The effect of dietary protein on reproduction in the mare. V. Endocrine changes and conception during the early post partum period

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ABSTRACT
Pregnant Anglo-Arab and Thoroughbred mares (n = 24) were divided randomly according to age and breed into 4 groups of 6 mares each from approximately 6 weeks before their expected foaling date. Diets received by the 4 groups varied in essential amino-acid and total protein contents. Serum progestagen, FSH and LH concentrations were determined from the day of parturition until foal heat and during the 1st oestrous cycle following foal heat. Serum progestagen, FSH and LH concentrations did not differ between the treatment groups. Progestagen concentrations were high (7 ± 0.5–16.4 ng/mL) at parturition but decreased rapidly within 48 h. As progestagen concentrations decreased LH concentrations increased from Days 3–6 post partum to reach maximum values at, or the day after ovulation. FSH concentrations declined 3–4 d after parturition and increased 2–3 d before ovulation at foal heat. The duration of elevated progestagen concentrations during the luteal phase of the subsequent oestrous cycle affected the interovulatory period. A 12–14 d FSH cyclical releasing pattern occurred. Season/photoperiod affected the resumption of normal oestrous cyclicity during the post partum period. The duration of the 1st oestrous cycle after foal heat in mares fed a low-quality protein diet showed a greater range (13–30 d) compared to mares fed a high-quality protein diet (18–26 d).

Key words: equine, FSH, LH, post partum period, protein nutrition, serum progestagens.


INTRODUCTION
The duration of pregnancy of c. 11 months in the Thoroughbred mare implies that the mare must conceive within 30 days post partum in order to produce a foal each year. The post partum interval is described as the time after parturition when the uterus undergoes involution to the state where pregnancy can be maintained. Late pregnancy, parturition and the early post partum period are characterised by major changes in the reproductively and endocrine systems. Foal heat occurs within 7–9 d of parturition in more than 90 % of mares. Although mares will ovulate and conceive during foal heat, several studies indicated a reduced pregnancy rate and embryonic loss that may be associated with the actual state of the uterine environment.

The reproductive efficiency of lactating mares is affected by both season/photoperiod and nutrition. Approximately 28 % of Thoroughbred mares that foal early in spring do not show signs of oestrus. Optimal energy intake of lactating mares is important because this is the period when conception should take place. The bodily condition of the mare at foaling is therefore important, as mares in poor condition at foaling may maintain condition at the expense of milk production. Conception is improved when mares are maintained in a positive energy balance during lactation and by limiting the period of suckling by the foal. Studies have indicated that a protein deficiency in the diet might contribute to early embryonic loss irrespective of the mare’s bodily condition.

The purpose of this study was to determine the normal clinical, morphological and endocrine changes that occur in the reproductive functions of the mare during the post partum period until conception. The effects of season at the time of foaling and dietary protein quality and intake were also investigated.

MATERIALS AND METHODS
Pregnant Thoroughbred and Anglo-Arab mares (n = 24) were kept on pasture until 6 weeks before their expected foaling dates and were then allocated randomly according to age and breed to 1 of 4 dietary groups of 6 mares each. The 4 diets consisted of: Group 1: tef hay (6 kg) and cubes (4 kg); Group 2: lucerne hay (6 kg) and cubes (4 kg); Group 3: tef hay (4 kg), cubes (6 kg) and fishmeal (200 g); Group 4: lucerne hay (6 kg), cubes (4 kg) and fishmeal (200 g). These 4 feeding regimens were chosen to obtain a wide range of dietary amino-acid contents. Tef hay is generally used as a source of roughage in large parts of South Africa. Its protein content is low compared to lucerne hay. The crude protein and amino-acid intake of mares in the different dietary groups has been described. The cubes were offered in 2 equal portions in the morning and afternoon. After the mares had foaled, the cubes were increased to 6 kg fed 3 times a day in equal portions and the hay to 7 kg. While the fishmeal remained at 200 g/day. Each mare and foal received their cubes separately, but during the morning feed, when the mares received their fishmeal, the foals were prevented from eating with the mares. Throughout the trial the mares finished all feed offered to them every day. At 3 months of age the foals received 1 kg of cubes and c. 2 kg of roughage per day. Each group of mares was kept in a 30 × 50 m paddock. The foaling date was recorded in all cases. Daily teasing commenced 2 d after parturition and continued until 90 d of pregnancy. The ovaries of the mares in oestrus were palpated daily. The ovaries of mares that had not shown signs of oestrus by 10 d post partum were palpated weekly until signs of oestrus became evident. During examinations the size and consistency of the ovaries and follicles present were determined as described previously. A Squibb Model 2000 scanner with a 3 MHz probe was...
used for all ultrasonographic examinations.

