Uterine secretion of prostaglandin F2α stimulated by different doses of oxytocin and released spontaneously during luteolysis in cattle

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Abstract – The objectives were to determine the involvement of oxytocin (OT) in the stimulation of prostaglandin F2α (PGF) secretion during luteolysis in cattle. On days 16–17 of the oestrous cycle, catheters were inserted into the aorta abdominalis of heifers for OT or saline infusion and into the jugular vein for blood sample collection. The following day, heifers were assigned to one of three experimental groups (Gr): Gr I – 10 IU OT (n = 4); Gr II – 20 IU OT (n = 4); Gr III – 50 IU OT (n = 4). Blood samples were collected every 10 min during a 1-h control period before treatment and every 5–10 min for 2 h after OT treatment. In Gr IV (n = 5), a catheter was inserted into the jugular vein on day 15 of the cycle and blood samples were collected every 15 min for 12 h on days 16–19. Plasma concentrations of progesterone, PGF metabolite, 13, 14-dihydro-15-keto-prostaglandin F2α (PGFM) and OT were determined. Within 5 min of infusion of 10 or 20 IU OT, peripheral concentrations of OT (7–12 pg/mL) increased by about 200 and 350–500 pg/mL, respectively. These doses did not affect plasma concentrations of PGFM or progesterone within 1.5 h. Fifty IU of OT increased its maximal peripheral concentration to 1 500 pg/mL, which is over 20 times greater than that observed physiologically. Concentrations of plasma PGFM in Gr III increased from basal concentrations (50–65 pg/mL) to 150–250 pg/mL (P < 0.01) within 10–30 min. During luteolysis, PGFM pulses ranged between 250 and 600 pg/mL on days 16–19 of the cycle (Gr IV), whereas coincident pulses of OT, and those appearing between spikes of PGFM, were never above 75 pg/mL. Only 50 % of OT pulses coincided with pulses of PGFM, and 54 % of PGFM pulses coincided with a pulse of OT. Results indicate that luteolytic PGF secretion in cattle is not directly dependent upon ovarian OT. © Inra/Elsevier, Paris

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Résumé – Sécrétion utérine de prostaglandine F2-α stimulée par différentes doses d’ocytocine, et libérée spontanément durant la lutéolyse chez les bovins. Les objectifs étaient de déterminer le rôle de l’ocytocine (OT) sur la régulation de la sécrétion de prostaglandine F2-α (PGF) pendant la lutéolyse chez les bovins. Aux jours 16 ou 17 du cycle oestrus, des cathéters étaient insérés dans l’aorte abdominale de génisses pour infuser de l’OT ou du sérum physiologique, et dans la veine jugulaire pour collecter des échantillons de sang. Le jour suivant, les génisses étaient réparties au hasard dans un des trois groupes (Gr) expérimentaux : Gr I - 10 UI OT (n = 4) ; Gr II - 20 UI OT (n = 4) ; Gr III - 50 UI OT (n = 4). Des échantillons de sang étaient collectés toutes les 10 min pendant une période témoin de 1 h, et toutes les 5–10 min pendant 2 h après le traitement OT. Dans le Gr IV (n = 5), un cathéter était placé dans la veine jugulaire le quinzième jour du cycle et des échantillons de sang étaient collectés toutes les 15 min pendant 12 h aux jours 16, 17, 18 et 19 du cycle. Les concentrations plasmatiques de progestérone, de 13, 14-dihydro-15-citoprostaglandine F2-α (PGFM) et d’OT ont été mesurées. L’infusion de 10 ou 20 UI d’OT augmente les concentrations périphériques d’OT qui atteignent environ 200 et 350–500 pg/mL en 5 min. Ces doses n’affectent pas en revanche les concentrations plasmatiques de PGFM ou de progestérone. Une dose plus élevée d’OT (50 UI) porte la concentration maximale de 1 500 pg/mL, ce qui est plus de 20 fois supérieur à celle observée physiologiquement. Dans le Gr III, les concentrations plasmatiques basales (50–65 pg/mL) de PGFM augmentent jusqu’à 150–250 pg/mL (P < 0,01) en 10–30 min. Durant la lutéolyse, des pulses de PGFM atteignant des niveaux de 250–600 pg/mL étaient détectés entre j 16 et j 19 du cycle (Gr IV), tandis que les pulses d’OT coïncidaient, et ceux apparaissant entre les pics de PGFM, n’étaient jamais supérieurs à 75 pg/mL. Seulement 50 % des pulses d’OT coïncidaient avec des pulses de PGFM, et 54 % des pulses de PGFM coïncidaient avec un pulse d’OT. Les résultats indiquent que la sécrétion lutéolytique de PGF chez les bovins ne dépend pas directement de l’OT ovarienne. © Inra/Elsevier, Paris

ocytocine / progestérone / lutéolyse / prostaglandine F2-α / corps jaune / bovins

