Is progesterone the key regulatory factor behind ovulation rate in sheep?


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Abstract

Ovarian antral follicles in the ewe grow in an orderly succession, producing 3 to 4 waves per estrous cycle. In prolific sheep, some large antral follicles from the second-to-last wave of the estrous cycle are added to the ovulatory follicles emerging just before estrus to give a higher ovulation rate; it is feasible that regression of these follicles is prevented by an increase in serum concentrations of FSH or LH pulsatility at proestrus. Proliptic sheep tend to have a shorter luteal phase than nonproliptic ewes and there is a great deal of evidence that luteal progesterone (P4), in addition to regulating LH release, may govern the secretion of FSH heralding the emergence of follicular waves. The specific purpose of this study was to determine whether or not extending the duration of the luteal phase in proliptic sheep to that typically seen in nonproliptic breeds would alter the follicle wave dynamics and ovulation rate. In 2 separate experiments, exogenous P4 (7.5 mg per ewe intramuscularly) was administered on day 11 at PM and day 12 at AM (day 0 = first ovulation of the interovulatory interval studied) in moderately proliptic Rideau Arcott × Polled Dorset ewes (experiment 1, n = 8) and highly prolific Olkuska ewes (experiment 2, n = 7; TRT), whereas the equinumerous groups of animals served as controls (CTR). Transrectal ovarian ultrasonography was performed daily, and jugular blood samples were drawn twice a day from day 9 until the next ovulation. Progesterone injections resulted in relatively uniform increments in serum P4 levels, but the mean duration of the interovulatory interval did not differ (P > 0.05) between TRT and CTR groups of ewes in either experiment. The mean ovulation rate post-treatment was 1.6 ± 0.2 vs 3.2 ± 0.4 (experiment 1, P < 0.001) and 3.2 ± 0.8 vs 4.0 ± 1.0 (experiment 2, P > 0.05) in TRT vs CTR, respectively. The number and percentage of ovulating follicles from the penultimate wave of the interovulatory interval studied was 0.25 ± 0.16 vs 1.75 ± 0.45 (P < 0.01) and 25.0 ± 16.4% vs 75.0 ± 16.4% (P < 0.05) in experiment 1, and 0.50 ± 0.30 vs 1.60 ± 0.40 (P < 0.05) and 13.8 ± 9.0% vs 53.4 ± 16.7% (P < 0.05) in experiment 2, for TRT vs CTR, respectively. In summary, administration of P4 at the end of diestrus decreased the incidence of ovulations from the penultimate wave of the estrous cycle in both the moderately and highly prolific strains of sheep, but it reduced the ovulation rate only in moderately prolific ewes.

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1. Introduction

Sheep have a broad spectrum of fertility, which makes them a perfect model species to study the factors controlling ovarian function and ovulation rate. Lamb productivity of the ewe is genetically predetermined and it can vary among breeds from ~1 in typically monovular breeds (eg, Merino del Pais) to >5 in the carriers of fecundity genes such as FecB or Booroola genotypes [1,2]. However, in spite of the fact that products of certain genes may program this tremendous biodiversity [3], the hormonal control of ovarian follicle kinetics and ovulation rates in sheep is not completely understood. The physiological regulation of ovulatory follicle development remains unclear [1], and continued investigations into the mechanisms governing ovulation and fertility have been central in reproductive biology across species [3].

