Stimulation of ovarian oxytocin secretion and uterine prostaglandin release by exogenous progesterone early in the cycle of the ovarian auto-transplanted ewe

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The present study was undertaken to determine whether the administration of progesterone, early in the oestrous cycle, had an influence on ovarian oxytocin secretion and on peripheral concentrations of the prostaglandin F₂α metabolite 13,14-dihydro-15-keto PGF₂α (PGFM) in the ovarian auto-transplanted ewe. Twelve ewes with ovarian auto-transplants (n = 6 per group) were randomly assigned to receive an i.m. injection of progesterone (12.5 mg) or vehicle, twice a day, on days 1, 2 and 3 of the oestrous cycle. Beginning on day 7, blood samples were collected at intervals of 1 h from the ovarian and contralateral jugular veins for up to 70 h. Ovarian oxytocin secretion rate and jugular concentrations of PGFM and progesterone were determined by radioimmunoassay. The number of ewes that showed pulses of both ovarian oxytocin and PGFM was significantly (P < 0.05) greater in progesterone-treated ewes than in control ewes. In progesterone-treated ewes, the average number of ovarian oxytocin pulses per ewe was 9.66 ± 5.5 (mean ± SD) and the interval between pulses was 7.18 ± 5.8 h. The mean amplitude and amount of oxytocin released, as calculated by the area under the curve of ovarian oxytocin pulses, were 6.27 ± 1.98 ng min⁻¹ and (10.05 ± 8.91 ng min⁻¹)τ, respectively (where τ is the number of hours between the last time point before and the first time point after a significant increase in hormone concentration was detected by the Pulsar program). The mean amplitude and area under the curve of PGFM pulses were 317.22 ± 5.65 pg ml⁻¹ and (383.36 ± 1.77 pg ml⁻¹)τ, respectively. The average number of pulses of plasma PGFM observed per ewe was 5.8 ± 1.9 and interpulse interval for plasma PGFM pulses was 10.32 ± 8.7 h between day 7 and day 9 after oestrus. These data indicate that administration of progesterone during the first 3 days of the oestrous cycle results in the premature release of ovarian oxytocin and uterine prostaglandin F₂α.

Introduction

Administration of progesterone early in the oestrous cycle of ewes (Ginther, 1968, 1969; Ottobre et al., 1980; Lawson and Cahill 1983) and cows (Ginther, 1970; Garrett et al., 1988) shortens the inter-oestrous interval by stimulating the earlier release of prostaglandin (PG) F₂α from the uterine endometrium which is responsible for the premature regression of the corpus luteum (Ginther, 1970; Ottobre et al., 1980; Garrett et al., 1988). Moreover, removal of a uterine horn adjacent to the luteal ovary (Ginther, 1968; Woody and Ginther, 1986) or infusion of indomethacin, an inhibitor of the synthesis of prostaglandins, into the uterine horn adjacent to the luteal ovary from day 8 to 11 prevents premature luteal regression after treatment with progesterone (Lewis et al., 1977).

Ovarian oxytocin also plays an important role in luteal regression in ewes by stimulating the secretion of the uterine luteolytic factor, PGF₂α (Fairclough et al., 1980, 1984; Flint and Sheldrick, 1983; Hooper et al., 1986; Moore et al., 1986) during the later stages of luteolysis. The ability of oxytocin to stimulate uterine PGF₂α secretion also depends on exposure to progesterone during the luteal phase of the cycle. Homanics and Silvia (1988) demonstrated that oxytocin can stimulate uterine secretion of prostaglandin F₂α in ovariectomized ewes after the animals have been exposed to progesterone for 7–10 days. Conversely, the release of the prostaglandin F₂α metabolite 13,14-dihydro-15-keto PGF₂α (PGFM) on days 12–15 of the oestrous cycle, after an oxytocin challenge on day 12, was prevented in ewes treated with the progesterone receptor

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Materials and Methods

Animal husbandry

All protocols were approved by the Animal Experimentation Ethics Committees of Victoria University of Technology (AECC 95/025) and CSIRO (Division of Animal Production).

