Testosterone related to age and life-history stages in male baboons and geladas

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A R T I C L E  I N F O

Article history:
Received 26 April 2009
Revised 11 August 2009
Accepted 12 August 2009
Available online 25 August 2009

Keywords:
Androgen
Fecal steroid
Hormone
Life history
Maturation
Method validation

A B S T R A C T

Despite significant advances in our knowledge of how testosterone mediates life-history trade-offs, this research has primarily focused on seasonal taxa. We know comparatively little about the relationship between testosterone and life-history stages for non-seasonally breeding species. Here we examine testosterone profiles across the life span of males from three non-seasonally breeding primates: yellow baboons (Papio cynocephalus or P. hamadryas cynocephalus), chacma baboons (Papio ursinus or P. h. ursinus), and geladas (Theropithecus gelada). First, we predict that testosterone profiles will track the reproductive profiles of each taxon across their respective breeding years. Second, we evaluate age-related changes in testosterone to determine whether several life-history transitions are associated with these changes. Subjects include males (>2.5 years) from wild populations of each taxon from whom we had fecal samples for hormone determination. Although testosterone profiles across taxa were broadly similar, considerable variability was found in the timing of two major changes: (1) the attainment of adult levels of testosterone and (2) the decline in testosterone after the period of maximum production. Attainment of adult testosterone levels was delayed by 1 year in chacmas compared with yellows and geladas. With respect to the decline in testosterone, geladas and chacmas exhibited a significant drop after 3 years of maximum production, while yellows declined so gradually that no significant annual drop was ever detected. For both yellows and chacmas, increases in testosterone production preceded elevations in social dominance rank. We discuss these differences in the context of ecological and behavioral differences exhibited by these taxa.

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Introduction

The steroid hormone testosterone (T) is known to affect many vertebrate life-history traits and has been implicated as a mediator of life-history trade-offs (Hau, 2007; Ricklefs and Wikelski, 2002; reviewed in Zera and Harshman, 2001). For example, the increase in production of T when males reach puberty and begin to seek out mating opportunities is at the same time associated with costs, such as reduced immune function (McGlothlin et al., 2007).

One model for T-behavior trade-offs, known as the “challenge hypothesis”, proposes that variation in male T across life-history stages reflects differential allocation to mating and parenting behavior (Wingfield et al., 1990). Specifically, the challenge hypothesis predicts that high T facilitates inter-male competition at times in the life cycle when males need to compete for receptive females or the resources necessary to attract such females. However, because high levels of T may interfere with paternal behavior (e.g., Goymann et al., 2007; Gray et al., 2006; Muller et al., 2009; Nunes et al., 2000, 2001), T levels should decrease when males care for offspring. Formulation of the challenge hypothesis was based on monogamous, seasonal birds with a high degree of paternal care and cycles of mating and care within each year. For such seasonal species with relatively short and intense cycles of mating, the period of interest for investigating T-mediated trade-offs is within each breeding season (temperate birds: e.g., McGlothlin et al., 2007; Wingfield et al., 1990; tropical birds: e.g., Hau et al., 2008; reptiles: e.g., Wack et al., 2008; and mammals: e.g., Brockman et al., 2001; Cavigelli and Pereira, 2000; Malo et al., 2009; Moss et al., 2001; Ostner et al., 2002, 2008).

More recently, the hypothesis has been modified to apply to non-seasonally breeding species (Archer, 2006; Muller and Wrangham, 2004). For non-seasonal species, T changes across the year are less
informative than T changes across the life span (e.g., Bribiescas, 2006; Crawford et al., 1997; Ellison et al., 2002; Martin et al., 1977). Therefore, with respect to T-mediated trade-offs for non-seasonal species, T should be up-regulated at the start of reproductive maturity, maintained throughout the breeding years, and down-regulated once males no longer breed or when they focus on paternal behaviors. For non-seasonal species, many of the potential trade-offs extend across life-history stages that can take years for long-lived organisms. With the exception of several studies of human and captive non-human primates (e.g., Bribiescas, 2001, 2005, 2006; Crawford et al., 1997; Ellison et al., 2002; Martin et al., 1977), investigation of T profiles across the entire life span is rare (but see chimpanzees (Pan troglodytes): Seraphin et al., 2008; and mandrills (Mandrillus sphinx): Setchell and Dixon, 2002).

Here we examine T profiles across the life span of males in wild populations of three long-lived, non-seasonally breeding primate taxa (see Methods section): yellow baboons (Papio cynocephalus or P. hamadryas cynocephalus), chacma baboons (Papio ursinus or P. h. ursinus), and geladas (Theropithecus gelada). Specifically, we evaluate age-related changes in T and examine whether several life-history transitions (or “maturational milestones”) are associated with these changes. Despite the rarity of mammalian paternal care, baboon and gelada males are known to invest in some degree of paternal care (Beckner and Bergman, 2008; Buchan et al., 2003; Dunbar, 1984; Moscovice et al., 2009; Palombit et al., 2000). Therefore, we expect that trade-offs between the high levels of T optimal for mating and the low levels of T optimal for parenting will result in T modulation for these three taxa, and that this modulation will reflect differences among them in their respective life histories.

