Monitoring ovarian function and pregnancy in Eld’s deer 
(Cervus eldi thamin) by evaluating urinary steroid 
metabolite excretion

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Summary. Direct radioimmunoassays (RIA) for urinary oestrone conjugates and 
pregnanediol-3α-glucuronide (PdG) were used to study ovarian activity patterns and 
pregnancy in Eld’s deer. In 2 does, urinary metabolite patterns were compared to 
temporal patterns of plasma LH, oestradiol-17β and progesterone. Preovulatory LH 
peaks occurred coincident with behavioural oestrus, and plasma progesterone secretion 
paralleled PdG excretion. Although plasma oestradiol-17β levels fluctuated between 5 
and 10 pg/ml throughout the oestrous cycle, no preovulatory oestrogen surge was 
observed. Based on PdG excretion, non-conception oestrous cycles averaged 21.5 ± 
2.1 days (± s.e.m., n = 65); however, 2 of 13 does exhibited prolonged oestrous cycles 
(30.1 ± 4.4 days; range 14–62 days, n = 14) characterized by sustained PdG excretion. 
Excluding these 2 females, the mean oestrous cycle was 18.5 ± 0.3 days (range 14–23 
days, n = 51). Behavioural oestrus (12–24 h duration) was observed in 42 of 65 cycles 
(64.6%), and always corresponded with intercyclic troughs in PdG excretion (2–5 days 
duration). Mean gestation duration (n = 10) was 33.5 ± 0.4 weeks. PdG concentrations 
increased (P < 0.05) by Week −32 (3rd week of gestation), plateaued between Weeks 
−31 and −25, increased (P < 0.05) markedly by Week −22 and then rose steadily 
until parturition, declining (P < 0.05) rapidly thereafter. Mean excretion of oestrone 
conjugates remained low until Week −30, increased (P < 0.05) steadily to Week −24 
(P < 0.05) and then returned to baseline by Week −17. Increased (P < 0.05) oestrone 
conjugates concentrations were detected again by Week −4 followed by a rapid 
increase to peak pregnancy levels by Week −1, declining (P < 0.05) precipitously after 
parturition. The results confirm that the Eld’s deer is seasonally polyoestrous with 
onset (January–March) and cessation (August–October) of regular, cyclic ovarian 
activity coinciding with increasing and decreasing daylengths, respectively. Urinary 
PdG excretion accurately reflects cyclic ovarian activity and markedly elevated concen-
trations of this metabolite provide an accurate index of pregnancy. The simultaneous 
monitoring of oestrone conjugates appears useful for estimating the stage of pregnancy 
and predicting parturition onset.

Keywords: Eld’s deer; oestrogen; progesterone; urinary steroids; LH; seasonality

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Introduction

Although once ranging from Manipur in eastern India to Indochina and southern China (between 3° and 25°N latitude), the Eld’s deer (Cervus eldi) is endangered and exists primarily in small herds maintained in captivity (Wemmer & Grodinsky, 1988). Behavioural studies suggest that does are seasonally polyoestrous and have an oestrous cycle of ~17 days in length. Oestrous behaviour generally is observed in late winter or early spring, and gestation lasts ~35 weeks with 80% of births occurring between September and November (Wemmer & Grodinsky, 1988). Although basic captive management protocols have been established for the Eld’s deer, reproductive endocrinology of the species has not been studied.

Hormonal analysis of serial blood samples has been used to study reproductive patterns in red deer (Guinness et al., 1971; Kelly et al., 1982), roe deer (Sempere, 1977; Hoffman et al., 1978; Schams et al., 1980), white-tailed deer (Plotka et al., 1977a, b, 1980), fallow deer (Asher, 1985; Asher et al., 1986), Père David’s deer (Curlewis et al., 1988), reindeer and caribou (McEwan & Whitehead, 1979). Because of their nervous behaviour and susceptibility to the manipulatory stresses associated with capture and restraint, cervid species are difficult to study. In white-tailed deer (Plotka et al., 1983) and fallow deer (Asher et al., 1989) does, a substantial proportion of serum progesterone is secreted from the adrenal glands. This suggests that long-term, sequential blood sampling, which often is impractical for monitoring reproductive patterns in wildlife species, can result in data confounded by the stresses associated with animal handling.