Numerous studies have been conducted in an attempt to determine the regulation of ovarian follicular development in domestic animals, including small ruminants [1,4,5]. Antral follicles in the ewe emerge in an orderly succession, producing 3 to 4 follicular waves per 17-day estrous cycle [1,2]. Each wave consists of 1 to 4 follicles attaining an ostensibly ovulatory size before regression or ovulation and is preceded by a transient rise in FSH secretion [1,2]. From earlier studies of temporal relationships between FSH secretory patterns and wave emergence in ewes, it appears that FSH controls the initial stage of antral follicle growth, from emergence to the end of the growth phase, but ensuing follicular development and demise are independent of FSH [1,2,6]. Recurrent pulses of LH also promote the growth of large ovine follicles and prolong the static phase of their lifespan [7]. In prolific ewes, some large antral follicles from the penultimate wave of the estrous cycle are added to the ovulatory follicles emerging just before estrus to give a higher ovulation rate [1,2,8,9]. Prolific sheep tend to have a shorter luteal phase than nonproliﬁc breeds, and there is a great deal of evidence supporting the notion that luteal progesterone (P4), in addition to regulating pulsatile release of LH, may govern the secretion of FSH [4,5]. Therefore, it is feasible that the onset of regression of some penultimate wave follicles in prolific ewes is delayed or prevented by the next increase in serum concentrations of FSH or gradual increase in LH pulsatility at proestrus.

Thus, our general supposition was that luteal P4 was the driving regulatory force behind ovulation rates in ewes. To test if this theory stands true, our study was designed to monitor antral follicle development and assess the ovulation rate with transrectal ultrasonography or diagnostic laparoscopy in cyclic ewes in which elevated, midluteal phase levels of P4 were maintained through administration of the exogenous gestagen at the expected end of diestrus. We hypothesized that maintaining high P4 concentrations for 1 to 2 more day(s), as seen in nonproliﬁc breeds of sheep [1,2,10–12], would alter follicular wave development and ovulation rate in prolific sheep to resemble those in nonproliﬁc ewes. Two independent experiments, with identical designs, were conducted on a moderately prolific crossbred ewes (Rideau Arcott × Polled Dorset crosses), with 2 to 3 ovulations per cycle [13; experiment 1], and on a highly prolific sheep strain (Olkuska breed), in which certain individuals were the carriers of high fecundity genes and all animals had consistently produced 3 to 6 lambs per breeding [14,15; experiment 2].

2. Materials and methods

All experimental procedures performed on live animals were in compliance with the guidelines of the Canadian Council on Animal Care or with the European Community (EC) directives for animal experimentation and had been formally approved by the local animal care or bioethics committees.

2.1. Experiment 1

This experiment was performed during the breeding season (October–November) on 16 clinically healthy, nulliparous Rideau Arcott × Polled Dorset ewes aged 11 to 28 mo, with a mean (± standard error of the mean [SEM]) body weight of 67.9 ± 8.2 kg. Animals were housed at the Ponsonby Research Station near Guelph, ON, Canada (latitude: 43° 83′ 30″N) under natural conditions of photoperiod and ambient temperature. The ewes were kept in sheltered dry lots, fed daily maintenance rations of alfalfa pellets and hay, with water and cobalt-iodized salt licks available ad libitum. Estrus was detected with a 2-electrode vaginal impedometer (Electronic Oestrus Detector; Dramiński Electronics in Agriculture, Olsztyn, Poland) recording changes in vaginal mucous impedance that had been validated for the present application in sheep [16,17]. Two intramuscular injections of 2 mL of Prostaglandin F2α alpha (Pgf2α) (5 mg/mL; Lutalyse; Pharmacia Animal Health, Orangeville, ON, Canada) were initially given 9 d apart to 22 animals. Detection of estrus continued until the vaginal mucus impedance declined below 40 Ω (onset of luteolysis) and then increased above 40 Ω (ovulation = day 0) in the first 16 ewes. Subsequently, the ewes were randomly allocated to the treatment and control groups (n = 8 in each group). The treatment group animals received 2 injections of 7.5 mg of progesterone (P4; Chiron Founding Pharmacy; Guelph, ON, Canada) in 0.5 mL of corn oil intramuscularly, in the evening of day 11 and in the morning of day 12; control ewes received injections of the vehicle only (sterilized corn oil). Corn oil was used as the vehicle to allow for a slow release of P4 after the treatments applied 12 h apart [4].