Twelve, Border Leicester Merino cross ewes underwent ovarian auto-transplantation as described by Goding et al. (1967). Briefly, the right ovary was removed and the left ovary and its vascular pedicle were auto-transplanted into a jugular-carotid neck loop. The ovarian artery and the utero-ovarian vein were then anastomosed to the carotid artery and the jugular vein, respectively. During blood sampling, the ewes were housed individually in metabolism cages in a temperature-controlled room at 20°C and fed a ration of 800 g of pelleted food consisting of hammer-milled lucerne (60%) and oats (40%) once a day.

Experimental protocol

Ewes with auto-transplanted ovaries are not naturally cyclical. Therefore, oestrus was induced synchronously in the transplanted ewes by three i.m. injections of 125 μg synthetic PGF2α (Estrumate; ICI Pty Ltd, Sydney, NSW) given 15 days apart. After the third injection, oestrus was detected by inspection of the ewes twice a day for the presence of crayon marks after mating with a ram fitted with a sire-o-sine harness (Radford et al., 1960). The day that the ewe displayed oestrous behaviour was designated day 0. The twelve ovarian auto-transplanted ewes (n = 6 per group) were randomly assigned to receive an i.m. injection of 12.5 mg progesterone in 1 ml peanut oil (progestin; Intervet Pty Ltd, Sydney, NSW) or vehicle, twice a day, at intervals of 12 h, on days 1, 2 and 3 of the induced oestrous cycle.

Cannulation of ovarian and jugular veins

Cannulation of blood vessels was performed as described by Downing (1994) at least 12 h before the start of blood sampling. Briefly, cannulation was performed under local anaesthesia (10% lignocaine hydrochloride spray: xylocaine; Astra Pharmaceuticals, Sydney, NSW). The ovarian vein was cannulated by inserting a polyvinyl catheter into the jugular vein exteriorised in the skin loop, using a 12 G hypodermic needle as a trocar. The tip of the catheter was positioned at the junction of the ovarian and jugular veins. An additional polyvinyl catheter (50 cm) was inserted into the contralateral jugular vein to a distance of 10 cm. The free ends of both catheters were fitted with a blunted 19 G needle and connected to a three-way stopcock. The catheters were secured by suturing to the skin and wrapping elastoplast strips around the tubing at the site where it entered the neck. The catheters were filled with heparinized saline (1000 IU ml⁻¹).

Blood sampling

Blood samples (1.5 ml) were collected from the contralateral jugular vein at intervals of 1 h for 70 h starting on day 7 after oestrus. Blood was centrifuged at 1900 g for 15 min and the plasma collected and stored at −20°C until assayed for PGFM and progesterone concentrations. The procedure used for blood sampling was as follows: the heparinized saline was first removed from the contralateral jugular catheter and collected into a syringe fitted to the three-way stopcock connected to the catheter. The blood sample was then taken and placed in a heparinized glass tube and the catheter refilled with heparinized saline (50 µl ml⁻¹).

Ovarian venous blood (2 ml) was collected at the same time using the method described by Downing (1994). Briefly, the ewe was restrained to the side of the metabolism cage using a halter. A sphygmomanometer cuff was placed on the upper part of the skin loop and inflated above maximum venous pressure to obstruct the carotid artery and jugular vein. The forefinger and thumb were used to occlude the jugular vein in the lower part of the skin loop. The blood sample was then taken as described above. Every 4 h, samples of ovarian venous blood (approximately 5 ml) were taken instead of the 2 ml ovarian venous samples but following the same procedure. These samples were collected into heparinized 15 ml graduated centrifuge tubes after clearing the ovarian/jugular vein catheter of heparinized saline and allowing around 10 drops of ovarian venous blood to pass through the stopcock. The time taken to collect this sample was measured using a stopwatch.

Blood flow (ml min⁻¹) was calculated from the time required to collect a known volume of ovarian venous blood. The packed cell volume (PCV) was determined for each timed ovarian sample and the plasma flow (ml min⁻¹) was calculated by multiplying the blood flow by (100–PCV) divided by 100. Ovarian blood samples were assayed for oxytocin, and the secretion rate of oxytocin (ng min⁻¹) was obtained by multiplying the plasma flow (ml min⁻¹) by the concentration of oxytocin in the ovarian venous plasma (ng ml⁻¹).

