Hormonal correlates of natal dispersal and rank attainment in wild male baboons

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ABSTRACT

In many mammals, maturational milestones such as dispersal and the attainment of adult dominance rank mark stages in the onset of reproductive activity and depend on a coordinated set of hormonal and socio-behavioral changes. Studies that focus on the link between hormones and maturational milestones are uncommon in wild mammals because of the challenges of obtaining adequate sample sizes of maturing animals and of tracking the movements of dispersing animals. We examined two maturational milestones in wild male baboons—adult dominance rank attainment and natal dispersal—and measured their association with variation in glucocorticoids (GC) and fecal testosterone (T). We found that rank attainment is associated with an increase in GC levels but not T levels: males that have achieved any adult rank have higher GC than males that have not yet attained an adult rank. This indicates that once males have attained an adult rank they experience greater energetic and/or psychosocial demands than they did prior to attaining this milestone, most likely because of the resulting participation in both agonistic and sexual behaviors that accompany rank attainment. In contrast, natal dispersal does not produce sustained increases in either GC or T levels, suggesting that individuals are either well adapted to face the challenges associated with dispersal or that the effects of dispersal on hormone levels are ephemeral for male baboons.

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1. Introduction

In mammals, maturational milestones such as puberty, adult dominance rank attainment and natal dispersal mark important transitions associated with the onset of reproductive activity (Adkins-Regan, 2005; Jack et al., 2014; Pusey, 1987). These transitions critically depend on a set of coordinated hormonal and socio-behavioral changes (Finch and Rose, 1995). Further, they require the mobilization of resources from individuals’ energy stores, and physiological preparation for reproduction (see discussion in Belthoff and Dunlop, 1995, 1998; Nunes et al., 1999; Ronce and Clobert, 2012). Steroid hormones, specifically glucocorticoids (GC) and testosterone (T) are presumed to play key roles in these processes because of their involvement in energy mobilization and reproduction (Dixson, 2012; Nelson, 2011; Sapolsky et al., 2000). Adult dominance rank attainment—the stage at which a maturing male consistently begins to outrank at least one adult male in agonistic encounters—represents an important maturational milestone in many primate species, because individuals (especially males) are often unable to acquire mating opportunities prior to adult rank attainment (Albore et al., 2006; Altmann and Altmann, 1995b; Beehner et al., 2009; Charpentier et al., 2008; Hamilton and Bulger, 1990; Van Noordwijk and Van Schaik, 2001). In some primates, males are unable to reproduce until they attain a high dominance rank, but in other species, the attainment of an adult rank per se is sufficient (reviewed in Altmann et al., 2003; Bulger, 1993; Cowlishaw and Dunbar, 1991).

Dispersal is nearly ubiquitous across the animal kingdom, enhancing gene flow between populations, limiting inbreeding, and allowing individuals to distribute themselves over available resources (Clutton-Brock, 1989; Greenwood, 1980; Johnson and Gaines, 1990; Lawson Handley and Perrin, 2007; Paradis et al., 1998). Dispersal represents a major maturational milestone for many mammals, because individuals often do not reproduce prior to dispersal (Greenwood, 1980; Pusey and Parker, 1987; Pusey, 1987). Many mammals—including baboons (Papio cynocephalus), the subjects of this study—exhibit male-biased dispersal, so that social groups consist of families of female kin.
with immigrant males who tend to be unrelated to other group members other than their own offspring (Greenwood, 1980; Pusey, 1987; Clutton-Brock, 1989).

Adult rank attainment and natal dispersal both represent a potent combination of social, reproductive, and energetic challenges, possibly among the greatest challenges that an animal will experience in its lifetime. For instance, dispersing primates may face increased mortality risk as a result of predation, partly because they may often be alone (Alters and Altmann, 1995a; Cuárd, 1997; Fernández-Duque, 2009; Isbell and Jack, 2009). Dispersing individuals may also experience nutritional stress resulting from lack of familiarity with existing resources in new areas (Pusey and Packer, 1987). Some dispersing individuals travel long distances before they encounter a new group, potentially depleting energy stores during dispersal (see discussions in Dufy and Bethloff, 2001). Successful dispersal and rank attainment may also involve fighting with conspecifics, resulting in injuries or occasionally death (see discussion in Isbell and Jack, 2009; Van Noordwijk and Van Schaik, 2001). In the aggregate, the challenges and changes associated with these maternal events are likely to be reflected in hormonal changes.

