et al., 1985). These high prolactin concentrations (about 12 ng ml\(^{-1}\)) suggest that the testicular inactivity of dispersing males (van Jaarsveld & Skinner, 1991) and the gonadal inactivity of prepubertal spotted hyaenas (Lindeque, 1981) may be correlated with hyperprolactinaemic conditions (McNeilly, 1980; Haynes & Howles, 1981). The decline in mean serum prolactin titres observed in the resident natal males, between puberty and dispersion, is accompanied by a transition from the prepubertal to pubertal testis. In support of this view, the positive correlation recorded between prolactin and androstenedione concentrations in male cubs changes to a positive correlation between prolactin and testosterone in the resident natal males (van Jaarsveld, 1990).

Among the reproductive categories, lactating spotted hyaenas are prominent for their low serum prolactin concentrations (Fig. 4a). These low prolactin concentrations in lactating animals follow the pattern previously observed in rabbits (Oryctolagus cuniculus) and brown hares (Lepus europaeus) (McNeilly & Friesen, 1978; Caillol et al., 1990). In contrast, prolactin concentrations are high for the duration of lactation in cats (Felis domesticus) (Banks et al., 1983) and blue foxes (Mondain-Monval et al., 1985). This species-specific difference has been attributed to differences in the suckling patterns; rabbits and hares suckle their young only once a day in contrast to the frequent suckling observed in cats (Caillol et al., 1990). Similarly, although spotted hyena cubs depend on milk for 4-6 months and are not weaned until 14-18 months (East et al., 1989), the lactating females are often forced to move large distances in search of food and, consequently, mothers may be away from their cubs for up to 5 days (Kruuk, 1972; East et al., 1989). This results in an irregular and sometimes infrequent suckling pattern in this species. These reduced prolactin concentrations observed during lactation could explain previous reports concerning the lack of a distinct lactational anoestrus in this polyoestrous aseasonal breeder (Lindeque, 1981; Lindeque & Skinner, 1982; Lindeque et al., 1986). Histological and endocrine data suggest that follicular development can be resumed during lactation or shortly thereafter in this species (van Jaarsveld et al., 1992) and provides a mechanism whereby spotted hyaenas, which usually produce only two pre-
social young in each litter (van Jaarsveld et al., 1988) and have an extended lactation period (East et al., 1989), can significantly increase population recruitment rates. In support of this view, negative correlations between prolactin and progesterone in lactating females, and between prolactin and oestradiol in this species, suggest that prolactin secretion declines when ovarian activity increases (van Jaarsveld, 1990). This may also indicate feedback regulation of stress-induced prolactin by progesterone, as recorded in rats (Deis et al., 1989).

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