Hormonal analysis

PGFM assay. Plasma PGFM concentrations were measured by radioimmunoassay as described by Burgess et al. (1990). The antiserum was raised in sheep against PGFM conjugated to porcine gelatine and was kindly supplied by R. I. Cox (CSIRO, Blacktown, NSW). At a final dilution of 1:50 000, the cross-reactivity of this antiserum with 13,14-dihydro-15-keto-PGF2α, 13,14-dihydro-15-keto-PGF2α, 15-keto-PGF2α and PGF2α was 100, 4.6, 1.9, 0.94, 0.33 and 0.02%, respectively, and was < 0.01% with antagonist mifepristone (RU 486) in the early to mid-luteal phase of the oestrous cycle (Thomas et al., 1985; Morgan et al., 1993).

Whether premature PGF2α release in response to early progesterone administration is due to a direct effect on the uterine PGF2α secretory system or is caused indirectly through the release of ovarian oxytocin is still uncertain. The aim of the present study was to determine whether PGF2α is released prematurely in response to early progesterone administration and, if so, whether such release is mediated via the premature release of ovarian oxytocin.
PGF<sub>2a</sub>, PGE<sub>2</sub>, 6-keto-PGF<sub>1α</sub>, 6-keto-PGE<sub>1α</sub>, PGE<sub>1</sub>, PGD<sub>2</sub>, PGB<sub>2</sub> and thromboxane B<sub>2</sub>. The sensitivity of the assay was 0.022 pmol per 200 µl. The intra- and interassay coefficients of variation were 9.35% and 10.4%, respectively.

**Progestosterone assay.** For the progestosterone assay, 200 µl of jugular venous plasma was extracted with 2 ml n-hexane as described by Rice et al. (1986). The antiserum was raised in sheep against progesterone-11α-BSA (generously provided by J. Malecki, Regional Veterinary Institute, Department of Agricultural and Rural Affairs, Bairnsdale, Victoria). At a final dilution of 1:9000, the crossreactivity of the antiserum with progesterone, 11α-hydroxy-progesterone, 5α-pregnane-3α-ol-20-one, 5β-pregnane-3α-ol-20-one and corticosterone was 100, 43.8, 15.9, 10.0 and 1.05%, respectively. Crossreactivity was < 1.0% with 11-deoxy cortisol, 5α-pregnane-3α,17α-diol-20-one, 5β-pregnane-3α,17α, 20α-triol-20-one, 5β-pregnane-3α,17α,20α-triol and 5α-pregnane-3β-ol-20-one. 0.7% with 17α-hydroxy-progesterone, < 0.4% with dehydroepiandrosterone, 0.3% with 20α-hydroxy-pregnen-3-one and < 0.2% with cortisol. The sensitivity of the assay was 0.22 pmol per 200 µl. All samples were measured in one assay and the intra-assay coefficient of variation was 16.5%.

**Oxytocin assay.** For the assay of oxytocin, 300 µl of the standards or ovarian plasma samples were mixed with 3 ml of chilled (-20°C) absolute ethanol using a modification of the method described in the Australian Laboratory Services Oxytocin Radioimmunassay Kit. After centrifugation at 1900 g for 20 min at 4°C, the supernatant was decanted and evaporated to dryness with filtered air at 37°C using a sample concentrator. The residues of these solutions were reconstituted in 300 µl of 0.05 mol phosphate buffer 1 -1 (pH 7.4). Standards and samples were incubated for at least 30 min at 4°C. One hundred microlitres of 125I tracer (10 000 c.p.m., Du Pont, Melbourne, Vic) was added to duplicate aliquots of 100 µl of extracted standards and samples, followed by 100 µl of antiserum diluted in assay buffer. The assay was incubated overnight at 4°C. Separation of the bound oxytocin was achieved by the addition of 0.05 ml (1 mg) bovine immuno-globulin serum (Calbiochem, Sydney, NSW) and 1 ml of 15% polyethylene glycol 6000 (Crown Scientific, Melbourne, Vic) at 4°C and then centrifuged at 3300 g for 20 min. The supernatant was aspirated and the radioactivity present in the precipitate was quantified using a Packard Crystal Gamma-Counter. The antiserum to oxytocin (GI, 137/1), was raised in sheep and was kindly donated by A. P. F. Frint (University of Nottingham, Sutton Bonington, Notts) and was used at a final dilution of 1:37 000. The crossreactivity of this antiserum with oxytocin, isotocin and melanocyte-stimulating hormone was 100, 0.79 and 0.04%, respectively. The crossreactivity was < 0.02% with mesotocin, pressinoic acid, prolactin, arginine-vaspressin and lysine-vasopressin. The sensitivity of the assay was 1.1 ng per 100 µl, the intra- and interassay coefficients of variation were 11.80% and 14.48%, respectively.