Hormones and socio-sexual behavior operate in a feedback loop, each responding to and contributing to the regulation of the other (Bercovitch and Ziegler, 2002; Nelson, 2011). For instance, a stressful event—an event that has the potential to perturb physiological homeostasis (Sapolsky, 2004)—leads to activation of the hypothalamic-pituitary-adrenal axis and the adrenomedullary system, which in turn leads to downstream secretion of catecholamines (epinephrine and norepinephrine) and glucocorticoids (GC) (e.g., cortisol and corticosterone) (Elenkov and Chrousos, 2002; Sapolsky et al., 2000). Secretion of these hormones in turn stimulates or suppresses various cardiovascular, metabolic, and immune system processes (e.g., heart rate, blood pressure, glucocorticosis, lipopolysaccharide, inflammation), all of which may have immediate or downstream consequences for behavior. A short-term release of GC during acutely stressful events is adaptive and aids in survival. On the other hand, chronic release of GC is thought to be immunosuppressive and consequently leads to increased susceptibility to disease and infection (Sapolsky et al., 2000; Sapolsky, 2004; Selva et al., 2011).

Testosterone (T) is produced in the testes and regulated by the hypothalamic-pituitary-gonadal axis (Hirschenthaler and Oliveira, 2006). Testosterone regulates the production of gametes, enhances anabolism, and maintains musculoskeletal performance (Adkins-Regan, 2005; Bardin, 1996; Bribiescas, 2001). Testosterone has previously been associated with behaviors such as territoriality, mate guarding/consortship, aggression, and mating display (Beehner et al., 2006; Oyango et al., 2013a; Wickens and Dixson, 1992; Wingfield et al., 1990). The 'challenge hypothesis,' designed to explain the link between T and competitive behavior in seasonal breeders, predicts that T levels rise in response to social challenges, particularly those associated with competition for mates, and specifically that T levels can be rapidly elevated when critically needed to promote aggressive competition in a reproductive context (Wingfield et al., 1990). Research in seasonally breeding birds has supported this hypothesis by showing that T is low in the absence of mating behavior or mate competition and elevated during aggressive episodes and mating (Wingfield et al., 2000). Studies in both seasonally and non-seasonally breeding nonhuman primate species have also broadly supported the challenge hypothesis (e.g., chacma baboons (Papio hamadryas ursinus): Beehner et al., 2006; howler monkeys (Alouatta palliata): Cristóbal-Azkarate et al., 2006; ring-tailed lemurs (Lemur catta): Gould and Ziegler, 2007; mous-tached tamarin monkeys (Saguinus mystax): Huck et al., 2005; tufted capuchin monkeys (Cebus apella nigrizus): Lynch et al., 2002; bonobos (Pan paniscus): Marshall and Hohmann, 2005; chimpanzees (Pan troglodytes): Muller and Wrangham, 2004; red-fronted lemurs (Eulemur fulvus rufus): Ostner et al., 2002; reviewed in Muller, 2016). Testosterone is energetically costly to maintain at high levels because of its anabolic effects (e.g., muscle mass building). These energetic costs will lead to decreased investment in immune function and increased susceptibility to disease (Muehlenbein and Bribiescas, 2005; Zuk and McKeen, 1996).

Studies that focus on the link between hormones and maturational milestones are uncommon in wild mammals because of the challenges of obtaining adequate sample sizes of maturing animals and of tracking the movements of dispersing animals. Only a few studies have looked at the endocrine correlates of dispersal in mammals (yellow baboons: Alberts et al., 1992; chacma baboons: Beehner et al., 2006; Bergman et al., 2005; spotted hyenas (Crocuta crocuta): Holekamp and Smale, 1998; white-faced capuchin monkeys (Cebus capucinus): Jack et al., 2014; chimpanzees: Kahlenberg et al., 2008; Belding's ground squirrels (Spermophilus beldingi): Nunes et al., 1999; Muriqui monkeys (Brachyteles arachnoidea): Strier and Ziegler, 2000 and crested macaques (Macaca nigra): Marty et al., 2017). Some studies of GC or T in relation to dispersal have compared hormone levels in immigrant males with those in resident males, whereas others have compared hormone levels in the same individuals before and after dispersal, or at different stages after immigration. For example, in spotted hyenas, immigrant males—those that have immigrated within the previous 7 days—have higher circulating T than natal males even after controlling for the effects of age (Holekamp and Smale, 1998). Alberts et al. (1992) found unusually high GC and T levels in a particularly aggressive male baboon immediately after he immigrated into a new group. In chacma baboons, T levels appear to rise in anticipation of future mating opportunities and rank rises (Beehner et al., 2006; Bergman et al., 2005). Fecal GC levels are higher in immigrant male chacma baboons in the month following immigration than they are in subsequent months of residency (Bergman et al., 2005), and the same is qualitatively true in graycheeked mangabeys (Lophocebus albigena) (Arlot et al., 2009). Among chimpanzees, recent immigrants to a community have higher GC levels than natal residents (Kahlenberg et al., 2008). Fecal GC levels are higher in immigrant male crested macaques in the first few days after immigration and return to baseline levels quickly (Marty et al., 2017). However, a longitudinal study in black howler monkeys (Alouatta pigra) showed that immigrant males did not differ in their GC and T levels 2 weeks before and after immigration (Van Belle et al., 2009). In Muriqui monkeys, cortisol levels did not differ between natal enigmatic females and recent immigrant females (Strier and Ziegler, 2000). Notably, only one study to our knowledge has looked at T variation around the time of adult rank attainment, in three primate species (chacma baboons, yellow baboons, and geladas (Theropithecus gelada): Beehner et al., 2009). In that study, the acquisition of adult ranks in males occurs at the onset of adult T levels but before peak T levels (Beehner et al., 2009).