Blood samples were drawn twice daily via jugular venipuncture into 10-mL coagulant-free vacutainers (Becton Dickinson, Rutherford, NJ, USA). The samples were permitted to clot for 12 to 24 h at room temperature and then centrifuged at 1,500 × g for 10 min. Serum was harvested and stored at –20°C until analyses at the later date.

Daily transrectal ultrasonography of ovaries used a portable Aloka 900-SSD echo-camera (Aloka; Tokyo, Japan) connected to a rigid 7.5-MHz biplane rectal transducer. The linear-array component of this transducer was used to visualize all antral follicles ≥2 mm in diameter and luteal structures. All images were displayed on the viewing screen of the echo camera at magnification ×1.5 to ×2, and the relative position of individual antral follicles was sketched on ovarian charts. In addition, all ultrasonographic images were recorded on a DVD recorder.
(DVR-510H; Pioneer Electronics of Canada, Oakville, ON, Canada) for subsequent analyses of ovarian data. Daily ultrasonographic examinations of ovaries were carried out by the same experienced operator from the time of the second PgF2α injection until ovulation detection (day 0) and again from day 9 until the next ovulation of the inter-ovulatory interval studied. Ovulations were detected by a disappearance of large antral follicles (≥5 mm in size) followed by a formation of corpora hemorrhagica approximately 24 h after follicle rupture [18]. Seven days after the estrus post-treatment, all animals underwent laparoscopic examinations of ovaries using the Tele Pack Vet X Led video endoscope (Karl Sorz GmbH & Co KG, Tuttingen, Germany) to enumerate corpora lutea. Approximately 10 min before endoscopy, each ewe was given an intramuscular injection of 0.4 mL of Rompun (20 mg/mL of xylazine; Bayer, Toronto, ON, Canada) and at the start of the procedure, each animal received 1.5 to 2.0 mL of local subcutaneous (s.c.) anesthetic (1% lidocaine hydrochloride; Zoetis, Kirkland, QC, Canada) to the sites of trocar punctures in the abdominal wall.

Serum samples were analyzed by the Prairie Diagnostic Services, University of Saskatchewan, Saskatoon, SK, Canada, using validated RIAs to determine concentrations of FSH [19], P4 [20], and estradiol (E2; [21]). The sensitivities of assays defined as the lowest concentration of a hormone that was capable to significantly displace radiolabeled hormone from the binding with the primary antibody were 0.1 ng/mL, 0.03 ng/mL, and 1 pg/mL for FSH, P4 and E2 assays, respectively. For reference sera with mean FSH concentrations of 0.92 and 3.45 ng/mL, the intra-assay coefficients of variation (CVs) were 6.6% and 9.3%, respectively. For reference sera with mean E2 concentrations of 8.1 and 21.2 pg/mL, the intra-assay CVs were 8.6% and 11.1%, respectively. For reference sera with mean P4 concentrations of 0.27 or 1.23 ng/mL, the intra-assay and interassay CVs were 6.3% and 5.1% and 13.0% and 3.8%, respectively. The ranges of standards were as follows: (1) ovine FSH (ofFSH): 0.06 to 16.0 ng/mL (ofFSH-SiAFP-RP-2); (2) E2;17β: 1.0 to 50 pg/mL; and (3) P4; 0.03 to 10.0 ng/mL.

2.2. Experiment 2

The second experiment was conducted during the month of November in the field research station of the Agricultural University of Kraków, Poland, situated in Bielany (latitude: 50°03′31″N), and it employed 14 multiparous Polish Olkuska ewes aged 3 to 5 yr and weighing between 52 and 73 kg. The animals were housed in a barn under natural conditions of light and ambient temperature. All ewes received daily food rations formulated to provide 100% of nutritional requirements [22]; water and mineral licks were available ad libitum. Among 7 animals allocated to each group, 2 ewes had been identified as carriers of a high fecundity gene (F-gene homozygous), 3 were heterozygous and 2 were noncarriers of the F-gene [14]. Experimental design, various methods of data acquisition and drugs used were all identical to those described for experiment 1, except for the procedures and pharmaceuticals detailed in the following paragraph.