Mares were mated during the 1st oestrus following foal heat. If a mare did not show signs of oestrus within 12 d post partum she was mated during the 1st oestrous period. Mares were mated when a 30–50 mm, soft, thin-walled follicle was palpable and daily thereafter until ovulation occurred, after which mating ceased.

Blood samples were collected on the day of parturition and every 2nd day thereafter until signs of oestrus were evident. Samples were collected daily during oestrus, and then every 3rd day until the following oestrous period. This procedure was followed until pregnancy was diagnosed. Blood samples were collected and the serum stored as described previously.

All mares and foals were weighed on the day of foaling and weekly thereafter until the foals were weaned at 6 months.

Hormone assays

Total unconjugated serum progesterone

Serum samples were analysed for total unconjugated progesterones. The radio-immunoassay procedure was principally as described by Youssefnejadian et al. using an antibody generated in sheep according to the method of Odell et al. against 11 α-hydroxyprogesterone hemi-succinate BSA as modified by Faure. In addition, the antibody shows significant cross-reactivity with α-pregnane-3,20-dione, and to a much lesser extent with other pregnane derivatives of progesterone that are known to occur in the pregnant mare. Cross-reactivity with major adrenocortical C18 and C19 steroids is for all practical purposes nonexistent.

Follicle-stimulating hormone (FSH)

FSH concentrations were determined using the Amerlex-M FSH (Code DM 3070/3071) kit (Amersham, UK). Use of this antibody was based on the results of Alexander et al.

Luteinising hormone (LH)

LH concentrations were determined using the method described by Niswender et al. as modified by Visser.

Statistical analysis

Analysis of variance was conducted using the LSML 76 computer programme.

RESULTS

The mean body weight of mares on the day of foaling was: Group 1, 437 kg; Group 2, 472 kg; Group 3, 441 kg; Group 4, 454 kg. The mean birth weight of the foals of the mares in each group was: Group 1, 47.3 kg; Group 2, 47.2 kg; Group 3, 51.2 kg; Group 4, 46.2 kg.

The number of days from parturition until 1st signs of oestrus (foal heat), the duration of the oestrous periods, time of ovulation, diameter of follicles and the duration of the oestrous cycle following foal heat are shown in Table 1 for the lactating mares in the 4 dietary groups. The results that include the 2 mares (No. 8 in Group 2 and No. 20 in Group 4) that did not show signs of foal heat or oestrus until 80 d and 64 d post partum respectively, are shown separately in Table 1.

No differences in the diameter (mm) of the largest follicle at the onset of oestrus and the diameter of the largest follicle on the day before ovulation were found between groups.

In Table 2 a comparison is made between the onset of 1st signs of oestrus and the time of ovulation in mares that foaled during August and September (transitional period) and mares that foaled during October to December (physiological breeding season), irrespective of their nutritional group. Those that foaled in the transitional period showed the 1st signs of oestrus on average 24 d post partum, compared to 9 d in those that foaled in the physiological breeding season, and on average the 1st ovulations occurred 27.1 d and 14.6 d post partum.
respectively. The mean duration of the 1st oestrous period for the early and late foalers was 5.1 d and 6.8 d respectively.

No differences in serum progestagen concentrations were found during the time from parturition to foal heat among the 4 nutritional groups. The mean serum progestagen concentrations of mares (n = 18) that ovulated on Days 6, 9–11, 12–13 and 15–19 post partum are shown in Fig. 1.

The progestagen concentrations of mares during the 1st oestrous cycle after foal heat did not differ between the nutritional groups, and therefore the results of mares in all 4 dietary groups were pooled.

During foal heat, serum progestagen concentrations increased within 2 days post-foaling. Serum progestagen concentrations were 7–20 ng/ml at 4–6 days post-ovulation and were maintained at these concentrations for 13–15 days, after which they decreased to reach baseline values c. 2 days before the next ovulation. After pooling the results, 4 different patterns of progestagen secretion were identified according to the duration of the interovulatory period, namely A: 17–19 d; B: 20–22 d; C: 24–25 d, and D: 29–30 d (Fig. 2).

Persistent CLs were identified in 2 mares. Mare No. 8 (Group 2: lucerne hay and cubes), which foaled in mid-August, ovulated on Day 9 post partum without showing signs of oestrus for the following 80 days and had low (<5 ng/ml) serum progestagen concentrations during this period (Fig. 3). Mare No. 20 (Group 4: lucerne hay, cubes and fishmeal), which foaled in September, showed no signs of oestrus but ovulated on Day 11 post partum. She then maintained low (<5 ng/ml) serum progestagen concentrations until signs of oestrus and she ovulated on Day 66 post partum (Fig. 3).