1.1. Maturation and reproduction in male baboons

To provide context and background for the goals of this analysis, we review here the general sequences of maturational events that male baboons experience, and their functional consequences for males' physical and social lives. Among the wild baboons in the Amboseli basin of southern Kenya, the subjects of this study, males experience testicular enlargement (puberty) at a median age of 5.41 years (Fig. S1; Oyango et al., 2013b). Testicular enlargement signals physiological maturity and the ability to produce viable sperm (Bercovitch and Goy, 1990; Plant, 1994). However, unlike females, who conceive their first offspring about 1 year after puberty (menarche), males experience a 2 year period of reproductive quiescence after testicular enlargement— an adolescent phase that is generally termed subadulthood. Adolescence involves a growth spurt that, by the age of 7–8 years, results in male body size approximately double that of adult females (Altmann and Alberts, 2005).

Once males attain this large body size, they are able to agonistically challenge adult males and take a place in the adult male dominance hierarchy; this milestone, the attainment of adult rank, occurs at a median age of 7.45 years in Amboseli and signals the transition from
subadulthood to adulthood (Fig. S1). Natal dispersal, the second milestone of interest in this analysis, often occurs immediately after adult rank attainment, but may occur well before or well after it (Fig. S1). In addition, some males in Amboseli engage in reproductive activity via mate guarding and copulation (generally referred to as consorting in primates) before dispersing, although most do not (Alberts and Altman, 1995a).

In Amboseli, males never mate guard before they attain an adult rank, and usually begin mate guarding very soon afterwards (Alberts and Altman, 1995b; also Fig. S2). As in some other baboon populations, this effect is not dependent upon males attaining a rank in the top three positions (defined as high-ranking) of the adult male dominance hierarchy. Both high and low-ranking male baboons can and do engage in mate-guarding episodes and father offspring, but low-ranking males are not as successful in obtaining mate-guarding opportunities as high-ranking males, on average. In our data set, approximately half of the males attained a high rank in the 6-month period after adult rank attainment, and approximately half did not (Fig. S3; Alberts et al., 2006, 2003, see also Bercovitch, 1988, 1986; Bulger, 1993). Specifically, the extent of reproductive skew is density-dependent. In small groups, high-ranking males have a large advantage; in large groups, high-ranking males have only a small advantage. On average in this population, the highest ranking male attains 20% of available mate guarding opportunities (Alberts et al., 2003).

1.2. Goals, hypotheses, and predictions

In the present study, we examined two maturational milestones in wild male baboons that are associated with the transition from subadult hood to adulthood: adult dominance rank attainment and natal dispersal. We measured the association between the timing of these milestones and variation in fecal glucocorticoids (FGC) and testosterone (TT) (see Gesquiere et al., 2005; Onyango et al., 2013b for similar analyses for age at puberty). Our study focused on the developmental period after puberty and into early adulthood (5–10 years) to eliminate pubescent males and also to eliminate males that were long past maturational process. Specifically, we modeled fecal steroid concentrations as a function of maturational status (which of the two milestones of interest had been attained) and other variables known to affect hormone concentrations. We tested the hypothesis that psychosocial and energetic demands on male baboons will be substantially higher after dispersal and/or rank attainment than before them. Animals may experience increased nutritional stress and other energetic and social demands as they adjust to their new environment after dispersal, and males are likely to experience substantially increased reproductive activity following rank attainment. This hypothesis yielded two predictions that we tested. 1) We predicted that individuals that had attained one or both of these milestones would have higher levels of FGC than those that had not. 2) We predicted that individuals that had attained one or both of these milestones would have higher levels of TT than those that had not as a consequence of greater mating and agonistic activity after milestone attainment.