Excretory patterns of urinary oestrone conjugates and/or pregnanediol-3a-glucuronide (PDG), determined by direct radioimmunoassay (RIA), have been found to reflect reproductive activity accurately in the scimitar-horned oryx, bongo, okapi, giraffe (Laskutoff et al., 1982, 1983, 1986), Indian rhinoceros (Kasman et al., 1986), black rhinoceros (Ramsay et al., 1987), tapir (Kasman et al., 1985) and domestic horse (Evans et al., 1984; Kirkpatrick et al., 1988). This approach eliminates the stress associated with frequent venepuncture and permits daily, long-term tracking of endocrine activity. The present study applied these techniques to documenting the circannual reproductive patterns of the female Eld’s deer, a tropical, ‘long-day’ breeding species. A longitudinal protocol was developed to identify seasonal reproductive activity, define the length of the oestrous cycle and diagnose pregnancy and parturition onset.

Materials and Methods

Animals and sample collection. Thirteen hand-reared, female Eld’s deer (2–8 years of age; 50–90 kg body weight) were housed at the National Zoological Park’s Conservation and Research Center, Front Royal, VA (38°N latitude). Does were maintained individually in stalls (3.4 × 4.6 m) connected to outdoor enclosures (3.6 × 36.6 m) and were exposed to normal fluctuations in photoperiod. All females were in visual and olfactory proximity to each other and one or more intact males. Diet consisted of alfalfa hay and herbivore chow (15.5% crude protein, 3.0% crude fat, 16.0% crude fibre; Ralston Purina Co., St Louis, MO, USA) and free access to a mineral block and water. Does were observed twice daily (early morning, late afternoon) for behavioural signs of oestrus which included scent marking of technicians with preorbital gland secretions and assuming a humped-back, lordotic stance when pressure was applied to the hindquarters.

Urine samples (1–5 ml) were collected daily (07:00–11:00 h) by midstream catch or by aspirating urine pooled on enclosure floors. Eight non-pregnant females were sampled for periods ranging from 36 to 64 weeks. Ten females were monitored at different intervals during pregnancy ranging from 4 to 37 weeks before parturition through 7 weeks post partum. Samples were frozen immediately after collection (−20°C), stored for 1–7 days, thawed, centrifuged briefly (1500 g, 10 min) to remove particulate matter and then assayed.

For validation purposes, urinary hormone excretion patterns were compared with temporal hormone profiles in the peripheral circulation. Serial blood samples (8 ml) were collected by jugular venepuncture from 2 females sedated with xylazine (Rompun®; Haver, Shawnee, KA, USA; 5–10 mg/female, administered i.m.). Sampling was initiated in August, and proceeded at 48 h intervals until 2 behavioural oestrous periods were observed for each doe. Plasma was collected after centrifugation and immediately stored frozen (−70°C) until assayed.

Creatinine assay. To compensate for variations in urine concentration, urinary creatinine (Cr) was determined using a modified Jaffe reaction (Taussky, 1954). Diluted urine samples (100 μl, 1:100) were combined in duplicate (in 96-well microplates) with 50 μl each picric acid (0.40 M) and NaOH (0.75 M). After a 20 min incubation, absorbance
(490 mm) was measured on an MR600 microplate reader (Dynatech, Chantilly, VA, USA). Hormone levels were divided by creatinine concentrations and values expressed as mass units/mg Cr. Urinary creatinine concentrations averaged 1.37 ± 0.05 mg Cr/ml urine (n = 3362 samples) and inter- and intra-assay coefficients were 10.3% (n = 20) and ≤5%, respectively.