1. INTRODUCTION

Hormonal processes associated with the initiation of luteolysis in cattle are not completely understood [34, 39]. Although uterine prostaglandin F2-α, (PGF) is accepted as the natural luteolysin in cattle [14], the trigger for its secretion remains unknown. It has been suggested that ovarian OT initiates PGF secretion [7], and that this effect depends on the exposure of the endometrium to physiological concentrations of progesterone during dioestrus [8, 17]. In cattle the exposure to progesterone, which lasts for 11–14 days depending upon the length of the oestrous cycle [18], is crucial for uterine synthesis of OT receptors [31]. However, a negative correlation has been demonstrated between concentrations of circulating progesterone and concentrations of uterine oxytocin receptor and hence, the degree of responsiveness to oxytocin [16]. These data suggest progesterone concentrations should be decreased in order for the uterus to become sensitive to an OT challenge. This is in agreement with data suggesting that ovarian OT may not be essential for initiation of luteolysis in cattle, since depletion of luteal OT by 68–82 % with noradrenaline on days 12 or 16 of the cycle [19] did not affect luteolysis. In addition, the complete blockade of OT receptors with a specific OT antagonist from day 15 until day 22 of the cycle [22], did not affect luteolysis. The aim of this study was to investigate the effect of increasing doses of exogenous OT on PGF secretion on day 17 of the oestrous cycle (the time of luteolysis), and to compare peripheral concentrations of OT produced by these treatments to those resulting from endogenous secretion of this peptide on days 16–19. The PGF metabolite, 13, 14-dihydro-15-
keto-prostaglandin F2α (PGFM) was used as an indicator of PGF secretion in this study.

2. MATERIALS AND METHODS

2.1. Animals and treatment

The experiments were carried out in accordance with the principles for the care and use of research animals. Mature nulliparous heifers with a palpable corpus luteum (CL) were injected i.m. with 500 μg of a PGF analogue, Oestrophan (Spofa) to induce luteolysis. On day 17 after signs of oestrus were observed, catheters were inserted into the aorta abdominale [20] of each heifer for OT or saline infusion, and into the jugular vein for blood sample collection. The following day, heifers were assigned randomly to one of four experimental groups (Gr): Gr I - 10 IU OT (n = 4); Gr II - 20 IU OT (n = 4); Gr III - 50 IU OT (n = 4); Gr IV - control. Blood samples were collected every 10 min for 1 h before infusion of OT, followed by collection every 5 min for 30 min, and every 10 min for a further 1 h after OT infusion. Oxytocin (Richter, Hungary) was infused in 5 mL of saline, followed by 5 mL of saline to flush the cannula.

In Gr IV (n = 5) jugular blood samples were collected every 15 min for 12 h, from 0800 to 2000 hours, on days 16–19 of the oestrous cycle to characterize PGF and OT secretion during spontaneous luteolysis. Clotting of blood samples (8 mL) was prevented by the addition of 76-gM EDTA and 11-pM acetylsalicylic acid.

2.2. Radioimmunoassays

Concentrations of PGFM were determined as described by Homanics and Silvia [11] and the antiserum (WS4468 BD 6/23-7/21) was characterized by the same authors. Sensitivity of the assay was 12 pg/tube. Plasma concentrations of progesterone were determined by a method validated in this laboratory [20] using a rabbit progesterone antiserum (IFP₄) characterized previously [21]. The sensitivity of the procedure was 15 pg/tube (0.3 ng/mL of plasma). Intra- and inter-assay C.V. were 8.1 and 15.9 %, respectively. The relationship between real (x) and determined (y) amounts of four different concentrations of progesterone added to plasma samples is expressed by the linear regression equation (y = 1.034 x – 0.13). Oxytocin was extracted from plasma using a 3-fold excess of ice-cold acetone. Rabbit OT antiserum (R-1) was characterized previously [19]. The efficiency of extraction was 85 %, and final data were corrected for procedural losses. The sensitivity of the method was 1.3 pg/tube (3 pg/mL). Intra- and inter-assay C.V. were 7.5 and 14.6 %, respectively. Precision of the procedure based on four different concentrations of added mass is expressed by the linear regression equation (y = 0.99x + 0.14). Estradiol-17β was extracted from 1 mL of plasma using 5 mL of diethyl ether and RIA was performed using rabbit antiserum (BSz/88/754) characterized by Szafranska and Tilton [40]. Efficiency of extraction was above 80 % and data were corrected for the procedural losses. Sensitivity of the method was 2 pg/mL and the intra- and inter-assay C.V. were 5.5 and 11.2 %, respectively.