2. Methods

2.1. Study population

Our study focused on members of a well-studied wild baboon population in the Amboseli region of Kenya. This baboon population is composed of yellow baboons, Papio cynocephalus, that experience some admixture with neighboring populations of olive baboons, Papio anubis (Alberts and Altman, 2001; Charpentier et al., 2012; Tung et al., 2008b). The Amboseli baboon population has been studied for the last four decades; this research has produced longitudinal, individual-based behavioral and demographic data, as well as fecal samples used for the measurement of hormone metabolites. Demographic, behavioral, and life-history data (births, emigration, immigration and maturation events) are collected on a near-daily basis on known individuals in the study population (Alberts and Altman, 2012) and are stored in Base, the Amboseli baboon project database (Plinc et al., 2016). During the study period (January 2000 to November 2013), the number of study groups varied, as groups occasionally fissioned, and some groups were dropped from continuous observations. Number of study groups at any given time varied from 4 to 6 (Table 1).

2.2. Study subjects and maturational milestones

The subjects of this study were all males that were (i) born into Amboseli study groups and known from birth (hence their ages were known to within a few days), and (ii) males for whom hormone data are available for the period before and after they attained adult rank and/or dispersed. The resulting sample includes 114 males born between October 1990 and July 2007, attained adult dominance rank and/or dispersed between January 2000 and November 2013.

As described above, we consider the attainment of adult rank to signal the transition for males from subadulthood—a period of rapid growth but reproductive quiescence after puberty—to adulthood (Alberts and Altman, 1995b; Altman and Alberts, 2005). Adult rank attainment occurs at a median age of 7.45 years (range 6.2 to 8.7 years; Onyango et al., 2013b; Fig. S1). The attainment of a dominance rank among adult males, even if it is not a high rank, is a pre-requisite for obtaining reproductive opportunities in this population (Fig. S2). The other maturational milestone we considered here, natal dispersal, is usually attained after testicular enlargement (maturational milestone of puberty), but 5% of males disperse before testicular enlargement. Natal dispersal occurs at a median age of 6.76 years (age range of post-pubertal natal dispersal is 6.81–13.42 years; Onyango et al., 2013b). Our study focused on the two post-pubertal maturational milestones: the attainment of adult dominance rank and dispersal from the natal group to a non-natal group. Rank attainment and natal dispersal are frequently, but not always, attained within a few weeks or months of each other in our study population, making them highly associated (X^2 = 1809.6, p < 0.001, Cramer's V = 0.781; Fig. S1; Alberts and Altman, 1995b; Onyango et al., 2013b).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Description of predictors for hormone levels.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Predictor variable</td>
<td>Description</td>
</tr>
<tr>
<td>Male ID</td>
<td>Male identity.</td>
</tr>
<tr>
<td>Social group</td>
<td>The social group the individual was in at the time of fecal sample collection. We observed between 4 and 6 social groups at any given time during the course of the study (see Study population, Methods).</td>
</tr>
<tr>
<td>Year</td>
<td>Year of sample collection.</td>
</tr>
<tr>
<td>a) Random effects</td>
<td>Social group</td>
</tr>
<tr>
<td>Age</td>
<td>Age in years at fecal sample collection (continuous). Age at sample collection ranged from 6.0 to 10 years.</td>
</tr>
<tr>
<td>Season</td>
<td>Whether the fecal sample was collected during a wet or dry season (wet, dry); binary.</td>
</tr>
<tr>
<td>Number of adult males</td>
<td>Number of adult males in the individual's social group at the time of fecal sample collection. Number of adult males in a group: 9.7 ± 4.36 (mean ± SD), range 1–21.</td>
</tr>
<tr>
<td>Dominance rank</td>
<td>Ordinal dominance rank of the individual at the time of fecal sample collection. This varied considerably, including as it did both subadults and adult males. Range: 1–29.</td>
</tr>
</tbody>
</table>
Male dominance ranks were determined by assigning wins and losses in dyadic agonistic encounters between males. Males are considered to win agonistic encounters in which their opponent gives only submissive gestures while they give only aggressive or neutral (nonsubmissive) gestures (Alberts et al., 2003; Hausfater, 1975). This procedure of assigning wins and losses allows the construction of a square matrix of interactions in which entries below the diagonal (which would represent wins by the lower-ranking animal) are few or zero; males are assigned ordinal dominance ranks based on their position in this matrix. Ordinal dominance ranks based on these interactions were assigned monthly to each male in each study group, with the highest rank status as 1. Males were assigned an ordinal dominance rank whether or not they had attained a dominance rank among adult males. For instance, in a group with 7 fully adult males, the highest-ranked subadult male (i.e., a male who had never successfully challenged an adult male in an agonistic encounter but won encounters with all other subadults) held an ordinal rank of 8, with lower-ranking subadult males assigned successively higher ordinal rank numbers. In other words, the male's maturation status (whether or not he had attained a rank among adult males) and his ordinal rank (the rank number that he was assigned) were two separate pieces of information about the male's rank, both of which we considered in our analyses.

