Effect of conceptus removal and intrauterine administration of conceptus tissue on luteal function in the cow

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Summary. Two experiments were conducted to determine the day on which the bovine embryo first affects luteal function.

Experiment I was to determine the sole effect of flushing the uterine lumen and involved 12 non-inseminated heifers flushed 9, 14 and 16 days after estrus (4 animals per group). The same procedure was used to remove conceptuses from 15 pregnant heifers on days 9 (n = 4), 14 (n = 5) and 16 (n = 7) after the onset of estrus. Progesterone concentrations were measured daily throughout the cycle corresponding to conceptus removal (or flushing alone) and throughout the preceding cycle. In non-inseminated animals and when the conceptuses were removed on days 9 or 14, neither the progesterone pattern nor the inter-estrous interval was altered. By contrast, when the embryo was removed on day 16, the time of luteolysis was delayed by 4 to 7 days and the heifers were observed to be in estrus 26 to 29 days after AI.

In experiment 2, 21 virgin heifers were distributed over four treatment groups. Animals in groups 2 to 4 received the following treatments twice a day between day 15 and day 19 of the estrous cycle in the horn ipsilateral to the corpus luteum: group 2 (n = 3), 0.25 ml saline; group 3 (n = 3), two 12-day-old conceptuses; group 4 (n = 5), one 16-day-old conceptus. Progesterone concentrations were determined from day 13 of the cycle to 4 days after the following estrus. No antiluteolytic effect was found with intrauterine administration of either saline alone or day-12 embryos. In contrast, administration of day-16 conceptuses through the cervix (group 4) lengthened the estrous cycle by 6 to 7 days.

The results of these experiments suggest that the antiluteolytic and/or luteotropic factor(s) originating from the conceptus is fully potent by day 16. Embryonic mortality occurring on day 16 or later may induce an extension of corpus luteum lifespan.

Introduction.

Early pregnancy maintenance in the cow, as in other mammals, requires extended corpus luteum lifespan which is supported by luteotropic and/or antiluteolytic signals produced by the conceptus. In the ewe and the cow, evidence of luteotropic factors has been found both in both in vitro and in vivo

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experiments (Beal et al., 1981; Bulman and Lamming, 1978; Lukazewska and Hansel, 1980). Nevertheless, in these species, the mechanism by which the conceptus prevents luteolysis is not yet completely elucidated. In the ewe, PGE2 or other prostanoids (Henderson et al., 1977; Lewis et al., 1978; Pratt et al., 1979; Marcus, 1981) and a protein of trophoblastic origin (Martal et al., 1979) may be involved in corpus luteum maintenance. In bovines, Northey and French (1980) reported that the embryonic signal(s) may be given between days 15 and 17, but the chronological aspects of the interaction between the conceptus and ovary, as well as its consequences on corpus luteum function and estrous cycle length, were not completely determined.

The aims of the present study were to investigate the effects of (1) conceptus removal from the uterus at different stages following conception and of (2) transcervical intrauterine administration of frozen conceptuses of precise ages on corpus luteum function and (3) to determine the possible chronological relationship between conceptus death and the alteration of estrous cycle length.

**Materials and methods.**

*Animals and preliminary treatments.*

Sexually mature heifers (18 to 24 months old) of the French Friesian (FFPN) or FFPN × Limousine breeds, kept in a free stall area and fed essentially with maize silage, were used in this study. Before the beginning of the experiments, estrus was synchronized using a combination of a progesterone intravaginal device (Prid, Ceva France) left in place for 12 days and an intramuscular injection of 0.5 mg of prostaglandin F2α analog (Cloprostenol: Uniardine, ICI France) administered two days before PRID removal. After the synchronization of estrus and throughout the subsequent estrous cycles heifers were routinely checked for estrus twice a day (from 7 to 9 a.m. and 6 to 8 p.m.). Only heifers exhibiting standing estrus were used in these experiments.

**Experiment 1: Effect of the flushing procedure and of conceptus removal on luteal function.**

Fifty-two heifers were distributed into two treatment groups. The animals in group 1 were used to study the effect of flushing the uterus on the length of the estrous cycle. In this group 12 heifers which had not been previously inseminated were flushed 9, 14 and 16 days (4 animals per day) after the onset of estrus (fig. 1) by the transcervical procedure initially described by Ozil et al. (1979). The catheter was introduced through the cervix into the uterine horn ipsilateral to the corpus luteum. A total volume of 300 ml of phosphate-buffered saline (PBS) was injected in small fractions (30 to 50 ml) and collected.