We have three lines of inquiry. First, as a physiological validation and in accordance with T profiles from other vertebrate species, we test the prediction that juvenile males have significantly lower fecal T metabolites than adult males. Further, based on profiles of T across the human male life span, T for all three taxa should exhibit an inverse U-shaped pattern, exhibiting a rise at or around maturity and a decline as the animals senesce.

Second, we make the general prediction that the T profiles for males will follow the reproductive profiles of each taxon across their respective breeding periods. In particular, we expect T to remain elevated during ages when males are reproductively active and to return to pre-reproductive levels when mating activity declines and parental care increases. Although all three taxa are non-seasonal breeders, they differ in the timing of reproductive and paternal behavior. At one extreme, geladas have a unimale system (polygynous), in which one male (“leader male”) has sole reproductive access to females that make up a one-male unit. Gelada males have (1) a single tenure as a leader, (2) no reproductive access to females before obtaining a unit (when they live as “bachelor males” in all-male groups), and (3) no reproductive access to females after losing their unit (when they live as “follower males” in their former unit). For gelada males, changes in mating activity across the life span are qualitative and reproductive tenure is discrete. Furthermore, once leader males relinquish their unit and become follower males, then and only then do they engage in protective parenting behavior (Dunbar, 1984)—presumably to protect their offspring from infanticide (Beckner and Bergman, 2008).

In contrast, yellow and chacma baboons exhibit a multimale social structure with a mating system that is polygynandrous to varying degrees. Male dominance rank mediates temporary reproductive access to fertile females. Specifically, dominance rank functions as a maturational marker, (1) enlargement of testes, is available for only one taxon (yellow baboons) and has been shown to precede significant increases in fecal T metabolites (Gesquiere et al., 2005). Therefore, we examine the relationship in these three taxa between T profiles and four additional maturational markers: (2) timing of natal dispersal, (3) acquisition of adult dominance rank, (4) first sexual consortship, and (5) acquisition of highest rank.

Methods

Because the taxonomic level of the different Papio groups remains uncertain (Jolly, 1993), we avoid this debate altogether by referring to yellow and chacma baboons throughout as different “taxa” (see also Barrett and Henzi, 2008)—whether they are considered different species or subspecies does not affect the results presented here. Subjects for this study include all males aged 2.5 years and older from each study population from whom we had fecal samples for hormone determination.

Yellow baboon data collection

The data for yellow baboons come from multiple groups in the Amboseli Basin, Kenya. Because individual life-history data for members of these study groups cover more than three decades (Alberts et al., 2003, 1996; Alberts and Altmann, 1995a; Altmann and Alberts, 2003; Altmann et al., 1988; Pereira, 1988; Shopland, 1987), birthdates are known (within a few days) for all immature and many mature males. Ages of immigrant males for whom birthdates are not known were estimated using an established protocol based on body size and other age-related physical characteristics when these males first appear in one of the study groups. Timing of male maturational milestones for yellow baboons used in this study is taken from previous studies on the population (see summary in Charpentier et al., 2008). Male dominance ranks were determined by assigning wins and losses for all dyadic agonist encounters between males, as described in Hausfater (1975) and Alberts et al. (2003).
As part of the continuing Amboseli baboon research, repeated fecal samples are collected opportunistically from all group members. Because the testosterone RIA kit previously used in our laboratory (Equate 125I Testosterone RIA kit; SolidPhase, Portland, ME) was discontinued, we validated a subset of our samples using a new T RIA kit (Diagnostics Systems Laboratories; Beckman Coulter, Webster, TX). For this validation, we used 2570 samples from 125 different males, for an average of 21 fecal samples per male, with at least 20 different males per age category (see age categories below). All data collection procedures adhered to the Institutional Animal Care and Use Committee guidelines of Princeton University and the laws of Kenya.

Fecal sample collection, storage, and extraction were performed as described previously (Beehner et al., 2006b; Gesquiere et al., 2005, 2007; Khan et al., 2002; Lynch et al., 2003). In brief, freshly deposited samples were mixed thoroughly, placed in 95% ethanol, and stored in a charcoal refrigerator (−20–25 °C) until shipped to the University of Nairobi (every 2 weeks), where the ethanol was evaporated and the samples were freeze-dried. Following freeze-drying, samples were stored at −20 °C until shipped to Princeton University. After transport, each fecal sample was sifted to remove vegetative matter, and 0.2 g of fecal powder was extracted into 2 ml 90% methanol using a multipulse vortexer for 30 min. Following extraction, samples were run through a prepped Oasis cartridge (Waters, Milford, MA) for further purification. Prior to assay, all samples were stored at −20 °C.