**Statistical analyses**

Pulses of ovarian vein oxytocin and jugular PGFM that were statistically significant were identified using a Pulsar program (Merriam and Wachtler, 1982). Assay noise was estimated by regression analysis of the standard deviation for the duplicate determinations and the mean at each point. Baseline was calculated from the contribution of circadian rhythms or other longterm trends, but not from fluctuations of shorter duration. The amplitudes of the ovarian oxytocin and peripheral PGFM pulses were calculated by subtracting baseline values and then rescaling in terms of SD units, by dividing the rescaled values by an estimate of assay noise. The amplitude in the rescaled pulses was then identified by applying height and duration criteria specified by user-defined cut-off points [G(1)] for pulsles. These calculations were repeated until two iterations produced the same values for pulses or until the preset limit of six iterations were completed. The quadratic (a), linear (b), and constant (c) terms for pulsles were as follows: for oxytocin, a = 0.00, b = 11.8, c = 0.00; and for PGFM, a = 0.00, b = 9.35, c = 0.00. The G(1) values for pulses with n = 1, 2, 3, 4 or 5 points above the baseline were selected by an empirical approach using a selected data series for each hormone from one animal which was then applied to all animals in the study. The following G(1) values were selected for oxytocin pulses: G(1) = 6.0, G(2) = 4.1, G(3) = 3.0, G(4) = 2.37, G(5) = 1.89; and for PGFM pulses: G(1) = 7.1, G(2) = 6.5, G(3) = 3.8, G(4) = 3.0, G(5) = 2.4. The values reported for intervals between ovarian oxytocin and peripheral PGFM pulsles were determined by calculating the time that elapsed between the highest values of sequential pulses. Coincident episodes in the secretion of oxytocin and PGFM were defined as those that showed an increase in the value of the PGFM pulse coincident with a defined oxytocin pulse. The plasma secretion rates of oxytocin and concentratons of PGFM pulses were determined as ng min⁻¹ and pg ml⁻¹, respectively, while the duration of that pulse was designated as τ, being the number of hours between the last time point before and the first time point after a significant increase in hormone concentration was detected by the Pulsar program. The areas under the curve of significant ovarian oxytocin and peripheral PGFM pulsles were calculated for each ewe and were expressed as (ng min⁻¹τ) and (pg ml⁻¹τ), respectively. Estimated parameters included overall mean concentrations, basal value, number of pulses, pulse amplitude, area under curve for the pulse and inter-pulse interval, and were carried out using the Pulsar analysis program.

The number of ewes that showed episodic pulses of oxytocin and PGFM were compared using a Chi-Squared test. The mean (± SD) concentratons of progesterone, basal oxytocin and PGFM between experimental and control animals were compared using Student’s unpaired t test.

**Results**

The concentration of progesterone in the peripheral plasma of the control ewes during the sampling period was 2.11 ± 0.62 ng ml⁻¹ (mean ± SD) which was not significantly different (P > 0.05) from that of progesterone-treated ewes. 2.32 ± 0.90 ng ml⁻¹. Progesterone concentrations remained raised during the sampling period, indicating the presence of a functional corpus luteum in both groups (Figs 1 and 2).