In group 2 forty heifers were inseminated and the uterus was subsequently flushed by the same procedure on days 9 (n = 18), 14 (n = 6) and 16 (n = 16) after AI. The presence of a conceptus was determined after collection of the flushing and, in these females, 4 day — 9, 5 day — 14 and 7 day — 16 conceptuses were found. All heifers were subjected to AI and uterine flushings
after the second estrus following synchronization of estrus. All animals were inseminated twice with the semen of one bull. The first AI was performed 12 hours after the beginning of estrus and the second AI 12 hours later.

For heifers flushed on day 9 (inseminated or not), plasma samples were obtained daily for progesterone determination throughout two consecutive cycles from the end of the estrus synchronization regimen to day 5 after the estrus following the flushing procedure (fig. 1). Heifers flushed on days 14 and 16 (inseminated or not) were bled during two consecutive cycles, i.e. between days 0 and 5 and between days 12 and 14, respectively, to 5 days after the following estrus (fig. 1).

Experiment 2: Intrauterine administration of conceptus tissue.

The conceptuses used in this experiment (thirty 12-day-old conceptuses and approximately, fifty 16-day-old conceptuses) were obtained from 30 superovulated cows either by the transcervical procedure described previously or after collection of the genital tracts at the slaughterhouse. The embryos were sorted according to morphological criteria. All the conceptuses or pieces of conceptus which were degenerated were discarded. After collection, the conceptuses were immediately sucked into French straws (0.25 ml) and stored in liquid nitrogen (−196 °C) until used.
The treated heifers were divided into 4 groups. Group 1 (n = 10) received no intrauterine treatment; in groups 2 to 4 the heifers were injected in the horn ipsilateral to the corpus luteum twice a day (9 a.m. and 6 p.m.) between days 15 and 19 with either 0.25 ml of saline solution (group 2, n = 3), two 12-day-old conceptuses in 0.25 ml of saline solution (group 3, n = 3), or 0.25 ml of 16-day-old conceptus tissue (group 4, n = 5), respectively.

The saline solution and day-12 conceptuses were introduced with a 4 mm Ø transfer catheter (Rusch RFA) with a subterminal opening. Day-16 conceptuses were introduced with a 4 mm Ø inseminating catheter (IMV France) with an axial opening.

The treatment solutions were administered intrauterinely after epidural anesthesia with 4 ml of a solution of propoxycaine (Silvocaine N.D.).

Blood samples were obtained daily from each heifer for progesterone determination from day 13 after the onset of the first estrus following the synchronization of estrus until day 5 after the onset of the second estrus.

Blood sampling and progesterone radioimmunoassay. — Peripheral blood (15 ml) was collected from the jugular vein into heparinized vacutainers (Beckton Dickinson, France). Plasma was immediately separated by centrifugation and stored at —20 °C until progesterone assay.

An assay procedure previously described and validated by Thibier and Saumande (1975) was used to analyse single aliquots of plasma. Antiserum 111/6 (Specific antisera LTD 1981, UK) raised in goat against progesterone — 11 α succinyl BSA was used at a final dilution of 1/10 000. This antiserum displayed more than 12 % cross-reactivity with 17 α-hydroxyprogesterone. The two steroids were separated by sephadex LH 20 microcolumn chromatography. The accuracy and efficiency of this separation procedure were satisfactory as reported previously (Thibier et al., 1973). The sensitivity of this assay was estimated to be 0.05 ng/ml. The intraassay and interassay coefficients of variation were 15.4 and 17.3 %, respectively (10 determinations).

Analysis of data. — In both experiments, the length of the estrous cycle was estimated by the intervals between successive estruses (day 0 was designated as the first day of standing estrus). Estrus was further confirmed by complete luteolysis (i.e. progesterone concentrations lower than 1 ng/ml). In experiment 1 these data were collected over two consecutive cycles. Each heifer was its own control thus minimizing the effect of individual variability. It has been shown that the length of the estrous cycle and the area under the progesterone curve in successive cycles are reproducible (Yenikoye et al., 1981, Humblot et al., 1981).