Chacma baboon data collection

The data for the chacma baboons come from one wild-feeding group in the Moremi Game Reserve, Botswana. This group has been studied almost continuously since 1982 (Bugler and Hamilton, 1987; Cheney et al., 2004), and the ages of all natal males are known. The ages of immigrant males were estimated based on body size and tooth wear (Kitchen et al., 2003). If newly immigrated males appeared young and in their prime, they were assigned the median age at dispersal for this population (9.25 years, N = 26) at the time they entered the study group. Only natal emigrants who were later seen in a neighboring group were used to calculate median age at first dispersal. Dominance ranks of all males were calculated monthly based on the outcomes of dyadic interactions. Males were assigned an adult rank after achieving dominance over another adult male. First consortships were recorded after a male exhibited his first mate-guarding episode (see also Alberts and Altmann, 1995a).

As part of a 2-year study from 2001 to 2003, repeated fecal samples were collected from all adult males. Additionally, as part of a short-term study to target all age groups, repeated fecal samples were collected from males of all ages during August 2007. As for samples from the yellow baboons, chacma fecal samples had previously been analyzed with the Equate T RIA kit. Therefore, we validated chacma fecal samples with the new DSL T RIA kit as well. For the validation, we used 726 samples from 41 different males, for an average of 18 fecal samples per male. All data collection procedures adhered to the Institutional Animal Care and Use Committee guidelines of the University of Pennsylvania and the University of Michigan and the laws of Botswana.

Hormones were extracted from feces in the field using the method described by Beenhker and Whitten (2004). Specifically, fresh fecal samples were mixed thoroughly, an aliquot of the sample (~0.5 g) was placed in 10 ml of a methanol/aceton solution (4:1), and the solution was immediately homogenized for 1 min using a battery-powered vortexer (BioVortexer; BioSpec Products, Inc., Bartlesville, OK). The dry weight of all fecal samples was later determined (±0.001 g) using a battery-powered, portable scale (Ohaus Scout Pro, Pine Brook, NJ). Approximately 7 h later, 4.0 ml of the fecal homogenate was filtered through a 0.2-µm polytetrafluoroethylene (PTFE) syringeless filter (Whatman, Florham Park, NJ), and the filter was subsequently washed with 1 ml of methanol/aceton (4:1). We then added 7 ml of distilled water to the filtered homogenate, mixed the solution (by inverting it 10 times), and loaded it onto a prepped, solid-phase extraction cartridge (Sep-Pak Plus; Waters) followed by 2 ml of a sodium azide solution (0.1%) as a wash and preservative. All samples were stored dry on cartridges in separate sealed bags with silica beads (~2 g) at subzero temperatures (~ −10 °C) until transported to the University of Michigan for analysis. In the laboratory, steroids were eluted from cartridges with 2.5 ml 100% methanol and subsequently stored at −20 °C until the time of RIA.

Gelada data collection

The data for the geladas come from two bands of wild-feeding geladas in the Simien Mountains National Park, Ethiopia. Because daily observations on this gelada population began in January 2006, all ages of gelada males are necessarily estimated. We placed males in age categories based on (1) a combination of published descriptions of gelada age characteristics based on morphological traits (Dunbar and Dunbar, 1975) and (2) our observations of physical size and developmental markers (e.g., canine eruption) as compared to baboons (Papio) of known ages. Age categories were assigned independently by two observers, and both sets were in close agreement (age categories, estimated ages, and general characteristics describing each category can be found in Supplemental material). No overt dominance hierarchy among gelada leader males has been reported (Mori, 1979). Juvenile males may form temporary hierarchies with their peers and these hierarchies might later extend to all-male groups (Dunbar, 1984); however, no dominance data are available for either of these groups. Timing of first sexual consortships for gelada males was recorded as the first time a new leader male mated with one of the females in his unit.

Samples available for this study were derived from the first 6 months of collection. In total, we collected 328 fecal samples for hormone analysis from 100 different males, for an average of 3 samples per male (range: 1–10), with at least 10 individuals in each age category. Hormones were extracted from feces in the field using a method almost identical to that for the chacma baboons described above (with a few modifications to the volume of sample and solutions used). In brief, fresh fecal samples were thoroughly mixed, an aliquot of the sample (~0.1 g) was placed in 3 ml of a methanol/aceton solution (4:1), and the solution was immediately homogenized. The dry weight of all fecal samples was later determined (~±0.001 g). Approximately 7 h later, 2.5 ml of fecal homogenate was filtered through a 0.2-µm PTFE filter and washed with an additional 1 ml of methanol/aceton (4:1). We then added 7 ml of distilled water to the filtered homogenate, mixed the solution, loaded it onto a prepped Sep-Pak cartridge, and washed the cartridge with 1 ml of a sodium azide solution (0.1%). All samples were stored dry on cartridges in separate sealed bags with ~2 g silica beads. Samples were stored at ambient temperatures for up to 10 days until they could be transported to a freezer at a nearby lodge. Once frozen, samples remained at subzero temperatures (~ −10 °C) until transported to the University of Michigan for analysis. In the laboratory, steroids were eluted from cartridges with 2.5 ml 100% methanol and subsequently stored at −20 °C until the time of RIA.