2.3. Fecal sample collection, hormone extraction, validation and analyses

Fresh fecal samples for hormone analysis were opportunistically collected from individually known males, labeled with the individual male's ID and the date of sample collection, and stored in 95% ethanol. Protocols for extraction and measurement of IGC and FT metabolites are well-established and have been previously validated in research on this population (Beehner et al., 2008; Gesquiere et al., 2005; Khan et al., 2002). These samples were freeze dried, sifted, and weighed; they underwent methanol and solid phase extraction using Waters Oasis HLB cartridges. IGC and FT metabolites in the samples were measured in ng/g of dried feces using radioimmunoassays (RIA). RIA specific to IGC and FT metabolites have previously been validated for our population (Beehner et al., 2009; Gesquiere et al., 2011; Khan et al., 2002; Lynch et al., 2003). For FT assays, we have used different kits over the years, all of which produce highly correlated results. These kits have included the kit from Equate (SolidPhase, Portland, ME; See Lynch et al., 2003 for validation), the kit from Diagnostic Systems Laboratories (Beckman Coulter, Webster TX; Beehner et al., 2009; Gesquiere et al., 2011), and the kit from Pantex (Santa Monica, CA; Gesquiere et al., 2014). To render the FT concentrations directly comparable across kits, a correction factor was applied to the FT from the different kits (details in Gesquiere et al., 2011).

2.4. Data analysis

2.4.1. Mixed model analysis of variation in IGC and FT metabolites associated with adult rank attainment and natal dispersal

2.4.1.1. Response variables and modeling approach. Our data set included measures of hormone levels from 2955 fecal samples collected from the 114 unique subadult and adult males (aged 6–10 years) described above; most of these individuals were represented in our data set by multiple fecal samples (mean number of fecal samples per individual = 26, range 1 to 97). To test our predictions, we ran two general linear mixed models with Gaussian error distribution, one for each hormone. The response variables for the models were log-transformed IGC levels or log-transformed FT levels. Predictor variables are described below and in Table 1. To generate our mixed effect models, we used the 'glmmadmb' and 'lme4' package (Fournier et al., 2012) in the R Statistical software, version 3.2.3 (R Foundation for Statistical Computing, 2014). Visual inspection of the data for all models indicated no violations of the assumptions regarding the normality of residuals and homogeneity of error variances. The variance inflation factors for the predictors were measured using the 'car' package in R statistical software and were all ≤5.7 and Cook's distance measured by the 'influence.ME' package in R did not indicate influential cases.

2.4.1.2. Predictor variables. Our fixed effect of interest was the maturation status of the individual at the time the fecal sample was collected. Individuals were assigned to one of the following categories for maturation status: dispersed (he had dispersed but not attained rank, N = 114 fecal samples, 13 males), ranked (he had attained rank but had not dispersed, N = 209 fecal samples, 35 males), neither (he had neither dispersed nor attained rank, N = 1419 fecal samples, 101 males), and both (he had dispersed and attained rank, N = 1213 fecal samples, 50 males). Our models included three random variables; 1) the identity of the individual, to account for multiple samples from most individuals, 2) year of sample collection, to account for samples collected across multiple years and 3) the social group the individual was in at the time of sample collection, to account for multiple samples from each social group.

Age, season, and the male's ordinal dominance rank were included in all analyses as covariates because they are known to predict IGC and FT hormone levels in male baboons. Specifically, FT levels in Amboseli males are stable prior to puberty between 2 and 5 years of age, increase through early adulthood to a peak at 9.5 years, and then gradually decline with age throughout adulthood (Altmann et al., 2010; Beehner et al., 2009; Onyango et al., 2013b). In addition, IGC levels are higher during the dry season (June–October) than the wet season (November–May), while FT levels show the reverse pattern with higher levels during the wet season (Alberts et al., 2005; Altmann et al., 2002; Gesquiere et al., 2011). Following previous studies, we categorized each month of our study period as a dry month (June through October) or a wet month (November through May). With respect to dominance rank, high-ranking adult males have lower IGC and higher FT levels than lower ranking males, with the exception of the alpha male, who exhibits high levels of both IGC and FT (Gesquiere et al., 2011). We also included the number of adult males in the social group at the time of sample collection as a predictor of hormone levels because it acts as a proxy for male-male competition (a potential stressor). For example, (Marty et al., 2017) have shown that male crested macaques have higher IGC levels when they immigrate into groups with larger numbers of adult males.