**Oestrone conjugates assay.** Urinary oestrone conjugates were analysed with minor modifications of the RIA described by Shideler et al. (1983). Identification of specific oestrogen metabolites for Eld’s deer was not performed by chromatographic separation, so only oestrone conjugates immunoreactivity was assessed. Initially, urine samples were diluted (1:50) in phosphate-buffered saline (PBS; 0-1 M, 0-1% gelatin, pH 7.0), and an aliquant (non-pregnant and early pregnant, 150 µl; late pregnant, 20 µl) was adjusted to a final assay volume of 300 µl in Tris buffer (0-1 M-Tris, 0-9% NaCl, 0-1% NaH₂PO₄, 0-1% gelatin, pH 8.4). The antiserum which cross-reacts 100% with oestrone glucuronide and oestrone sulphate (anti-oestrone-3-glucuronide serum, 100 µl; 1:1500: D. Collins, Emory University, Atlanta, GA, USA) and [³H]oestrone sulphate (100 µl, 7000 c.p.m., sp. act. 55 Ci/mmol: Dupont-New England Nuclear, Wilmington, DE, USA) were combined with unknowns and standards (4-9-2500 pg/tube, Sigma Chemical Co., St. Louis, MO, USA) and incubated overnight at 4°C. Following the addition of 300 µl charcoal–dextran (0-0625% Norit A charcoal, 0-00625% dextran in 0-1 M-PBS, pH 7.0) and a 30-min incubation at 4°C, tubes were centrifuged (10 min, 1500 g), decanted into scintillation vials, combined with 5-0 ml Ready Solv HPb (Beckman Instruments Inc., Fullerton, CA, USA) and counted for 5 min.

Serial dilutions of Eld’s deer urine yielded displacement curves parallel to that obtained for oestrone sulphate standards. The mean ± s.e.m. recovery of oestrone sulphate (range; 4-9–2500 pg/tube) added to a pool of Eld’s deer urine (1:50) was 92.4 ± 3.9% (y = 0.98x - 32.97; r = 0.99; P < 0.001). Assay sensitivity was 4-9 pg/tube, inter-assay coefficients of variation for 2 separate internal controls were both 11.6% (n = 38) and intra-assay variation averaged ≤10%.

**Pregnanediol-3a-glucuronide assay.** Urinary PdG immunoreactivity was assessed using the methods of Mitchell et al. (1982) and Loskutoff et al. (1982). A sample of urine diluted (1:50) in PBS (non-pregnant, 100 µl; pregnant, 25 µl) was combined with 100 µl PdG antiserum which cross-reacts 100% with PdG and 6-7% with pregnadiol (02/2Zoo, 1:20 000) and 100 µl [³H]PdG (7000 c.p.m., sp. act. 42 Ci/mmol) supplied by Courtauld Institute of Biochemistry, London, UK. Urines and standards (19-5–5000 pg/tube, Sigma Chemical Co.) were incubated overnight (4°C) and antibody-bound and free steroid were separated after a 45-min incubation with 300 µl charcoal–dextran suspension and centrifugation for 10 min (1500 g). Supernatants were combined with 5 ml Ready Solv HPb and counted for 5 min.

Serial dilutions of Eld’s deer urine yielded displacement curves parallel to that obtained with standard preparations. Recovery of known amounts of PdG (range; 19-5–5000 pg/tube) added to a pool of diluted urine (100 µl, 1:50) gave a mean ± s.e.m. value of 95.3 ± 2.5% (y = 0.97x - 12.66; r = 1.0; P < 0.001). Assay sensitivity was 19-5 pg/tube and inter-assay coefficients of variation were 7.4% (n = 55) and 12.6% (n = 57) for 2 separate internal controls, and intra-assay variation was ≤10%.

**Luteinizing hormone assay.** Plasma LH was analysed using the procedure of Niswender et al. (1969). Serial dilutions of Eld’s deer plasma yielded displacement curves parallel to that obtained with NIH-LH-S18 standards. The mean ± s.e.m. recovery of LH (range, 0-03–4 ng/tube) added to 100 µl pooled Eld’s deer plasma was 98.2 ± 3.1% (y = 0.98x - 1-51; r = 0.99; P < 0.001). Assay sensitivity was 0-03 ng/tube. The inter- and intra-assay coefficients of variation were 8.5% (n = 6) and ≤10%, respectively.