Testosterone RIA

For all three taxa, we used the same assay methods and the same T primary antibody. We assayed all samples for T metabolites using a modified protocol from a commercially available T RIA kit (Diagnostics Systems Laboratories; Beckman Coulter). All assays used the standards, the primary antibody, the labeled testosterone, and the precipitant solution provided by the kit. Working buffer was a charcoal adsorbed human serum similar to the buffer in which the standards were diluted (American Biological Technologies, Inc., Seguin, TX). The primary antibody from the DSL T kit cross-reacts 100% with testosterone, 6.6% with 5α-dihydrotestosterone, 2.2% with 5-androstan-3β, 17β-diol,
1.8% with 11-oxotestosterone, 0.9% with androstenedione, and 0.6% with 5α-dihydrotestosterone. Cross reactivity of the antiserum with all other steroids is less than 0.5%. All samples were run in duplicate, and the results are expressed as ng/g dry fecal matter.

Method validation

First, to evaluate the effectiveness of extracting T metabolites from primate feces, we determined recovery for each primate taxa with each method. For the yellow baboons, 10,000 cpm of \(^{125}\)I testosterone was added to 0.2 g of dry feces (Lynch et al., 2003). After incubation for 1 h at room temperature, we proceeded with methanol extraction and solid-phase extraction as described above. For the chacma baboons and geladas, we thoroughly mixed a large mass of feces in a plastic cup and measured out 10 aliquots of 0.5 ml wet feces. We then added 18,000 cpm \(^{125}\)I testosterone tracer to each aliquot. After incubating aliquots for 1 h at room temperature, we placed aliquots in 3 ml MeOH/acetone solution (4:1) and extracted each using the same methods described for each taxon above. Following elution, we measured recovery of the radioactivity with a gamma counter. Recovery results for each taxa are listed in Table 1.

Second, we validated the DSL antibody for each taxon. We ran serial dilutions of baboon and gelada fecal extract pools to check for parallelism with the standard curve. We also determined mean assay accuracy (observed/expected × 100) for each taxon by spiking each standard with an aliquot from the respective fecal pool and running them as samples. We calculated intra- and inter-assay coefficients of variation for the assay for all three taxa. Results for parallelism, accuracy, and precision for each taxon are listed in Table 1.

Third, as a physiological validation to our methods, we compared T concentrations between juvenile and adult males for all three taxa. Although we did not expect age-related changes to be identical for all three taxa, we did expect a consistent difference in which juvenile males of each taxon would exhibit lower concentrations of T metabolites than the fully adult males.

Age categories

Generally, primate juveniles are defined by maturational markers such as independence from their mother (marking the start of juvenility) and puberty (marking the end). Because these maturational markers vary for each taxon in this study, we broadly define “juveniles” as males between the ages of 2.5 years (when males of all three taxa are independent from their mother) and 4.5 years (when none of the three taxa have reached puberty). We define adults using similar criteria; “adults” in this study comprise males older than 8.0 years (when all three taxa have reached full adulthood). Males between juveniles and adults are in various stages approaching adulthood, and these differences are taxon specific. Thus, for convenience herein, we broadly use the term “subadult” for all males aged between juveniles and adults (i.e., ages 4.5–8.0 years). We recognize that the subadult biological category is different for each of these taxa.

The ages of most yellow and chacma males were known to within a few days (for males born into study groups) and estimated based on physical characteristics and date of immigration (for immigrating males). By contrast, the ages of all gelada males were estimated based on developmental stages distinguished by physical markers (see Supplementary material). Therefore, to facilitate the comparison of profiles across all three taxa, we placed yellow and chacma males into multi-year age categories corresponding to those for gelada males. We then calculated a mean T value for each category (with each individual contributing only one value). Testosterone profiles thus comprise a combination of longitudinal and cross-sectional data. After comparing T profiles based on the multi-year categories for all three taxa, we then compared T profiles using single-year categories for yellow and chacma males (single-year categories were not available for geladas).

Maturational milestones

Male maturational milestones examined in this study include (1) age at testicular enlargement (signaling puberty and the onset of subadulthood), (2) age at natal dispersal (when males leave their natal group and seek entry in another group), (3) age at attainment of adult dominance rank (signaling the beginning of adulthood), (4) age at first sexual consortship (the best measure available for age at first reproduction in male primates), and (5) age at attainment of highest dominance rank (presumably the time of maximum mating opportunities). Milestones are based on individual life-history data for all except acquisition of highest rank, which, of necessity, is based on cross-sectional data. We report median values for all five of these milestones for yellow males (Alberts and Altmann, 1995a,b; Charpentier et al., 2008; current study) and all but age of testicular enlargement for...