2.4.2. Categorical analysis of variation in IGC and FT metabolites associated with adult rank attainment and natal dispersal

Finally, following Bergman et al. (2005), we tested the possibility that hormone levels would be elevated in the month immediately following the attainment of a maturation milestone (adult dominance rank or natal dispersal) but not during later months. We ran independent analyses for each maturation milestone. Specifically, we grouped fecal samples into two classes, those that were collected within one month of maturation milestone attainment, and those that were collected more than one month after attaining either milestone (i.e., between 2 and 6 months post-milestone attainment). Then we identified subjects for whom we had fecal samples in both categories. Restricting the analysis to these males, we calculated the mean hormone concentration for each male in each category. For each milestone, we used the Wilcoxon test (a non-parametric equivalent of paired t-test) to examine whether mean log hormone levels (IGC and FT) differed between the two categories. This differed slightly from the approach taken by Bergman et al. (2005) because in their analysis, post-dispersal IGC levels did not vary across individuals, allowing them to disregard individual identity and perform an ANOVA. In our case, we had significant heterogeneity among individuals in post-milestone hormone levels: for post-dispersal IGC levels (F5.18=3.473, p < 0.0001, range = 1.4–2.2) and FT (F5.18=5.344, p = 0.001, range = 1.5–2.5). Reported ranges for IGC and FT levels are log
transformed values. Rank attainment analysis showed similar trends; post-rank attainment fGC levels ($F_{2, 94} = 1.568, p < 0.09, \text{range} = \{1.0-2.2\}$) and ft levels ($F_{3,14} = 1.943, p < 0.029, \text{range} = \{1.3-2.9\}$). Thus, we opted for the paired Wilcoxon signed rank test to control for these individual differences.

3. Results

3.1. Mixed model analysis of variation in fGC and ft metabolites associated with adult rank attainment and natal dispersal

Fecal glucocorticoid (fGC) levels were significantly higher in males with a status of ‘ranked’ or ‘both’ than in males with a status of ‘neither’. Fecal GC levels were 15% higher in individuals in the ‘ranked’ category compared to neither ($p = 0.005, \text{CI} = 5-23\%$) and 11% higher in individual in the ‘both’ category compared to neither ($p = 0.059, \text{CI} = 2-20\%$). In our fGC model, the random effects (i.e. individual identity, social group and year) accounted for 48%, 22%, and 20% of variance in fGC respectively. In contrast, males with a status of ‘dispersed’ (i.e. dispersed but not ranked) did not have higher fGC levels than males with a status of ‘neither’ (Table 2). Further, the effect estimate for ‘dispersed’ males was negative, indicating that the non-significance of this predictor was unlikely to be simply the result of limited statistical power. In combination these results suggest that the significant estimate for males in the ‘both’ category is driven by males that had attained rank (whether or not they had dispersed), and that rank attainment but not dispersal was associated with elevated fGC. In support of this inference, a visual comparison of fGC levels shows that individuals that had attained rank (i.e., those in the ‘both’ category plus those in the ‘ranked’ category) had higher fGC levels than those that had not (i.e., those in the ‘neither’ category plus those in the ‘dispersed’ category; Fig. 1A). However, a parallel visual comparison for the dispersal case showed no effect of dispersal: those that had dispersed (i.e., those in the ‘both’ category plus the ‘dispersed’ category) had fGC levels that were indistinguishable from those that had not dispersed (Fig. 1B). These comparisons support our conclusion that rank attainment status has a robust association with fGC, while dispersal status does not, and that the significant estimate for males in the ‘both’ category is driven by males that have attained rank (with or without dispersing).

Consistent with previous analyses, fGC levels were higher in the dry season; 5% lower in the wet season compared to the dry season ($p = 0.009, \text{CI} = 1-10\%$). We did not find any effect of age on fGC levels in the age range we examined (Table 2). Individuals in groups with a large number of males were more likely to have high fGC levels; our models indicate a 2% increase in a male's fGC for every additional male in his social group, ($p < 0.001, \text{CI} = 1-33\%$). In agreement with previous results from this population, high-ranking individuals generally had lower fGC concentrations than low-ranking males; our models indicate a 2% decrease in a male’s fGC for every increase in rank ($p = 0.001, \text{CI} = 1-33\%$).

Fecal T levels (ft) showed somewhat different patterns than fGC levels. Fecal T levels were statistically indistinguishable among the four categories of maturation status for males (Table 3). In other words, neither attaining an adult dominance rank nor dispersing left any lasting detectable effect on ft levels in male baboons (Fig. 1C, D).