**Oestradiol-17β and progesterone assays.** Plasma oestradiol-17β and progesterone were analysed in unextracted plasma using commercially available RIA kits (Radioimmunoassay Laboratories, Carson, CA, USA). For the oestradiol-17β assay, the antiserum was anti-6- keto-oestradiol-17β-6-oxime–BSA which cross-reacts 100% with oestradiol-17β, 20% with oestrone and 1-5% with oestriol. Serial dilutions of unextracted Eld’s deer serum (spiked with 1000 pg oestradiol-17β/ml) yielded displacement curves parallel to that obtained for oestradiol-17β standards prepared in Eld’s deer plasma. The mean ± s.e.m. recovery of oestradiol-17β (range 5-3000 pg/ml) added to 100 µl pooled Eld’s deer plasma was 92.7 ± 3.6% (y = 1.03x - 10.51, r = 0.99, P < 0.001). Assay sensitivity was 5 pg/ml; all samples with undetectable oestradiol-17β values were assigned a value of 5 pg/ml for analysis and presentation. The inter- and intra-assay coefficients were 8.5% (n = 6) and ≤10%, respectively.

For the progesterone analysis, antiserum was anti-11 hydroxyprogesterone-11-hemisuccinate–HSA which cross-reacts 100% with progesterone, 5-4% with 20α-dihydroprogesterone and 3-8% with desoxycorticosterone. Serial dilutions of unextracted Eld’s deer plasma yielded displacement curves parallel to that obtained for progesterone standards prepared in Eld’s deer plasma. The mean ± s.e.m. recovery of progesterone (range, 0-2–50 ng/ml) added to 100 µl pooled Eld’s deer plasma was 93.3 ± 3.1% (y = 0.9x - 1.31, r = 0.99, P < 0.001). Assay sensitivity was 0-2 ng/tube. The inter- and intra-assay coefficients of variation were 7.5% (n = 6) and ≤10%, respectively.

**Definitions and statistical analyses.** Oestrous cycle length was determined by measuring the interval between successive intercyclic nadirs in PdG excretion, and Day 1 of the cycle was defined as the first day PdG excretion returned to baseline concentrations (< 10 ng/ml Cr). Gestation duration was considered as the interval from observed copulation until the birth of an offspring. Statistics were performed using Statview 512 +™ (Version 1·1, BrainPower, Inc. 1986, Calabasas, CA, USA) on an Apple® Macintosh Plus Computer (Cupertino, CA, USA). Standard descriptive statistics including mean and standard error of the mean (s.e.m.) were used to describe hormonal metabolite values. Linear regression analyses were used to evaluate the correspondence between serum progesterone and urinary PdG. Urine
values were used to predict serum values collected on the same day and 1 day apart to account for a potential lag in urinary metabolite excretion. Because serum samples were not collected every day, paired values vary from 11 to 16 points per individual. Differences among mean weekly hormone concentrations were determined by 1-way analysis of variance, and multiple pairwise comparisons between weeks were analysed using Fisher’s Protected Least Significant Difference Test (Fisher, 1966).

**Results**

Urinary PdG values coincided with circulating progesterone collected on the same day in the 2 does subjected to urine and blood collection (Doe 1: \( r = 0.736, P < 0.01; \) Doe 2: \( r = 0.805, P < 0.001 \)). Urinary PdG was a better predictor of serum progesterone when the former values lagged by 24 h (Doe 1: \( r = 0.828, P < 0.002; \) Doe 2: \( r = 0.895, P < 0.001 \)). As demonstrated below, concentrations of oestrone conjugates in urine were ineffective for monitoring reproductive cyclicity of Eld’s deer. Plasma progesterone, urinary PdG and plasma oestradiol-17\( \beta \) and LH profiles from one female are presented in Fig. 1. Behavioural oestrus occurred when plasma progesterone concentrations were at nadir (~0.25 ng/ml). Following presumed ovulation, plasma progesterone eventually peaked at 1.8 ng/ml. The urinary PdG excretory profile was similar qualitatively to plasma progesterone; PdG concentrations increased to ~70 ng/mg Cr during the luteal phase, decreased to <10 ng/mg Cr during the follicular phase with basal levels also occurring coincident with oestrus. Plasma LH concentrations remained below 0.25 ng/ml during the follicular and luteal phase but increased approximately 6-fold during a presumed preovulatory peak (1.5 ng/ml) coinciding with behavioural oestrus. Plasma oestradiol-17\( \beta \) concentrations in both does fluctuated between 5 and 10 pg/ml during the sampling interval with no evidence of a pre-LH surge (Fig. 1).