The basal secretion rate of ovarian oxytocin for ewes treated with progesterone early in the cycle (1.77 ± 1.56 ng min⁻¹)
was not significantly \((P > 0.05)\) different from that in control ewes \((2.41 \pm 0.57 \text{ ng min}^{-1})\). Administration of progesterone during the first 3 days of the cycle resulted in a significant \((P < 0.05)\) increase in the number of treated ewes showing pulses of both oxytocin \((n = 6 \text{ versus } n = 2)\) and PGFM \((n = 5 \text{ versus } n = 1)\) when compared with the control ewes. There was also an increase in the number of oxytocin pulses in ovarian venous plasma and PGFM pulses in peripheral plasma over days 7–9 after oestrus when compared with control ewes \((58 \text{ versus } 10 \text{ and } 29 \text{ versus } 4, \text{ respectively})\). The average
Fig. 1. The effect of vehicle injections during the first 3 days of the oestrous cycle on (●) oxytocin secretion in ovarian venous plasma and concentrations of (○) the prostaglandin F$_{2\alpha}$ metabolite 13,14-dihydro-15-keto PGF$_{2\alpha}$ (PGFM) and (△) progesterone in peripheral plasma from individual ewes with an auto-transplanted ovary (a–f), on days 7, 8 and 9 after oestrus. Statistically significant episodes in the secretion of ovarian oxytocin and PGFM are indicated by (*) and (T), respectively. (△) synchronous episodes of secretion of both compounds.
The number of pulses of oxytocin per ewe was 9.66 ± 5.5 and the interpulse-interval was 7.18 ± 5.78 h (Fig. 2) in the progesterone-treated ewes. The mean amplitude and area under the curve for ovarian oxytocin pulses were 6.27 ± 1.98 ng min⁻¹ and (10.05 ± 8.91 ng min⁻¹)t, respectively. In four of the six control ewes, no detectable pulse of ovarian oxytocin was found during days 7, 8 or 9 of the oestrous cycle. In one of the other two control ewes, four pulses of ovarian oxytocin were observed (Fig. 1d) at the start of blood sampling, with an average amplitude of 6.66 ± 3.72 ng min⁻¹. In the other ewe (Fig. 1b), there were six pulses of oxytocin in ovarian venous plasma. The mean amplitude and the area under the curve for

![Graphs showing ovarian oxytocin pulses and progesterone and PGFM levels over time.](image)
Fig. 2. The effect of progesterone injections during the first 3 days of the oestrous cycle on (●) oxytocin secretion in ovarian venous plasma and concentrations of (○) the prostaglandin F$_{2\alpha}$ metabolite 13,14-dihydro-15-keto PGF$_{2\alpha}$ (PGFM) and (▲) progesterone in peripheral plasma from individual ewes with an auto-transplanted ovary (a-f), on days 7, 8 and 9 after oestrus. Statistically significant episodes in the secretion of ovarian oxytocin and PGFM are indicated by (*) and (T), respectively. (●) synchronous episodes of secretion of both compounds.
ovarian oxytocin pulses for this ewe were \(6.14 \pm 2.79 \text{ ng min}^{-1}\) and \((5.22 \pm 1.44 \text{ ng min}^{-1})\), respectively.

In five of six ewes given progesterone, at least one detectable pulse in plasma PGFM concentration was observed during the sampling period. In these ewes, the mean amplitude and area under the curve for PGFM pulses were \(317.22 \pm 145.65 \text{ pg ml}^{-1}\) and \((383.36 \pm 111.77 \text{ pg ml}^{-1})\), respectively. The average number of PGFM pulses observed per ewe was \(5.8 \pm 1.9\) and sequential pulses of PGFM occurred at an average of \(10.32 \pm 8.7\) h intervals between day 7 and day 9. In one of the progesterone-treated ewes (Fig. 2e) pulses of oxytocin were observed without a corresponding increase in plasma PGFM concentrations (maximum \(276.5 \text{ pg ml}^{-1}\)). Basal peripheral plasma concentrations of PGFM for ewes treated with progesterone early in the cycle (97.43 \(\pm 22.92 \text{ pg ml}^{-1}\)) were not significantly \((P > 0.05)\) different from those in control ewes (91.32 \(\pm 20.65 \text{ pg ml}^{-1}\)). No significant pulses in plasma PGFM concentrations were found in the vehicle-treated ewes apart from one ewe (Fig. 1b). In this ewe, four pulses of PGFM were observed in peripheral plasma with an average amplitude of \(359.08 \pm 118.14 \text{ pg ml}^{-1}\). The average area under the curve for PGFM pulses for this ewe was \((555.61 \pm 251.98 \text{ pg ml}^{-1})\).

In ewes that received progesterone early in the cycle, there were at least twice as many oxytocin pulses compared with PGFM pulses, and at least half (51.7%) of the plasma PGFM pulses occurred coincidently with a significant increase in ovarian oxytocin. In contrast, only 22.4% of ovarian oxytocin pulses were coincident with, or preceded, pulses of plasma PGFM concentrations.

Discussion
In the current study, the ovarian auto-transplanted ewe was used as a model to determine the secretion rate of ovarian oxytocin and peripheral plasma concentrations of PGFM and progesterone in ewes given progesterone early in the oestrous cycle.