3.2. Categorical analysis of variation in fGC and ft metabolites associated with adult rank attainment and natal dispersal

The paired Wilcoxon signed ranked tests revealed no statistically significant differences in either fGC levels ($W = 248, p = 0.41, N = 34$) or ft levels ($W = 208, p = 0.13, N = 34$) in the two post-rank attainment categories (one month post-rank attainment, versus 2-6 months post-rank attainment). Parallel analysis on the post-dispersal period revealed similar trends, i.e. no statistically significant differences in fGC levels in the post-dispersal categories ($W = 73, p = 0.216, N = 14$) or in ft levels ($W = 65, p = 0.463, N = 14$). These results indicate that individuals do not appear to experience elevated fGC or ft levels in the month following either milestone relative to later time periods.

4. Discussion

We asked whether two important maturational milestones in male baboons (natal dispersal and rank attainment) predict hormone levels in post-pubescent males between 6 and 10 years of age. Our well-supported result is that male rank attainment was associated with an increase in fGC concentrations; males that had attained an adult rank had higher fGC than males that had not yet attained an adult rank, when we controlled for the season, their age, their ordinal rank, and the number of males in the group. In male baboons, adult rank attainment is an important event, because males do not acquire reproductive opportunities until they achieve adult rank; but do so almost immediately afterwards, with a median lag time between these two events of 2.5 months (Alberts and Altmann, 1995b; Figs. S1 and S2). Males in this species need not attain one of the highest ranks in order to achieve mate guarding (consortship) opportunities (Alberts et al., 2006; Bercovitch, 1986; Bercovitch, 1988; Bulger, 1993), although adult rank attainment may signal the onset of a rapid rise to high dominance rank (Hamilton and Bulger, 1990). Because dominance rank is a major predictor of male mating and paternity success in baboons, as in many other primate species, attainment of adult dominance rank has important fitness consequences (Alberts, 2012; Alberts et al., 2006, 2003; Bulger, 1993; Cowlishaw and Dunbar, 1991). Our results suggest that once males have attained an adult rank they experience greater energetic demands and/or psychosocial stress than they did prior to attaining this milestone. This result is somewhat puzzling in light of the fact that males prior to rank attainment are presumably emerging from a prolonged growth spurt, but it is consistent with data on behavioral differences associated with rank attainment. Specifically, after rank attainment but not before, males engage in mate guarding episodes, which are known to be energy limiting and time consuming (Alberts et al., 1996; Packer, 1979).

Contrary to our expectations, natal dispersal did not predict fGC levels. We predicted that natal dispersal, like rank attainment, would involve psychosocial stress and energetic demands that are substantially higher than before dispersal, hence that fGC levels would be higher after dispersal. We posit that the apparent lack of an effect of natal dispersal on fGC levels reflects the fact that individuals are well adapted to face the long-term challenges associated with dispersal, and that the short-term stress associated with dispersal is relatively ephemeral and thus difficult to detect unless hormone levels are sampled within hours or a few days of dispersal (see Alberts et al., 1992; Arlet et al., 2009; Bergman et al., 2005; Marty et al., 2017). In addition, in this study most individuals dispersed to adjacent social groups that share the home ranges of their natal group. This may have removed key
of the idea that males' experiences upon immigration may be quite heterogeneous. Van Noordwijk and Van Schaik (1985) report two different types of immigrant males in long-tailed macaques: "unobtrusive immigrants" enter a new group low in the dominance hierarchy and form associations with females over a period of months before challenging high-ranking males, while "bluff immigrants" attempt to take over the highest dominance rank immediately upon entry. Demographic factors in the new group, such as the relative numbers of adult males and adult females, may also be variable, and this variability will influence aggressive and reproductive encounters. Thus, the variable nature of the dispersal experience—how far males travel, what demographic conditions they encounter in the new group, and how much they engage in aggression and reproduction after dispersal—may explain the absence of a clear, sustained pattern of change in hormone levels in post-dispersers compared to pre-dispersers. We were able to control for at least one of these variables—the number of adult males in a group—which acted as a proxy for male-male competition. Despite controlling for this variable, the effect of dispersal was still insignificant.