![Fig. 1. Concentrations of plasma LH, progesterone and oestradiol-17\( \beta \) and urinary PdG from a single female sampled for 1 month. Plasma samples were collected at 48-h intervals and urine samples were collected daily.](image-url)
Fig. 2. Annual non-conception PdG profiles from 3 Eld's deer does sampled daily. Arrows indicate observations of behavioural oestrus and annual changes in daylength are presented in (a). Two regularly cyclic females are represented in (b) and (c); a female with consecutive prolonged luteal phases is presented in (d).
Figure 2(a–d) depicts longitudinal PdG excretion profiles during non-conception oestrous cycles in 3 representative females monitored for 1 year. Based on variations in this metabolite, onset (January–March) and cessation (August–October) of regular cyclic hormonal activity coincided with increasing and decreasing daylengths, respectively (Fig. 2a). The overall mean of 65 oestrous cycles for the 8 females was 21.5 ± 2.1 days. However, 2 does consistently exhibited prolonged oestrous cycles (average 30.1 ± 4.4 days; range 14–62 days, n = 14) characterized by sustained elevations in PdG excretion (Fig. 2d). Excluding these cycles, the remaining females had a mean oestrous cycle duration of 18.5 ± 0.3 days (range 14–23 days, n = 51). Behavioural oestrus (duration ~12–24 h) was observed in 42 of 65 (64.6%) of the cycles, and in all cases coincided with intercyclic troughs in PdG excretion which lasted from 2 to 5 days. Only 2 does failed to exhibit signs of behavioural oestrus during the breeding season. The temporal decrease in PdG indicated that the onset of luteal regression began 3–5 days before the next observed oestrus. A maximum of 12 consecutive hormonal cycles was observed (range, 6–12) in a single individual, and no female exhibited regular metabolite fluctuations later than October.

During the breeding season, peak non-pregnant PdG concentrations ranged from ~40 to 180 ng/mg Cr in non-pregnant does. At 1–6 weeks before the first behavioural oestrus and normal duration luteal phase, does always exhibited 1–3 transient increases in PdG excretion (peak, ≤50 ng/mg Cr) of several days duration (range, 2–5 days, Fig. 2b, c, d). Females also tended to exhibit an irregular excretory pattern of PdG following the final oestrus of the season which often was characterized by transient increases in PdG excretion similar to those observed before the first oestrus (Fig. 2c).

Although urine from all 8 non-pregnant females contained detectable quantities of oestrone conjugates (range 1.2–57.9 ng/mg Cr), no cyclic patterns of oestrone conjugates excretion were observed. Daily urinary oestrone conjugates and PdG values in 2 non-pregnant females sampled for 3 months, beginning in March illustrated that only the latter metabolite was useful for tracking regular cyclic activity (Fig. 3).

Weekly mean PdG and oestrone conjugates values for the 10 pregnant does are presented in Fig. 4 with weekly values aligned to the day of parturition. Based on known copulation dates for 6 females, the mean duration of gestation was 33.5 ± 0.4 weeks (range 31.7–34.4 weeks). Between-animal variation in mean oestrone conjugates excretion over time was apparent (P < 0.003) while PdG levels were similar (P > 0.10) among individual pregnant females. Mean PdG concentrations increased (P < 0.05) from 18.7 ± 2.6 ng/mg Cr (Week −34, estimated time of conception) to 50.8 ± 6.3 ng/mg Cr by Week −32 (3rd week of gestation). PdG excretion plateaued between Week −31 and Week −25 (range 64.4–75.2 ng/mg Cr) before increasing markedly by Week −22 (156.0 ng/mg Cr, P < 0.05) and then steadily rising to peak concentrations by Week −2 (2666.0 ± 281.5 ng/mg Cr). After parturition, mean PdG concentrations declined abruptly to 167.0 ± 33.8 ng/mg Cr (Week 1, P < 0.05) and were indistinguishable from values in non-pregnant does by Week 7 post partum (22.4 ± 4.2 ng/mg Cr, P < 0.05).