### Table 1

<table>
<thead>
<tr>
<th>Recovery (^{125})I Testosterone (%)</th>
<th>Yellow baboons</th>
<th>N</th>
<th>Chacma baboons</th>
<th>N</th>
<th>Geladas</th>
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<th>All assays</th>
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<td>76.8*</td>
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<td>Fecal pool (~ 70 pg)</td>
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<td>12</td>
<td>4.76</td>
<td>10</td>
<td>4.29</td>
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\* Lynch et al. 2003.
chacma males (Table 2). For geladas, we report age ranges for dispersal and first sexual consortship. Age at testicular enlargement is not known for either gelada or chacma males, and dominance ranks are not available for gelada males because, to our current knowledge, gelada males do not have a formalized dominance hierarchy.

Data analysis

As expected, none of the hormone values for the three taxa were normally distributed. Therefore, we log-transformed values prior to all analyses to facilitate the use of parametric statistics. Additionally, to avoid bias from uneven fecal sample distribution across individuals, we calculated a mean hormone value for each male for each age group to avoid bias from uneven fecal sample distribution across individuals, all analyses to facilitate the use of parametric statistics. Additionally, T metabolites in all three taxa exhibited an inverse U-shaped pattern across age categories (Figs. 1a–c). First dispersal occurred at 6.0 years for yellows and 6.5 years into a single age category for statistical testing. Tukey’s post hoc tests: *p < 0.05, **p < 0.01, ***p < 0.001. The number of males in each age category is indicated above the y-axis.

Three-taxa testosterone comparison

Consistent with the physiological validation, T trajectories were broadly similar across age for all three taxa. Nonetheless, two notable differences were observed. First, yellow and gelada males attained adult T levels at an earlier age than chacma males did (Figs. 1a–c). Testosterone Z-scores for both yellow and gelada males changed from negative to positive between 6.5 and 8.0 years of age, whereas those for chacma males did not do so until 8.0–9.5 years. The ANOVA analysis of log T across age categories further supports this observation (ANOVA: yellows F(7,230) = 4.85, p < 0.001; geladas F(7,100) = 11.13, p < 0.01; chacmas F(6,50) = 2.98, p < 0.05; for post hoc tests between successive age categories, see Figs. 1a–c). None of the juvenile chacma age categories by themselves was significantly different from males had intermediate levels (Z-scores near zero, with the exception of yellows, see below).

Table 2
Median age (or age range, in years) that males reach hormonal and life-history milestones for yellow baboons, chacma baboons, and geladas.

<table>
<thead>
<tr>
<th></th>
<th>Yellow</th>
<th>N</th>
<th>Geladas</th>
<th>N</th>
<th>Chacmas</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hormones (testosterone production)</strong></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Onset of adult T levels</td>
<td>6.5–7.5</td>
<td>6.5–8.0</td>
<td>7.5–8.5</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Peak T levels</td>
<td>7.5–8.5</td>
<td>6.5–8.0</td>
<td>8.5–9.5</td>
<td></td>
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</tr>
<tr>
<td><strong>Life history (maturational milestone)</strong></td>
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<td></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Testicular enlargement</td>
<td>5.38°</td>
<td>96</td>
<td>4.5–6.5°</td>
<td>70</td>
<td>9.252°</td>
<td>26</td>
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<tr>
<td>Acquisition of adult rank</td>
<td>7.38°</td>
<td>48</td>
<td>N/A</td>
<td>8.67°</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>First sexual consortship</td>
<td>7.87°</td>
<td>31</td>
<td>6.5–8.0°</td>
<td>13</td>
<td>8.44°</td>
<td>6</td>
</tr>
</tbody>
</table>

* Source: Charpentier et al. 2008, known ages.
* Source: Dunbar and Dunbar, 1975 (p. 59), estimated ages.
* Based on cross-sectional data (see Fig. 1 for sample sizes for each age category).
N/A, not applicable.
* Source: Current data set, estimated ages.
* Source: Current data set, known ages.

Results

Physiological validation

Juvenile males (2.5–4.5 years old) had significantly lower T metabolites than adult males (>8.0 years old) for all three taxa (t-test: yellows, t(111) = −4.20, p < 0.001; chacmas, t(39) = −2.17, p < 0.05; geladas, t(68) = −5.90, p < 0.001). Additionally, T metabolites in all three taxa exhibited an inverse U-shaped pattern across age categories (Figs. 1a–c). In general, juveniles and subadults had the lowest T metabolites (negative Z-scores), adult males had the highest levels of T metabolites (positive Z-scores), and the oldest of the adult

Fig. 1. Testosterone Z-scores for the different age categories of (a) yellow males, (b) gelada males, and (c) chacma males. Although the figure depicts testosterone Z-scores, all statistical analyses were conducted on the log-transformed testosterone values. Due to the small sample size of juveniles for chacma age categories, we pooled juveniles and subadults aged 4.5–6.5 years into one category (chacmas only).
the 8.0- to 9.5-year age category; however, this was likely due to insufficient power with this data set (i.e. sample sizes were low for chacma males < 6.5 years). When we pooled chacma juvenile males and males from the 4.5- to 6.5-year category, we detected a significant difference from the 8.0- to 9.5-year age category for chacma baboons.