Results

Effect of the uterine flushing procedure. — In noninseminated animals (group 1), when uterine flushing was performed on either day 9, 14 or 16 of the

FIG. 2. — Effect of uterine flushing on days 9 (a, n = 4), 14 (b, n = 4) and 16 (c, n = 4) of the estrous cycle on plasma progesterone concentrations (X ± S.E.). Day of estrus, Day of uterine flushing; Control cycle, Cycle during which uterine flushing was performed.
estrous cycle, the progesterone patterns and concentrations as well as
interestrous* intervals (fig. 2 ; table 1) were very similar to those observed
throughout the control estrous cycle. Consequently, no effect of the flushing
procedure alone was found on luteal function at these three stages of the estrous
cycle.

Effect of conceptus removal. — When the conceptuses were removed from
the uterus either 9 or 14 days after conception, estrous cycle length was not
significantly different from that of the control cycle (table 1), indicating no
extension of corpus luteum lifespan (fig. 3 a and b).

By contrast, when embryos were removed 16 days after conception,
luteolysis and the following estrus occurred 7 days later than expected (fig. 3 c ;
table 1), Six heifers were then observed to be in heat 26, 26, 26, 27, 29 and 29
days after Al. The mean progesterone concentrations observed between days 20
and 23 after Al were higher (p < 0.001) than those observed during the same
period of the control cycle (fig. 3 c).

As shown by the individual progesterone profiles (fig. 4) of these six heifers,
complete luteolysis occurred from 5 to 9 days later than in the control cycle. After
conceptus removal, progesterone concentrations remained elevated and
subsequently sharply declined to levels lower than 1 ng/ml within 24 hours. One
additional heifer, from which a day-16 conceptus was removed, was not observed
in estrus and failed to undergo complete luteolysis during the whole sampling
period (fig. 4).

The length of day-16 conceptuses recovered ranged from 5 to 11 cm (mean
= 7.8 ± 1.9 cm). No evidence of a relationship between the size of the
conceptus and the delay of luteolysis was detected (r = 0.08).

**TABLE 1**

<table>
<thead>
<tr>
<th>Day</th>
<th>Treatment</th>
<th>N</th>
<th>Control cycle (mean ± S. D.)</th>
<th>U.F. or E.R. cycle (mean ± S. D.)</th>
<th>S.</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>U.F.</td>
<td>4</td>
<td>21.7 ± 0.5</td>
<td>21.0 ± 2.0</td>
<td>N.S.</td>
</tr>
<tr>
<td>14</td>
<td>U.F.</td>
<td>4</td>
<td>20.3 ± 0.6</td>
<td>21.6 ± 0.6</td>
<td>N.S.</td>
</tr>
<tr>
<td>16</td>
<td>U.F.</td>
<td>4</td>
<td>21.7 ± 0.5</td>
<td>22.2 ± 0.5</td>
<td>N.S.</td>
</tr>
<tr>
<td>9</td>
<td>E.R.</td>
<td>4</td>
<td>22.2 ± 1.5</td>
<td>21.2 ± 0.9</td>
<td>N.S.</td>
</tr>
<tr>
<td>14</td>
<td>E.R.</td>
<td>5</td>
<td>21.8 ± 0.8</td>
<td>21.0 ± 0.7</td>
<td>N.S.</td>
</tr>
<tr>
<td>16</td>
<td>E.R. (*)</td>
<td>6</td>
<td>21.2 ± 0.4</td>
<td>27.2 ± 1.5</td>
<td>p &lt; 0.001</td>
</tr>
</tbody>
</table>

(*) A conceptus was removed from one additional heifer, but complete luteolysis and estrus were
not observed during the whole sampling period (35 days).

FIG. 3. — Effect of conceptus removal on days 9 (a, n = 4), 14 (b, n = 5) and 16
(c, n = 6) after onset of estrus on plasma progesterone concentrations (X ± S.E.). ♦ Day
of estrus, ◻ Day of conceptus removal, O Control cycle, ● Cycle during which
the conceptus was removed. (*) Significant difference (p < 0.001) detected by paired t-test.
FIG. 4. — Individual progesterone patterns of 7 heifers from which day-16 conceptuses were removed. O—O Progesterone concentrations measured throughout the control cycle, ••••• Progesterone concentrations measured throughout the cycle during which the conceptus was removed.
Effect of intrauterine administration of conceptus tissue. — It was found by analysis of variance that mean estrous cycle length differed (p < 0.001) between the four treatment groups (table 2). Individual differences between groups were further investigated by a multiple comparison analysis.