Mean excretion of oestrone conjugates remained low during the first 5 weeks of gestation (Weeks −34 to −30, 9.6 ± 0.5 ng/mg Cr), increased steadily to Week −34 (43.5 ± 6.1, P < 0.05) and then returned to baseline from Week −17 (10.0 ± 2.1 ng/mg Cr, P < 0.05) to Week −5 (17.5 ± 1.3 ng/mg Cr). Concentrations again increased ~30-fold between Week −4 (57.3 ± 11.1 ng/mg Cr) and Week −1 (2225.0 ± 199.7 ng/mg Cr, P < 0.05) but declined precipitously immediately post partum (Week 1, 160.3 ± 33.7, P < 0.05), and were not different from values in non-pregnant does by Week 7 (24.7 ± 6.7 ng/mg Cr, P > 0.05).

A representative, longitudinal excretory profile for PdG and oestrone conjugates is depicted in Fig. 5 (−37 weeks before parturition through 14 weeks post partum) illustrating both the temporal and quantitative dynamics of hormonal excretion in a doe experiencing 2 non-conception reproductive cycles followed by conception and early pregnancy. After parturition, several 6–8-day transient increases in PdG were observed which were unaccompanied by behavioural oestrus.
**Fig. 3.** Urinary concentrations of oestrone conjugates and PdG in 2 representative females sampled over a 3 month interval. Arrows denote observations of behavioural oestrus.

**Fig. 4.** Weekly (mean ± s.e.m.) composite urinary PdG and oestrone conjugates profile for 10 does sampled throughout pregnancy. All values are aligned to the day of parturition.

**Discussion**

Longitudinal, parallel measures of urinary oestrone conjugates and PdG metabolites were useful for studying reproductive seasonality, pregnancy and parturition in the Eld’s deer. These data
confirmed that this species is seasonally polyoestrous and a spontaneous ovulator. Oestrous cycles commence in late winter/early spring, and the potential breeding season lasts 8–9 months (January–September). The overall mean cycle length for the Eld’s deer doe was 21.5 days which was within the range reported for red deer (18.3 days: Guinness et al., 1971; 21 days: Adam et al., 1985), black-tailed deer (22–29 days: Thomas & Cowan, 1975), white-tailed deer (28 days: Plotka et al., 1980), fallow deer (22.4 days: Asher, 1985) and Père David’s deer (19.5 days: Curlewis et al., 1988). However, there was considerable variation among does which could explain the variability in cycle length reported in earlier cervid studies. The prolonged luteal-phase increases in PdG excretion (34–62 days) observed in 2 Eld’s deer in our study were similar to the extended luteal phase durations reported in red deer (range 34–59 days: Guinness et al., 1971; 35 days: Adam et al., 1985) and Père David’s deer (45–60 days: Curlewis et al., 1988), presumably reflecting the presence of persistent corpora lutea (CL). For the Eld’s deer, excluding these 2 does, the mean cycle length was reduced to 18.5 days which was more consistent with the observations of Wemmer & Grodinsky (1988) who based their estimates on behavioural patterns of sexual receptivity. Oestrous behaviour coincided with troughs in PdG excretion (2–5 days duration) which was analogous to observations of baseline circulating progesterone at the time of oestrus in white-tailed deer (Plotka et al., 1980), fallow deer (Asher, 1985) and Père David’s deer (Curlewis et al., 1988).