Second, the duration of elevated T profiles was shorter for gelada and chacma males than for yellow males. Testosterone Z-scores for gelada males dropped quickly after 9.5 years (to nearly zero) and scores for chacma males were negative after 11.0 years (Figs. 1b and c). In contrast, Z-scores for yellow males, while dropping slightly after 9.5 years, remain positive even in males 13 years and older (Fig. 1a). Once again, post hoc tests from the ANOVA support these observations (for post hoc tests between successive age categories, see Figs. 1a–c). Gelada males exhibited significantly lower T levels by 9.5 years, and although chacma males did not exhibit significantly lower T levels until 13.0 years, Z-scores were negative by 11.0 years. By contrast, yellow males maintained high T metabolite levels and did not exhibit a significant drop between any pair of successive adult age categories.

Peak T levels for all three species coincided most closely with age at first sexual consortship (although resolution in geladas was coarse due to our estimated age categories). For both yellow and chacma males, acquisition of adult rank came after the onset of adult T levels and before the onset of peak T levels. Age of highest average rank followed peak T levels by 1 year.

Yearly testosterone comparison: yellows and chacmas

Yellow and chacma males followed a similar pattern of maturational, except that chacma males were approximately 1 year behind yellow males for both hormonal and maturational markers (Figs. 2a and b). Males of both taxa (1) exhibited a significant rise in T (Z-scores changing from negative to positive), (2) followed 1 year later by lifetime maximum T, which coincided with attainment of adult rank, (3) followed approximately 1 year later by a decrease in T levels (by at least −0.5 SD), which coincided with males’ highest dominance rank. The delayed rise in T for chacma males coincided with delays in all maturational milestones (dispersal, adult rank, and first sexual consortship) as compared to yellow males (or, where relevant, gelada males; Table 2). Chacma males also exhibited a drop in T levels after 11 years (as indicated by negative Z-scores), while yellow males continued to exhibit higher than average T levels until 17 years of age (as indicated by positive Z-scores).

Discussion

Despite broad similarity in testosterone profiles across age groups of all three taxa, we observed variability across taxa in the timing of two major testosterone transitions. One of these transitions was the attainment of adult levels of testosterone. While yellow and gelada males exhibited maximum testosterone between 6.5 and 8.0 years of age, chacma males did not reach maximum testosterone until 8.0–9.5 years of age. Yearly testosterone profiles indicated that the chacma delay in testosterone production was 1 year later than that for yellows. The other major transition was the drop in testosterone that males exhibited after an approximately 3-year period of maximal production. This drop was significant for geladas and chacmas but not for yellows. Although the testosterone of yellow males decreased gradually, levels did not fall to a consistently lower level until males reached 18 years of age. Consequently, for geladas and chacmas, the period of maximal testosterone production was discrete (i.e., higher than all other age categories by at least one standard deviation) while for yellow males, testosterone production gradually tapered off as males senesced. To understand these differences, we examined testosterone profiles for each taxon in relation to several maturational milestones.

We recall at this point that our main objective in this study was to describe taxon-level patterns in testosterone profiles across the male life span. We do not yet have available longitudinal, individual-based data across ages to partition variability into within and between taxa sources of variance, nor can we yet evaluate hormonal responses to specific social situations for the males involved. Rather, as a first step toward comparative endocrinology for wild populations, we take a broad look at overall hormone patterns and how male milestones map onto these profiles. Specific investigations into the relationship between testosterone changes and male developmental markers for individual males are important topics for the future. As such, the present analysis offers a framework for formulating and testing specific hypotheses for such subsequent studies.

Testosterone and maturational milestones

First, the three taxa exhibited considerable variability in the relationship between testosterone and natal dispersal. At one extreme, geladas dispersed well before the attainment of adult testosterone levels. Gelada “dispersal” from natal one-male units, however, may be qualitatively different from dispersal from natal groups in baboons because the first of these dispersal events in geladas occurs during the subadult (and even juvenile) stages (Dunbar and Dunbar, 1975). Juvenile and subadult gelada males repeatedly come and go from all-male bachelor groups, returning to their natal one-male unit each time, making it difficult to establish a final natal dispersal event. At the other extreme, chacmas dispersed about a year after attainment of adult testosterone levels. Natal dispersal for chacma males is delayed not only with respect to
testosterone but also relative to all male milestones for the other two taxa. A model of the ecological effects on dispersal (Alberts and Altmann, 1995a) considers the trade-offs that dispersing males of all ages face between opportunities for mating (which increase with population density) and possibility of mortality (which increases with predation rate). However, the model does not consider the temporal aspect of natal dispersal within the life-history trajectory of a taxon. Indeed, for both the yellow and chacma populations studied here, many males remain to breed for some time in their natal group (Alberts and Altmann, 1995a; Bulger and Hamilton, 1988), possibly because (as per the model) Amboseli yellow baboons live at relatively low densities (Altmann and Alberts, 2003), and Moremi chacma baboons experience relatively high predation (Cheney et al., 2004). Perhaps as a consequence of multiple ecological factors impinging on dispersal timing, we found no clear relationship between the attainment of adult levels of testosterone and dispersal for these taxa.