**TABLE 2**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>N</th>
<th>Interestrus Interval (mean ± S.D.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls</td>
<td>10</td>
<td>22.3 ± 1.0 (1)</td>
</tr>
<tr>
<td>Saline</td>
<td>3</td>
<td>19.0 ± 1.0 (2)</td>
</tr>
<tr>
<td>Conceptuses (day 12)</td>
<td>3</td>
<td>20.0 ± 1.0 (3)</td>
</tr>
<tr>
<td>Conceptuses (day 16)</td>
<td>5</td>
<td>27.0 ± 1.0 (4)</td>
</tr>
</tbody>
</table>

N = Number of heifers.
2 vs 1 and 4 vs 1, 2 or 3, significant difference p < 0.001.
3 vs 1, significant difference p < 0.01.
3 vs 2, N. S.

**FIG. 5.** — Progesterone profiles (X̄) of control heifers ⋄ (n = 10) and after repeated transcervical administration ‡ of: saline solution ▲ (n = 3), day-12 conceptuses Δ-----Δ (n = 3) and day-16 conceptuses □-----□ (n = 5).

Control animals, which received no intrauterine treatment, underwent complete luteolysis a mean 21 days after the preceding estrus. Luteolysis occurred sooner (fig. 5) in heifers treated either with saline solution alone \( (p < 0.001) \) or with day-12 conceptuses \( (p < 0.01) \).

By contrast, when day-16 conceptuses were administered, luteolysis occurred 5 to 8 days later than in the other three groups. Interestrous intervals for these five heifers were 26, 26, 27, 28 and 28 days (table 2). These data demonstrate the antiluteolytic action of day-16 conceptuses administered transcervically.

No significant rise in plasma progesterone concentrations was observed after the introduction of day-16 conceptuses: the levels were very similar to those observed during a normal luteal phase before luteolysis.

Discussion

Effect of the uterine flushing procedure and conceptus removal on luteal function.

In contrast to data reported by Elsden et al., (1976) and Crister et al., (1980), no effect of the uterine flushing procedure on luteal function was found at any of the three stages of the estrous cycle studied. However, Crister et al., (1980) performed their studies during the early luteal phase (between days 5 and 8). These divergent results suggest that the sensitivity of the CL to uterine handling may be different according to the stage of the estrous cycle. The fact that, in the present study, uterine flushing did not interfere with luteal function demonstrates that this method is suitable for mimicking embryonic death, at least at these early stages of conceptus development. The situation may be different during later stages of pregnancy. It was reported in the ewe (Dooley et al., 1974) that estrus and a decline in progesterone concentrations occurred earlier after embryonic mortality simulated by uterine flushing than that induced by a colchicine injection.

The results obtained after embryo removal in our study fully confirm those obtained by Northey and French (1980) and show clearly that 9 and 14-day conceptuses do not produce an antiluteolytic signal or, at least, produce it in quantities too small to extend corpus luteum lifespan.

By contrast, after removal of day — 16 conceptuses, luteolysis was delayed by 5 to 7 days in all animals. This delay was longer than that reported by Northey and French after removal of older conceptuses (17 or 19 days). The better expression of the antiluteolytic action of the conceptuses found here may be due to the fact that each animal was its own control, thus minimizing the effect of individual variability. When conceptuses were removed 16 days after AI, progesterone concentrations between days 20 and 24 were slightly lower than those detected in pregnant animals at the same stage (data not shown), but remained higher than 2 ng/ml up to 2 days before complete luteolysis. This typical pattern of change in progesterone concentrations in these heifers may reflect the lack of a conceptus factor which stimulates progesterone production by the corpus luteum, as suggested by Lukaszewska and Hansel (1980).
Moreover, from a practical viewpoint, embryonic death subsequent to day 15 would decreased the accuracy of early pregnancy diagnosis based on progesterone concentrations 21-24 days after breeding. Nevertheless, the incidence of such cases on cow fertility would be relatively low because the frequency of late embryonic mortality does not exceed 10% of the total number of Al's (Kummerfeld et al., 1978; Ball, 1978; Humblot, 1981).

Effect of intrauterine administration of conceptus tissue on luteal function.