The transient, short-lived elevations in PdG occurring before the first full-length luteal phase in Eld’s deer were similar to short waves of serum progesterone preceding first oestrus in red deer (Morrison, 1960), moose (Simkin, 1965), black-tailed deer (Thomas & Cowan, 1975), white-tailed deer (Plotka et al., 1977b), reindeer and caribou (McEwen & Whitehead, 1979), fallow deer (Asher, 1985) and Père David’s deer (Curlewis et al., 1988). For the black-tailed deer, these abbreviated elevations in progesterone have been attributed to ‘silent’ oestrus and luteinization (but not ovulation) of ovarian follicles (Thomas & Cowan, 1975). These transient waves of progesterone in fallow deer have been associated with the presence of CL which develop before first oestrus but fail to achieve or sustain full luteal function (Asher, 1985). The observation of transient waves of PdG excretion near the termination of the Eld’s deer breeding season coincided with similar serum progesterone data in fallow deer (Asher, 1985) and may reflect a gradually reduced sensitivity of the hypothalamo–pituitary–ovarian axis as a result of the reinitiation of short-day, photoperiodic cues. It is important that these short waves of luteal function, which appear necessary for triggering later, full reproductive function, can be detected by urinary PdG excretion, thereby offering an extremely valuable tool for elucidating the mechanisms of reproductive seasonality in non-tractable species.
In white-tailed deer, a broad 3–4-day increase in serum oestradiol-17β values has been reported during the follicular phase which is less distinct than the same hormone pattern reported for sheep and cattle (Plotka et al., 1977b). Although Plotka et al. (1977b) reported elevated mean oestradiol-17β concentrations for the 3-day period immediately before oestrus in white-tailed deer, daily concentrations did not vary significantly at any stage of the oestrous cycle, reflecting the absence of a convincing preovulatory surge. Similarly, in red deer (Kelly et al., 1982) and fallow deer (Asher et al., 1986), no significant preovulatory increase in serum oestradiol-17β coincided with oestrus. The absence of a preovulatory increase in plasma oestradiol-17β in the present study may reflect the relatively infrequent blood sampling protocol or an inadequacy in assay sensitivity. Nevertheless, longitudinal excretory profiles of oestrone conjugates were not useful for monitoring ovarian function in non-pregnant Eld’s deer, and this emphasizes the need to identify specific oestrogen secretion and excretion mechanisms in cervids.

Marked increases in PdG and oestrone conjugates excretion occurred during pregnancy with peak concentrations observed immediately before parturition. Increased progesterone production also has been reported for pregnant roe deer (Hoffman et al., 1978), red deer (Kelly et al., 1982), white-tailed deer (Plotka et al., 1977b), reindeer and caribou (McEwan & Whitehead, 1979), although there are important differences in qualitative secretory profiles among species. White-tailed deer (Plotka et al., 1977b, 1982) exhibit a relatively constant secretory pattern of progesterone and require luteal-derived progesterone for pregnancy maintenance. Likewise, because the amount of progesterone secreted by pregnant red deer (Kelly et al., 1982) is directly proportional to CL number, luteal tissue probably accounts for most of the progesterone produced during pregnancy in this species. In contrast, Eld’s deer exhibited PdG excretory patterns during pregnancy similar to serum progesterone profiles in roe deer, a species in which CL appear functional throughout gestation. Hoffman et al. (1978) reported that circulating progesterone in roe deer increases to a plateau during early pregnancy, is sustained for about 20 weeks and then increases again following embryonic attachment, making the roe deer the only artiodactyl known to experience obligate delayed implantation (Sempere, 1977). Whether the PdG excretory profile observed during early pregnancy in the present study reflects an abbreviated delay in implantation or a short delay in fetal-placental steroid biosynthesis is unknown. Biphasic elevations in oestrogen during pregnancy, similar to the increases in excreted oestrone conjugates observed in our study, have not been reported for other cervids. The early peak (−29 to −19 weeks) of oestrone conjugates in the Eld’s deer occurred coincident with the secondary rise in PdG following the 8-week plateau period. Perhaps these excretory profiles reflect differential secretion by the ovary and placenta.

Increased oestrogen production during late pregnancy is thought to be involved in the initiation of parturition in polyoestrous ungulates including sheep (Challis, 1971) and cattle (Edqvist et al., 1973; Smith et al., 1973; Dobson & Dean, 1974; Robertson, 1974). The increase in urinary oestrone conjugates observed in Eld’s deer (∼40-fold) during the last 5–6 weeks of pregnancy is temporally similar to increases in serum oestrogen concentrations measured in red deer (∼5-fold: Kelly et al., 1982) and white-tailed deer (∼7-fold: Harder & Woolf, 1976; ∼10-fold: Plotka et al., 1977a, b). In contrast, oestrogen levels in roe deer increased steadily during the last 6 months of gestation (∼2-fold: Hoffman et al., 1978). The sharp increase in oestrone conjugates observed in all pregnant Eld’s deer provided a reliable method of staging pregnancy and predicting impending parturition.

In summary, direct RIA of urinary hormonal metabolites provides a practical, stress-free approach for evaluating reproductive function in the female Eld’s deer. This strategy should permit more detailed studies of the mechanisms controlling seasonality in other stress-susceptible cervid species.

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