Serum LH and FSH concentrations were established only in mares in Group 1 (lowest quality protein) and Group 4 (highest quality protein), from parturition until foal heat and the following oestrous cycle until after ovulation. No differences in the LH concentrations between dietary groups that could be attributed to protein nutrition were found, and therefore the results were pooled to demonstrate the general secretion pattern. The mean pooled LH and progestagen concentrations from parturition until after the 2nd ovulation are shown in Fig. 4. The mean duration from foaling to ovulation at foal heat was 13.7 d. LH concentrations increased gradually from Day 6 post partum until 2 days before ovulation, when the LH concentrations began to peak. In 11 of the 18 mares the LH peak occurred on the day of ovulation, and in the remaining 8 during the following day. LH concentrations decreased to baseline levels within 48 h post-ovulation.

The serum FSH concentrations were pooled, as no differences were found between mares in the 4 dietary groups from parturition to 1st ovulation (Fig. 3).
Table 3: Conception rates of mares in 4 dietary groups.

<table>
<thead>
<tr>
<th>Dietary group</th>
<th>Oestrous period after parturition when mares were mated</th>
<th>Mated oestrous period per conception</th>
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<tbody>
<tr>
<td>1</td>
<td>1 (100 %) b 5 (80 %) 1 (0 %) 1 (0 %) — —</td>
<td>1.6</td>
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<tr>
<td>2</td>
<td>3 (33 %) 3 (33 %) 2 (50 %) 1 (0 %) 1 (100 %)</td>
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<td>3</td>
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<tr>
<td>4</td>
<td>5 (100 %) — — — — —</td>
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aFoal heat.
bNumber of mares (conception rate of mares mated during cycle).

Fig. 2: Mean serum progestagen concentration (ng/ml) in mares with interovulatory periods of (A) 17–19 d (n = 4), (B) 20–22 d (n = 5), (C) 24–25 d (n = 5) and (D) 29–30 d (n = 2) during the oestrous cycle following foal heat.

Fig. 3: Serum progestagen concentrations (ng/ml) of Mare 8 (Group 1) and Mare 20 (Group 4) that developed persistent corpora lutea after ovulation.
At parturition the mean FSH concentrations were relatively high (25 ng/ml) and increased (30 ng/ml) during the 1st 3 days post partum. In all mares that showed foal heat, FSH concentrations declined after Day 3 to reach minimum concentrations (15 ng/ml) by Day 8. The mean FSH concentrations increased at the onset of foal heat and reached high concentrations (>30 ng/ml) the time of ovulation. Thereafter concentrations declined 1–2 d after ovulation to reach low values (<20 ng/ml) 5–7 d after ovulation. An increase in FSH concentrations with maximum concentrations of approximately 33 ng/ml were found from Days 9–12 of the following cycle, after which the concentrations declined to minimum values (<20 ng/ml) by Day 18. Increases were recorded 2–3 d before the next ovulation (Fig. 5).

The conception rate of the lactating mares is shown in Table 3. Mares in Groups 3 and 4 conceived at the 1st mated oestrous period compared to the 1.6 (Group 1) and 2.5 (Group 2) mated oestrous periods per conception in the groups receiving the low-quality protein diets.

**DISCUSSION**

The total daily protein requirement of a mare with a body weight of 500 kg is 800–870 g/day during late pregnancy and...
1284–1422 g during the 1st 3 months of lactation\(^5\). In the current study the mean body weight of all the mares in the respective dietary groups was below 500 kg immediately after foaling. The total daily crude protein intake of the mares in the 4 dietary groups varied between 1000 and 1800 g/day during late pregnancy, while the lowest daily crude protein intake of 1200 g received by the mares in Group 1 during lactation is considered to meet their requirements\(^6\). It is thus clear that, according to the recommendations of the National Research Council, Washington DC, none of the mares suffered from a protein deficiency during any stage of this trial.