Repeated intrauterine administration of saline solution decreased corpus luteum lifespan by 1 or 2 days. This trend is in agreement with previous findings of Bartol et al., 1981a, who reported that repeated uterine manipulations and flushings increase prostaglandin F2α release. This may induce early luteolysis, especially if these treatments are performed at the end of the cycle when prostaglandin F2α receptor number and affinity in the corpus luteum are maximal (Bartol et al., 1981b).

After administration of 16-day conceptuses, luteolysis occurred a mean 26 days after the previous estrus. This time interval is 2 days longer than that reported by Northey and French (1980) who worked with a mixture of day-17 and day-18 conceptuses introduced into the uterine lumen by a surgical procedure between days 14 and 18 of the cycle. If such a difference could be considered significant, it would suggest that either day-16 conceptuses have greater antiluteolytic activity than day-17 or 18 conceptuses or that the transcervical administration of conceptuses interferes less than the surgical procedure with luteal function. The day-15 to day-19 period during which the embryos were administered might have affected luteolysis more than the day-14 to day-18 period. It may also be speculated that deep freezing was less detrimental to the antiluteolytic substance released by the conceptus than the homogenization procedure employed by Northey and French (1980).

The chemical nature of the signal(s) produced by the bovine conceptus remains unknown. PGE₂ or similar compounds have been suggested as candidates for this signal, but they need to be administered chronically to extend corpus luteum function (Magness et al., 1981; Huie et al., 1981; Gimenez and Henricks, 1983). Moreover, it was found from a preliminary experiment (Dalla Porta and Humblot, 1983) that PGE₂, administered by the same way as these conceptuses, was less potent than day-16 conceptuses in maintaining progesterone production by the corpus luteum. These observations suggest that other signals, such a proteins of trophoblastic origin (Martal et al., 1979; Heyman et al., 1982), may be necessary at the beginning of pregnancy. These proteins could also be involved in PGE₂ or prostacyclin production or transfer (Marcus, 1981) from the uterus to the ovary.

In conclusion, it has been shown that the antiluteolytic and/or luteotropic factor(s) originating from the bovine conceptus is fully potent on day 16 of gestation.

Acknowledgements. — The authors gratefully acknowledge the excellent technical assistance of J.-L. Schwartz.

Deux expériences ont été réalisées pour déterminer le jour auquel l’embryon commence à avoir un effet sur la fonction lutéale.

Dans l’expérience 1, 12 génisses non inséminées ont été collectées 9, 14 et 16 jours après les chaleurs (4 par groupe). La même méthode a été utilisée pour prélèver le conceptus chez 15 génisses gestantes 9 (n = 4), 14 (n = 5) et 16 jours (n = 7) après la fécondation. Les concentrations de progestérone ont été mesurées journalement pendant le cycle correspondant au retrait du conceptus (ou au lavage utérin seul) et pendant le cycle précédent. Chez les génisses non inséminées et quand les embryons ont été retirés 9 ou 14 jours après la fécondation la longueur du cycle sexuel comme les profils de progestérone n’ont pas été modifiés. En revanche, quand l’embryon a été retiré 16 jours après la fécondation, la lutéolyse est différée de 4 à 7 jours.

Dans l’expérience 2, 22 génisses ont été réparties en 4 groupes. Les génisses du groupe 1 n’ont reçu aucun traitement intra-utérin. Les animaux des groupes 2 et 4 ont reçu 2 fois par jour entre J_{16} et J_{19} du cycle dans la corne ipsilatérale au corps jaune les traitements suivants : groupe 2 (n = 3) 0,25 ml le sérum physiologique, groupe 3 (n = 3) deux embryons de J_{12} dans 0,25 ml de sérum physiologique, groupe 4 (n = 5) un embryon de J_{16}. Les concentrations de progestérone ont été déterminées à partir de J_{13} jusqu’au 4e jour après l’oestrus suivant. Aucun effet antilutéolytique n’a été trouvé après administration intra utérine d’embryons âgés de 12 jours. Par contre l’administration d’embryons de J_{16} a allongé la durée du cycle de 6 à 7 jours.

Les résultats de ces expériences suggèrent que le ou les facteur(s) lutéotrophiques ou antilutéolytiques provenant de l’embryon est (sont) actif (s) dès le 16e jour après la fécondation. La mortalité embryonnaire survenant à 16 jours ou plus tard provoque une extension de la durée de vie du corps jaune.

Références