Hooper et al. (1986) reported that the concentration of oxytocin in peripheral plasma ranged from 20 to 220 \(\text{pg ml}^{-1}\) until day 9 after oestrus and was much lower than those in utero–ovarian venous plasma (230–1020 \(\text{pg ml}^{-1}\)). Concentrations of oxytocin in ovarian venous plasma detected in the present study were 440–1430 \(\text{pg ml}^{-1}\) which are similar to those detected by Hooper et al. (1986) in the utero-ovarian vein (50–1499 \(\text{pg ml}^{-1}\)) over days 13–16 after oestrus. These observations indicate that, in the current study, oxytocin in ovarian venous plasma represents luteal rather than posterior pituitary secretion.

Administration of progesterone from day 1 to day 3 of the oestrous cycle resulted in a significant increase in the number of ewes showing pulses of both oxytocin and PGFM. There was also an increase in the number of oxytocin pulses in ovarian venous plasma and PGFM pulses in peripheral plasma over days 7–9 after oestrus in the progesterone-treated ewes when compared with the vehicle-treated ewes. The effect of progesterone treatment on the pattern of plasma PGFM concentration is consistent with previous reports in ewes (Ottobre et al., 1980) and cows (Ginther, 1970; Garrett et al., 1988). The data reported in these previous studies indicated that treatment with progesterone early in the cycle at a dose equivalent to that used in the present study advanced the time at which PGF₂<sub>19</sub> was released in intact animals and caused premature luteolysis. Previous reports had shown that uterine PGF₂<sub>19</sub> acts on the adjacent ovaries in intact animals through a local pathway by counter-current transfer from the uterine vein to the ovarian artery (Barret et al., 1971; McCracken et al., 1972) and causes regression of the corpus luteum. Uterine PGF₂<sub>19</sub> is cleared rapidly from the blood after one passage through the lungs as it is metabolized in the pulmonary vascular bed (Piper et al., 1970). The finding, in the study reported here, of intermittent pulses of PGFM, but an absence of luteolysis in ovarian auto-transplanted ewes, confirms previous views that uterine PGF₂<sub>19</sub> acts by a local counter-current mechanism to induce luteolysis in intact ewes (Barret et al., 1971; McCracken et al., 1972).

Peripheral progesterone concentrations are normally low (<1 ng ml⁻¹) during the first 4 days of the oestrous cycle (Webb et al., 1981; Garrett et al., 1988) and administration of exogenous progesterone during days 1–4 after oestrus to cyclic cows increases peripheral plasma concentrations of progesterone during days 2–5 after oestrus to concentrations comparable with those of control cows on day 5 to day 9 of the oestrous cycle (Garrett et al., 1988). In ewes, plasma progesterone concentrations normally start to fall on day 12 or day 13 after oestrus, or around the time of the first significant increase in the secretion of uterine PGF₂<sub>19</sub>. This would indicate that 7–8 days are required as a minimum exposure time of the uterus to progesterone before the uterus secretes PGF₂<sub>19</sub> (Baird et al., 1976; Homanics and Silvia, 1988). Previous studies have suggested that the duration of exposure of the uterus to progesterone plays a key role in the regulation of the timing of the initial increases in the release of PGF₂<sub>19</sub> (Baird et al., 1976; Ottobre et al., 1980). The results of the present study give further support to this premise, since the administration of progesterone on days 1, 2 and 3 of the oestrous cycle stimulated premature pulses of peripheral plasma PGFM secretion over days 7–9 after oestrus.

It is possible that progesterone also causes the premature release of uterine PGF₂<sub>19</sub> through ovarian oxytocin release. It is now well accepted that, in intact ewes, ovarian oxytocin stimulates uterine PGF₂<sub>19</sub> release during the late luteal phase of the oestrous cycle (Flint and Sheldrick, 1983; Hooper et al., 1986). The data obtained from the present study, therefore, raise the possibility that progesterone also acts to regulate the timing of the release of ovarian oxytocin, since the administration of progesterone to ewes early in the cycle resulted in a significant increase in the number of ewes that exhibited pulses of ovarian oxytocin compared with control ewes.