Fecal testosterone (FT) levels did not change systematically with either rank attainment or natal dispersal. Instead, our results show that FT levels between the ages of 6 and 10 years were relatively simply described by a quadratic function of age, and were higher in the wet season than in the dry season (Table 2). Pappano and Beehner (2014) also report seasonal effects on FT such that male geladas have higher T during the wet season compared to the dry season. Although we cannot rule out the possibility that our data set lacked the power to detect a subtle change in FT in response to these maturational milestones, we can say with confidence that the effects of attaining these milestones were certainly weaker than the effects of age or season. In addition, we found that the dominance rank position a male held at the time of fecal sampling did not predict his FT concentrations. This result contrasts
with previous research that demonstrates a link between high dominance rank and high FT levels in adult male baboons (Gesquiere et al., 2011). However, this contrast certainly stems from the fact that our study subjects here were not all adults; instead, subjects were experiencing a maturational process whose effects predominated over any effect of the rank that the male actually held. Male baboons experience a post-pubertal growth spurt between 6 and 8 years of age, during which they nearly double in body mass (Alberts and Altmann, 1995b; Altmann et al., 2010); this growth spurt is likely to be associated with the anabolic functions of FT (Nelson, 2011). Testosterone is an important predictor of growth during and after puberty in several primate species (chacma baboons and mangabeys: Bernstein et al., 2008; chimpanzees: Copeland et al., 1985; white-faced capuchins: Jack et al., 2014). In our population as in other primates, FT is upregulated at the onset of reproductive maturity, and begins to decline with age starting at around 9.5 years of age (Altmann et al., 2010; Beehner et al., 2009). Beehner et al. (2009) examined overall FT profiles and maturational milestones in both chacma and yellow baboons and found that male baboons generally attain adult rank after they achieve FT levels in the adult range, but before they achieve peak FT levels. This means that FT levels are already increasing in male baboons before they attain adult rank. Our results build on the study by Beehner et al. (2009) by demonstrating that after controlling for other predictors of FT levels, we did not find any effect of either maturational milestone on FT.

In conclusion, our results indicate that male baboons experience marked increases in fGC levels after rank attainment but not after dispersal, and that they experience no marked or persistent changes in FT levels associated with either adult rank attainment or natal dispersal. Our data combined with previous results in this population reveal that males experience a steady increase in FT beginning between 4 and 5 years of age, coincident with the onset of testicular enlargement, which continues throughout the subadult and early adult period, peaks around 9.5 years of age, and then begins to decline with age (Altmann et al., 2010; Beehner et al., 2009; Gesquiere et al., 2005; Onyango et al., 2013b). Fecal GC levels, too, increase with age across the age range studied here, but the attainment of adult dominance rank signals a marked change in these steroid hormones. Increased mate guarding (courtship) will generally produce increased energetic demands. Thus, our results suggest that the elevation in fGC after adult rank attainment results from increased energetic demands on newly adult males rather than increased psychosocial stress.

Additional research will be needed to elucidate more fine-grained or short-lived effects of dispersal and adult rank attainment on steroid hormones in this as well as other group-living species. For instance, fine-grained sampling has shown that male baboons that engage in more frequent mating exhibit higher T levels than those that engage in less mating (Onyango et al., 2013a). Similarly, fine-grained analyses of behavior with densely sampled hormone concentrations (i.e. daily or near-daily fecal samples) could potentially identify variation in fGC or FT associated with inter-individual differences in the attainment of maturational milestones. For example, individuals that travel longer distances should experience more energetic and psychosocial stress (and have higher GC levels) than those who disperse to neighboring areas, and individuals that experience more mating after rank attainment should experience higher GC levels than those who experience less. Near-daily repeated sampling of individuals during and immediately after the dispersal phase will be needed to reveal such inter-individual differences.

Acknowledgments

Mercy Akinyi was supported by a Franklin Mosher Baldwin Fellowship (L.S.B. Leakey Foundation) and a James, B. Duke Fellowship. We also gratefully acknowledge support from the National Science Foundation (grants DEB 0846286, DEB 0846532, DEB 0919200, IOS 0919200 and IOS 1456832) and the National Institute on Aging (grants R01AG034513-01, P01AG031719, and R21AG049396), for support of the long-term research. We also thank Duke University, Princeton University, the Chicago Zoological Society, and the Max Planck Institute for Demographic Research. We thank the Kenya Wildlife Services, Institute of Primate Research, National Museums of Kenya, National Council for Science and Technology, members of the Amboseli-Longido pastoralist communities, Tortillis Camp, Kër & Downey Safaris, Air Kenya, and Safarilink for their cooperation and assistance in Kenya. A number of people contributed to the long-term data collection over the years, and we are grateful to all of them for their dedication and contributions. Partial thanks go to the Amboseli Baboon Project long-term field team (R.S. Mututua, S. Sayialel, and J.K. Warutere), and to V. Somer and T. Wango for their untiring assistance in Nairobi. Karl Finch has provided expertise in database design and management and we are very grateful for his seminal contributions to the development of BABAB, the Baboon Project database. We also thank N. Lern, L. Maryott, and J. Gordon for database management. This research was approved by the IACUC at Princeton University and at Duke University and adhered to all the laws and guidelines of Kenya.

Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.ybehe.2017.07.005.

References