Second, the age at first sexual consortship was approximately the same time as that of peak testosterone production. However, under the predictions of the challenge hypothesis, sexual activity alone should not stimulate an exponential increase in testosterone production (Wingfield et al., 1990) but rather sexual activity in concert with male–male aggression, that, for baboons, occurs in the context of rank attainment within a dominance hierarchy. Because the age of first sexual consortship and the acquisition of adult rank were temporally similar for yellows and chacmas (<5 months apart for yellows and <3 months apart for chacmas), the present analysis is not fine tuned enough to sufficiently relate peak testosterone levels to either of these milestones.

Third, yellow and chacma males attained their highest dominance rank 1 year after peak testosterone production. This supports an earlier finding that baseline testosterone levels predict future dominance ranks, but rank changes in themselves do not affect testosterone levels (Beehner et al., 2006a). Our results support the challenge hypothesis in showing a close link between testosterone and a time period when we expect male–male contests, but the causality of this relationship (if any) is opposite the prediction. The challenge hypothesis was proposed as a feedback loop between the external social environment and the internal physiological one, with a social “challenge” initiating the cycle. However, for both yellow and chacma males, testosterone declines before males fall in rank—sometimes even up to 6 months beforehand (Beehner et al., 2006a). Unless there is an anticipatory decline in testosterone, our data indicate that testosterone production is not necessarily “socially modulated” (Wingfield et al., 1990)—or, at least not at the broad scale that we use in this study. Individual contests may indeed affect daily fluctuations in testosterone levels (i.e., winner-loser effects; Mazur and Booth, 1998; Mazur et al., 1992). However, we suggest that the more stable baseline testosterone levels that characterize life-history stages do not result from rank changes but, in fact, precede them.

Testosterone differences across taxa

Why do chacma males delay testosterone production, rank acquisition, and dispersal by at least a full year relative to yellow males? One explanation may relate to the high density of baboons in the Moremi chacma population (24 baboons/km²; Cheney, 1987; Cheney et al., 2004; Hamilton and Bulger, 1992; Hamilton et al., 1976). Males attempting to disperse to neighboring groups may face high resistance from the males already established in these groups. Males might overcome this resistance by achieving full adult body size prior to emigration since increased body size upon immigration to a new group could facilitate a more rapid ascent in the dominance hierarchy. Additionally, if males rise in dominance within their natal group, they may gain more “practice” at rank contests, whether this involves actual fighting or displaying (Kitchen et al., 2003). High predation may also have delayed Moremi chacma males’ life-history variables since dispersing males are certain to experience elevated mortality during transfer. Thus, pressures related to ecological factors in this area may have selected for a reproductive strategy that maximizes “maturational” in one’s natal group prior to dispersal. Chacma males dispersed and attained their highest dominance rank in the same year. This “strategy” is in sharp contrast to yellow males, and possibly even anubis (P. anubis or P. h. anubis) males; both yellow and anubis males disperse around 8 years of age (Charpentier et al., 2008; Packer, 1979; Packer et al., 1995), yet do not attain highest rank until nearly 2 years later for yellows (Table 2) and 3 years later for anubis (Packer et al., 2000).

At the other end of the life-history trajectory, why do yellow males continue to produce testosterone well beyond the period when they are high ranking? Low-ranking male yellow baboons have comparatively greater access to sexually receptive females than male geladas and chacmas (Alberts et al., 2003, 2006). Two components of this weak, dominance-based priority are that younger, non-alpha males are able to gain some fertile matings and that older, non-alpha males are also able to mate more than would be expected under a strict queuing scenario. This prolonged maintenance of higher testosterone levels into older ages for yellow males represents a departure, not only from chacma and gelada males but also from other known mammalian male profiles (Castracane et al., 1986; Crawford et al., 1997; Muehlenbein et al., 2001; Seraphin et al., 2008).

Testosterone and parental care

In many ways, male development for chacma baboons is more similar to that of geladas than it is to that of yellows. Compared to the testosterone profiles of yellow males, those of chacma and gelada males indicate a comparatively discrete increase to adult levels and decrease about 3 years later. This discrete period of elevated testosterone production is what we might expect for a species such as the gelada, with unimale reproductive units. That chacmas also fit this profile suggests that chacma male physiology may be tracking reproductive activity and/or parenting behavior better than the overall mating system. Although yellow and chacma baboons share a multimale, polygynandrous mating system, the prime “mating” versus “parenting” stages for chacmas and geladas may be more compartmentalized than those of yellows. Although older, low-ranking chacma males continue to take advantage of mating opportunities (Crockford et al., 2007), and they achieve some reproductive success (Cheney and Seyfarth, unpublished data) they also invest considerable time and energy in paternal care (Moscovice et al., 2009; Palombit et al., 2000, 1997). In many mammalian species, high levels of testosterone have been shown to be incompatible with paternal behavior (Gray et al., 2006; Muller et al., 2009; Nunes et al., 2000, 2001), and thus the chacma male sharp decline may be related to a shift in an overall mating strategy to a parenting one.