The results from the present study and other reports (Fairclough et al., 1980, 1983; Hooper et al., 1986) also show that oxytocin pulses in ovarian or utero–ovarian venous plasma, or oxytocin–neurophysin in jugular venous plasma frequently occur in the absence of any significant increase in utero–ovarian PGF₂<sub>19</sub> or peripheral PGFM pulses, and indicates that ovarian oxytocin release can occur independently of uterine PGF₂<sub>19</sub> release. Furthermore, the largest number of pulses with maximum ovarian oxytocin secretion rate was observed on day 7, whilst the largest number of pulses that
occurred with maximum concentrations of plasma PGFM were observed on day 8 of the oestrous cycle. The study reported here and that of Hooper et al. (1986) have shown that pulses of uterine PGF₂α or PGFM occur without simultaneous pulses of oxytocin. One possible explanation for this observation is that progesterone acts, at least in part, directly on the uterus to stimulate uterine PGF₂α release. Alternatively, oxytocin pulses may have been missed in the present study because of the sampling schedule, or may not have been detected given the stringent criteria used to identify significant pulses by the Pulsar program.

Hooper et al. (1986) reported that, in ewes, 97% of all pulses of uterine PGF₂α release were accompanied or followed by pulses of oxytocin in the ovarian vein, while only 56% of oxytocin pulses were coincident with pulses in uterine PGF₂α. The sampling frequency in the present study (every hour) resulted in a similar pattern, although in the present study, only 51.7% of plasma PGFM pulses were associated with pulses in ovarian oxytocin. This difference may be due to the different statistical criteria used to detect the significant pulses of oxytocin and PGFM. In addition, the interpulse interval was greater in the present study than in that of Hooper et al. (1986): oxytocin (1.6 ± 0.94 versus 2.5 ± 0.19) and PGF₂α (1.0 ± 0.3 versus 1.3 ± 0.2) pulses per 12 h. However, the area under the curve for ovarian oxytocin and PGFM pulses in the present study were higher and longer, covering more consecutive samples per pulse than in the study by Hooper et al. (1986). The observation showing several pulses in ovarian oxytocin without a corresponding increase in plasma PGFM concentrations could be due to uterine refractoriness to oxytocin at that time. Alternatively, it may reflect the variation in response, in individual animals, to progesterone treatment or to the oxytocin stimulus.

As indicated in previous studies, there is a significant effect of progesterone and oestriadiol on the pattern of plasma PGFM release in response to oxytocin. These studies have shown that low progesterone and high oestriadiol concentrations increase the plasma PGFM response to oxytocin on day 12 of a simulated oestrous cycle (Beard et al., 1994). Chronic treatment with progesterone also enhances uterine secretion of PGF₂α in response to an i.v. injection of oxytocin (Silvia and Homanics, 1988). The mechanism involved in the action of progesterone is unknown, although Eggleston et al. (1990) reported that treatment of intact ewes with progesterone early in the cycle induced a premature increase in the expression of mRNA encoding uterine prostaglandin H endoperoxide synthase. However, the mechanism(s) by which progesterone acts to regulate the luteolytic process in sheep is not fully understood.

It has been suggested that oxytocin receptors play an important role in regulating the duration of the oestrous cycle in sheep (McCracken et al., 1984), possibly through the downregulation of progesterone receptors and consequently the diminished inhibition of endometrial oestrogen and oxytocin receptors (McCracken et al., 1984; Watkins and Hamon, 1993). There is considerable evidence to indicate that concentrations of uterine oxytocin receptors are regulated by circulating concentrations of the steroid hormones (McCracken et al., 1984; Sheldrick and Flint, 1985), with oestriadiol acting to induce receptor formation and progesterone acting, early in the cycle, to inhibit oxytocin receptors. During the later stages of the oestrous cycle, progesterone appears to lose its inhibitory influence, possibly through downregulation of uterine progesterone receptors (Vallet et al., 1990; Zhang et al., 1992).

It is also possible that, in the study reported here, early progesterone treatment stimulated uterine oxytocin receptor concentrations, since on some occasions there were coincident pulses of ovarian oxytocin and PGFM at a time when oxytocin does not usually stimulate uterine PGF₂α release (Fairclough et al., 1984).

The present study is the first to indicate that ovarian oxytocin secretion is stimulated by the administration of progesterone during the first 3 days of the oestrous cycle, thus, raising the possibility that, in intact ewes, progesterone may directly or indirectly regulate ovarian oxytocin and uterine PGF₂α secretion systems and thus, the timing of luteal regression.

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