2.3. Data analysis

Magnitude of a PGFM response after OT infusion was defined as the maximum concentration of metabolite above baseline during the 1.5-h post-injection period. Baseline was defined as the average concentration of PGFM before injection of oxytocin. Pulses of PGFM and OT in heifers from Gr IV were identified by means of the Pulsar program of Merriam and Wachter [27] as modified for the personal computer by Gitzen and Ramirez [9]. Quadratic, linear and constant terms for calculating assay SD in Pulsar and G(n) values for defining pulses were the same as given earlier [38]. The relationship between OT and PGFM pulses were determined by calculating the number of pulses that appeared either concomitantly or within 1 h of the start of a pulse of another hormone.
3. RESULTS

Infusion of 10 or 20 IU of OT increased peripheral concentrations of this peptide to about 200 and 350–500 pg/mL, respectively, within 5 min of infusion; however, neither dose of OT affected PGFM concentrations (figure 1). When 50 IU of OT were given, an increase of OT to 1 500 pg/mL in peripheral circulation was observed. This increase in OT caused PGFM concentrations to increase ($P < 0.01$) from basal levels (35–65 pg/mL) to 100–150 pg/mL within 10–30 min (figure 1). Concentrations of progesterone during the experiment fluctuated and ranged from 2 to 6 ng/mL in heifers in groups I–III. Amplitude of the PGFM peak after OT treatment was independent ($P > 0.05$) of progesterone concentration in individual heifers.

![Figure 1. Influence of bolus infusion of 10 (n = 4), 20 (n = 4), and 50 IU (n = 4) of oxytocin (OT) into the aorta abdominalis on peripheral plasma concentrations of PGFM (mean ± s.e.m.) on day 17 of the oestrous cycle in heifers. PGFM concentrations after 50 IU of OT are given individually for each of four heifers due to varying temporal responses. Progesterone concentrations in these heifers fluctuated and ranged from 6 to 2 ng/mL.](image-url)
During spontaneous luteolysis in heifers of Gr IV, pulses of PGFM ranged between 250 and 600 pg/mL, depending upon day of the cycle. Coincident pulses of OT, and those which appeared between peaks of PGFM on days 16–19 of the cycle, never exceeded 75 pg/mL (figures 2 and 3). When jugular blood samples were collected at frequent intervals on days 16–19 of the cycle, 48% of all PGFM pulses occurred within the sampling interval of a pulse of OT, and 73% of OT pulses were coincident with pulses of PGFM. Thus, many pulses of one hormone were in the absence of the other. The total number of OT and PGFM pulses in the heifers studied was 15 and 23, respectively. Moreover, it should be noted that four PGFM pulses followed the OT peak and four OT pulses were observed after PGFM surges.

4. DISCUSSION

To significantly stimulate PGF secretion (as determined by analysis of PGFM) by means of exogenous OT, it was necessary to increase peripheral OT concentrations to about 1 500 pg/mL as shown in Gr III, and to even greater concentrations in a study reported by Howard and Britt [13]. This is over 20 times greater than physiological levels of this peptide in cattle ([22, 36]; current study). When OT plasma concentrations exceeded 4–8 times normal values (figure 1) after treatment with 10 or 20 IU, a PGFM response was not observed.

It has been suggested that uterine PGF secretion is switched on by ovarian OT [7, 39]. However, as long as plasma progesterone concentrations are high, OT receptors may not be replenished [16, 28] and in fact, specific binding sites for OT in the bovine corpus luteum are lowest on days 15–18 of the oestrous cycle [32]. Furthermore, analysis of OT and OT mRNA contents of bovine corpus luteum show that little or no OT or OT mRNA is present during luteolysis [4, 15, 24]. Therefore the larger amount of ovarian OT present in the early and mid-luteal phases may be rather luteotrophic in cattle, as found using a microdialysis system [29]. Thus, it can be assumed that an increase in uterine OT receptor numbers should be preceded by a decrease in progesterone concentrations and this rather precludes a controlling influence of luteal OT on the initiation of luteolysis.