Foal heat normally commences 7–12 d post partum\(^7\). Although there was no significant difference in the number of days from parturition until the onset of foal heat between groups, the mean duration of this period for mares in Group 1 (low-quality protein) was 7 d compared to 10 d in Group 4 (high-quality protein). The 2 mares that showed the 1st signs of oestrus 80 and 64 days after foaling, foaled early during the season (transitional phase) and the prolonged post partum anoestrous periods were probably due to the effect of season/photoperiod and not protein nutrition. The effect of season on the incidence of foal heat has been described in Thoroughbred and Percheron mares\(^8\). A higher incidence of failure to show foal heat was found amongst mares that foaled during early spring and also amongst high milk producers\(^9\). The pooled results (Table 2) indicated that the time of foaling (season) played an important role in the incidence of oestrus. In a previous study with barren mares kept in the same locality, only 10 % of the mares ovulated between 29 August and 20 September, while 70 % ovulated from the last week in September to mid-October\(^9\). When the 2 mares described above are excluded, the current results are in agreement with the literature\(^10\). Protein nutrition apparently had very little effect on the duration of the period from parturition to the onset of foal heat in the mares used in this study.

The average duration of foal heat (Table 1) of 8.8 d for Group 1 (low-quality protein) was approximately 3–4 d longer than that recorded in any of the other groups. Mares in Group 1 ovulated on average 7.3 d after the onset of oestrus, approximately 3 d later than the mares in the other groups. However, when the number of days from parturition to ovulation is compared, no differences were found among groups (Table 1).

Table 1 shows that the duration of oestrous cycles following foal heat was 3–4 d longer in the fishmeal-supplemented Groups 3 and 4 than in Groups 1 and 2, which did not receive fishmeal.

In addition, the duration of the oestrous cycles in the mares in Groups 3 and 4 showed less variation (22–28 d) than mares in Groups 1 and 2, where a range of 13–31 days was recorded. This might indicate improved endocrine function with regard to the production of progesterone and possibly LH, which is important for early CL stimulation after ovulation. This is in agreement with previous results, where high-quality protein decreased the time to 1st ovulation after the anovulatory period in barren mares\(^10\). However, the mean duration of the oestrous cycles was 20.8 d (13–31), which compares favourably with the 20.7 d reported for Thoroughbred mares in the southern hemisphere\(^10\).

During late pregnancy and at parturition the brain-pituitary-ovarium (BPO) axis and the uterus are exposed to high circulating concentrations of progesterone, oestrogen, prostaglandins and oxytocin. After parturition, not only must the uterus recover, but also the BPO axis of the positive and negative endocrine feedback systems to enable resumption of the ovarian cycle\(^2\). Total serum progestagen concentrations did not differ between groups from parturition until the onset of foal heat. The relatively high total serum progestagen concentrations (5.2–16.4 ng/ml) found on the day of parturition declined rapidly to concentrations of 2–3 ng/ml within the next 48 h, which is in agreement with previous findings\(^1,3,4\). Although the effect of protein nutrition on progestagen concentration during the post partum period is uncertain, it appears that in the mares that only came into oestrus 15–19 d post partum the progestagen concentrations remained at between 1.5–3.0 ng/ml until 12–14 days post partum. The possible suppression of GnRH secretion by this persistently elevated level of serum progestagen is unknown at this stage.

The serum progestagen secretion patterns as described and shown in Fig. 2 during the oestrous cycle following foal heat are in agreement with those described previously\(^5\). The mean serum progestagen concentrations of mares that exhibited interovulatory periods of 17–19, 20–22 d, 24–25 d and 29–30 d (Fig. 2) indicate that, as the duration of the oestrous cycles increases, mid-cycle progestagen concentrations remain elevated for a longer period during the luteal phase of the cycle, and that maximum progestagen concentrations occur 2–3 d later during the oestrous cycle (Fig. 2). This might initially be the result of under-stimulation of the luteal tissue by LH, leading to lower production of progesterone. Early embryonic loss before Day 12 has no prolonged effect on the duration of the oestrous cycle\(^11\).

The 2 mares that ovulated 6–11 days post partum and developed persistent corpora lutea only came into oestrus between 40–80 days after the 1st ovulation. Both mares foaled during the transitional period (August–September), at the time of the year when the incidence of ovulation amongst barren mares is reported to be low\(^1\). A similar decline in ovarian activity after parturition as well as ovulation during foal heat in mares that foaled during winter or early spring has been described\(^3\). However, the results of this study indicate that the protein intake and quality had very little effect on progesterone production during the early post partum period.