Taxon or population differences?

One factor that has not been addressed in this study is whether the male hormone and life-history patterns we report here are taxon specific or population specific. This raises the question of whether we are documenting intrinsic or extrinsic sources of variation. The overall hormonal profile may exhibit a fixed pattern within a taxon and tell us something about a taxon’s overall adaptive strategy. Alternatively, different populations may demonstrate facultative responses to environmental changes, and hormones (such as testosterone) can facilitate this behavioral flexibility (Oliveira, 2004). Offering some support to the latter hypothesis, captive yellow or yellow-olive hybrid males exhibited adult levels of testosterone much earlier (Altmann et al., 1988; Castracane et al., 1986; Muehlenbein et al., 2001) than the
wild yellow males in the current study. At present, no comparable hormone data are available from other wild populations of baboons or geladas to test these alternatives. Certainly, a replication of this study in other populations with different population dynamics and/or ecology would help resolve this issue (for details on the ecology of these three sites, see Amboseli (Alberts et al., 2005; Behrensmeyer, 2006), Moremi (Ellery et al., 1993), and Simien (Nievergelt et al., 1998))). For example, if the delay in testosterone production and other life-history stages observed in chacma baboons is due to the high population density in Moremi, then we predict a very different pattern for chacma baboons at lower densities (e.g., Drakensberg chacma population, South Africa).

A third possibility (which is obscured using the broad-scale approach we use here) is that testosterone variation represents enduring individual variation and thus reflects alternative life-history strategies. For example, a high parental investment strategy may exhibit a long-term commitment to one mate and parental care, accompanied by low mating effort, while a low parental investment strategy may exhibit a short-term commitment to one mate and no paternal care, accompanied by high mating effort (Gross, 1985, 1996; Thompson and Moore, 1992; van Rhijn, 1974). With respect to this third possibility, future studies that statistically examine the variability in developmental markers (relative to hormonal profiles) between taxa as well as between individuals within each taxa would be extremely useful for understanding the trade-offs between testosterone and life-history traits in wild populations.

Acknowledgments

The yellow baboon research was supported by NSF IBN-0322613, NSF BSE-0323553, RO3 MH65294, and the Chicago Zoological Society. We thank the Office of the President, Republic of Kenya, the Kenya Wildlife Services, its Amboseli staff and Wardens, the Institute of Primate Research, the National Museums of Kenya, and the members of the Amboseli-Longido pastoralist communities. In the field, thanks go to the Amboseli team who contributed to sample and data collection (R.S. Mututua, S. Sayialel, and J.K. Warutere). Thanks to T. Fenn for database assistance and to C. Markham, P. Onyango for their input on data analysis or for providing comments on earlier drafts. This research was approved by the Animal Care and Use Committee at Princeton University (IAUC protocol #1689, 9 November 2007) and adhered to all the laws and guidelines of Kenya (Kenya Research Permit MOEST 13/001/C351 Vol. II).

The chacma baboon research was supported by NIH MH62249, NRSA, the Leakey Foundation, the University of Pennsylvania, and the University of Michigan. We are grateful to the Office of the President of the Republic of Botswana and the Botswana Department of Wildlife and National Parks for permission to conduct this research. In the field, thanks go to T. Bergman, M. Mokupi, and A. Mokupi for sample and data collection, and in the laboratory, thanks go to S. Mohanthy for assay work, troubleshooting, and analysis. This research was reviewed and approved by the Animal Care and Use Committee at the University of Pennsylvania (IAUC protocol #09001) and the University Committee on Use and Care of Animals at the University of Michigan (UCUA protocol #09554) and adhered to all the laws and guidelines of Botswana.

The gelada research was supported by the National Geographic Society (NGS 8100-06), the Leakey Foundation, NSF BCS-0715179, and the University of Michigan. We are grateful to the Ethiopian Wildlife Conservation Department, the Amhara National Regional State Parks Development and Protection Authority, and the Wardens and staff of the Simien Mountains National Park for permission to conduct this research. In the field, thanks go to T. Bergman and H. Gelaye for sample and data collection, and in the laboratory, thanks go to S. Mohanty for troubleshooting and sample analysis. This research was approved by the University Committee on Use and Care of Animals at the University of Michigan (UCUA protocol #09554) and adhered to all the laws and guidelines of Botswana.

Finally, we would like to thank T. Bergman and two anonymous reviewers for their helpful comments on earlier versions of this manuscript.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.yhbeh.2009.08.005.

References