On days 13–16 of the oestrous cycle in ewes, 96% of PGFM pulses were observed concomitantly with OT pulses, whereas only 56% of oxytocin pulses coincided with pulses of PGFM if utero-ovarian vein blood was collected [12]. These authors concluded that OT stimulates uterine PGF secretion and not vice versa. A similar relationship between OT and PGF was found in ewes by Burgess et al. [2]. However, this conclusion omits the fact that over 44% of OT pulses from the ovary do not produce a pulse of PGF from the uterus. Furthermore other groups working with sheep have found opposite results. Moore et al. [30] observed that concentrations of PGFM in the utero-ovarian vein effluent increased before any increase was observed in secretion of OT during frequent bleeding. This observation agrees with data by Silvia and Raw [38] who found that frequency of PGFM pulses in ovariectomized ewes could be completely restored without OT support. However, the lower PGFM pulses observed in their studies may suggest that OT amplifies the course of induced luteolysis as a regulator of the amplitude of pulsatile PGF secretion, as shown in heifers [23].

In the present study, 27 and 52% of pulses of OT and PGFM, respectively, were released without concomitant secretion of the other hormone during 12-h periods in Gr IV (figures 2 and 3). Very similar data were obtained in recent studies.
Figure 2. Plasma concentrations of progesterone, oxytocin, and PGFM in heifer #1 and 2 on days 16–19 of the oestrous cycle. Pulsatile secretion of PGF (+) and oxytocin (*) was observed on days 18–19 of the cycle. Blood samples were taken from the jugular vein every 15 min for 12 h.
Figure 3. Plasma concentrations of progesterone, oxytocin, and PGFM in heifer #3 and 4 on days 16–19 of the oestrous cycle. Pulsatile secretion of PGF (+) and oxytocin (*) was observed on days 18–19 of the cycle. Blood samples were taken from the jugular vein every 15 min for 12 h.
Observations by Fields et al. [5] also seem to agree with these results. They showed that from days 7–14 there was a 69% decline in the number of large cells containing oxytocin-laden secretory granules in the cow. This occurred before the synthesis of uterine OT receptors and luteolytic pulses of PGF. These results help to explain why depletion of OT from the bovine CL by up to 82% on days 12 or 16 of the oestrous cycle did not affect the time of luteolysis [19]. It can be assumed that even less than 20% of OT may be enough to induce luteolysis. However, to prove that OT affects PGF secretion it was necessary to treat cow with 20–100 IU of OT [13, 25]. Moreover, continuous blockade of OT receptors in heifers from day 15 until oestrus did not affect luteolysis compared to control [22]. This supports recent data [1, 10] which demonstrated that luteal regression can occur in the absence of OT release from the luteal cells.

Collectively, these and other data seem to disagree with the theory that ovarian OT can trigger uterine PGF secretion and initiate luteolysis in cattle. Previously, it was observed that continuous infusion of OT in sheep blocked the synthesis of OT receptors [6]. However, injection of a PGF analogue during OT infusion caused a rise in OT receptors to values comparable to those in control animals. Thus, it is possible that during spontaneous luteolysis, PGF either decreases progesterone concentrations, allowing OT receptors to increase, or it stimulates OT receptor synthesis directly as suggested by Chan et al. [3] and by Sheldrick and Flick-Smith [37].

Data by Homanics and Silvia [11] and Lamming et al. [26] indicate that progesterone pre-treatment of sheep for over 10 days stimulates PGF synthesis and secretion. Furthermore, they suggest that the length of progesterone exposure controls the timing for the down-regulation of uterine progesterone receptors which, in turn, controls OT receptors. Similarly, studies in cattle [8, 17] have shown that duration of exposure to progesterone at concentrations above 2 ng/mL plays an important role in controlling the appearance of PGF pulses. Usually, the period of progesterone dominance in cattle lasts 11–14 days, depending upon oestrous cycle duration. Parkinson and Lamming [33] have observed pulses of PGF in cattle, theoretically of luteal origin [35], 2–3 days before the drop in progesterone concentrations in blood at luteolysis. Thus, it can be hypothesized that the influence of progesterone in controlling the secretion of low level PGF pulses occurs well-before the time of luteolysis, as assumed earlier [3, 11].

In conclusion, results of the current study indicate that the initiation of luteolytic PGF secretion in cattle is not directly dependent upon ovarian OT, as has been suggested in sheep. However, these observations do not exclude a possible involvement of ovarian OT in a luteolytic process already underway. Influence of OT on amplitude of PGF pulses [23] may be such an example. Thus, OT may be only one of a mixture of physiological factors involved in the start of functional luteolysis.

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