No differences in the time of the LH peak secretion or the maximum concentrations were found between dietary Groups 1 and 4 during ovulation at foal heat or during the following oestrous period. It is accepted that the photoperiod might have influenced LH secretion, as has been described\(^1\). Current results are in agreement with previous reports of low LH concentrations at parturition\(^6\). LH concentrations are reported to increase gradually from Day 6 post partum until 2 days before ovulation, followed by a sharp increase to reach peak concentrations just after ovulation\(^10\). As shown in Fig. 4, the increase in LH coincides with low progestagen concentrations at 4–6 days post partum. Serum oestrogen concentrations that increase 5–10 days post partum\(^1\) stimulate secretion of LH. According to the results of the individual mares in this study, the LH peak was recorded on the day of ovulation or the day following ovulation, which is in agreement with current literature\(^1\).

Although no differences in the concentrations of FSH were found among mares in the different dietary groups, the observed decrease in FSH concentrations after parturition and the subsequent increase before oestrus and ovulation during foal heat have been described previously\(^1,3,4\). It is evident from the results of the individual mares that a relationship exists between the time when FSH concentration begins to decline and the time of ovulation. It therefore indicates that in cases where foal heat is delayed, the typical FSH secretion pattern does not occur at the same early stage. A similar stimulatory effect of high-quality protein nutrition on the production of progestagen is considered to meet the requirements of lactation.
dietary protein on FSH production found in barren mares’ was not observed in this group of mares during the post partum period. The overriding effect of season/photoperiod over protein quality of the diet might have been the reason for this finding.

The 10–14 day cyclical secretion pattern of FSH found in the lactating mares in this study (Fig. 5) is in agreement with previous reports. FSH stimulates follicular growth and a FSH peak normally occurs a few days after ovulation. Oestradiol produced by the developing follicles stimulates the LH surge and subsequently ovulation. In Equidae the mid-cycle FSH peak results in follicular development and the formation of mid-cycle follicles. These follicles generally undergo regression as LH concentrations, which increase slightly during this period but do not reach peak concentrations, are insufficient to stimulate follicular maturation and ovulation.

Only 56% of the mares that were not supplemented with fishmeal (Groups 1 and 2) conceived when mated during the oestrous period following foal heat, compared to the 100% conception rate in the mares supplemented (Groups 3 and 4). However, all mares mated in Groups 1 and 2 eventually conceived during subsequent oestrus periods.

**CONCLUSION**

Inclusion of high-quality protein in the diet of lactating mares resulted in an increased conception rate. The incidence of abnormally long or short oestrous cycles was reduced in mares receiving fishmeal supplementation. Serum progesterone, FSH and LH concentrations were unaffected by protein intake and quality. It would appear that the normal endocrine function of the brain-pituitary-ovarian axis is resumed within 6–18 days post partum with the possible exception of mares foaling during early spring. The effect of season/photoperiod influenced the post partum breeding results of mares that foaled during early spring detrimentally. Uterine involution may be delayed in mares fed low-quality protein diets resulting in a decrease in conception rate, an increase in the number of services per conception and, more importantly, in Thoroughbreds an increased possibility of lowered pregnancy rates at the end of the restricted breeding season.

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**Book review — Boekresensie**

**International aquatic animal health code (2nd edn)**

**Diagnostic manual for aquatic animal diseases (2nd edn)**

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The growing international trade in aquatic animals and aquatic animal products poses a significant risk of spreading diseases of importance in aquaculture. Promotion of animal health on a global scale is the mandate of the *Office International des Épizooties* (OIE). In recognition of the unique problems posed by aquaculture, a Fish Diseases Commission was established by the OIE to formulate guidelines for the control of aquatic animal diseases. These take the form of an *International Aquatic Animal Health Code* and accompanying *Diagnostic Manual for Aquatic Animal Diseases*.

The *International Aquatic Animal Health Code* is aimed at regulatory bodies involved in veterinary health certification and import control. It provides detailed information on both the principles and the practical application of certification, including excellent models of international health certificates for fish and shellfish. A section on import risk analysis is included as well as chapters on import and export procedures. Background information is given on all the internationally important diseases of finfish, shellfish and crustaceans. An appendix deals with destruction of pathogens by disinfection.

The companion *Diagnostic Manual for Aquatic Animal Diseases* concentrates on diagnostic procedures for notifiable and other significant diseases of finfish, shellfish and crustaceans. Background information on host range, clinical signs and epidemiology is given for all the diseases and some chapters include sections on control measures, vaccines and pathology when applicable. Chapters on sampling procedures and other practical aspects of health surveillance programmes are incorporated. Comprehensive references complete each section.

Both the *International Aquatic Animal Health Code* and *Diagnostic Manual for Aquatic Animal Diseases* are clearly written and contain a wealth of information on a wide range of topics related to aquatic animal disease control. Although unlikely to be of use to the general practitioner, anyone involved in health certification and import control of aquatic animals will find them invaluable.

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