Progesterone for injections was purchased from Sigma Life Science (Sigma–Aldrich, St. Louis, MO, USA). Estrus was detected twice daily (2 × 20 min), starting 2 d after the second injection of PgF2α (Lutalyse; 10 mg/ewe), with a fertile ram fitted with an apron preventing penetration during mounting. Transrectal ovarian ultrasonography used the Aloka ProSound 2 scanner (Hitachi Aloka Medical; Tokyo, Japan) equipped with a stiffened (plastic extension) 7.5-MHz linear-array transducer. Ultrasound images of both ovaries were digitally acquired and recorded directly on a computer (at a resolution of 640 × 480 pixels). Ewe undergoing endoscopy received a single intramuscular injection of 0.4 mL of Nefrasin vet. (containing 20 mg/mL of xylazine hydrochloride; aniMedica Polska, Gdynia, Poland) and local s.c. injections of 1 to 1.5 mL of polocainum hydrochloricum (2%) with adrenaline (0.005%; Biowet, Drzawel, Poland). Blood sera were assayed by the Animal Sciences Research Center at the University of Missouri. All samples were analyzed for P4 concentrations in a single assay [23] with the standard curve range from 0.005 to 40 ng/mL and detection limit of 0.1 ng/mL; an intra-assay CV was 2.0%. Estradiol concentrations were determined with 3 assays [24]; intra-assay and interassay CVs were 4.3% and 4.4%, respectively; assay range was 0.05 to 12 pg/mL and minimum detectable concentration was 0.01 ng/mL. Serum concentrations of FSH were determined in a single RIA [24] with sensitivity of 0.1 ng/mL and concentrations of a standard (ofSH AFP5679c) ranging from 0.1 to 15.0 ng/mL; an intra-assay CV was 2.0%.

2.3. Data analysis

A cycle detector program (or the threshold adaptive technique [25]) was used to determine the peaks of successive FSH and E2 fluctuations during the period from days 11.5 to 16.5; the number and concentrations of peaks detected in individual ewes were computed. This method uses an iterative algorithm that scans serial data to identify increases and decreases greater than a preset threshold value (determined using intra-assay CVs). One complete cycle or fluctuation (nadir-to-peak-to-nadir) is regarded as an increase that is greater than threshold followed by a decrease, which is also greater than threshold.

A follicular wave was defined as a group of follicles that grew synchronously from 2 or 3 mm to ≥5 mm in diameter before regression or ovulation [2]. The day of wave emergence, the number of all follicles, and the number and proportion of ovulatory follicles in the penultimate and last wave of the estrous cycle studied were noted. In addition, all antral follicles of the wave were analyzed for their maximum diameter, duration of the growth phase, static phase and regressing phase, as well as the mean growth and regression rate. The growing phase was defined as the time taken to grow from 2 to 3 mm to the maximum diameter attained before regression or ovulation, and the regressing phase was the number of days taken to regress from its maximum diameter to 2–3 mm. The time between the end of growth and the onset of regression or ovulation was the static phase of follicular lifespan. The number and diameter of all ovulating follicles were recorded.

All data sets were initially screened for outliers using the Dixon Q-test. Single-time point mean values were compared between the treated and control ewes by
Student t test. For 2 data sets with nonhomogenous vari-
ances, the Mann–Whitney rank sum test was performed.
Statistical differences among serial hormonal and ovarian
data were assessed by 2-way repeated-measures analysis of
variance (ANOVA, general linear model procedures)
using SigmaPlot (version 11.0 for Windows, 2010 Systat
Software, Inc Richmond, CA, USA). Proportions were
compared by a $\chi^2$-test (Brandt–Snedecor formula [26]).
The results were expressed as mean ± SEM, and the statistical
significance was regarded as $P < 0.05$. The present experi-
ments used different ultrasonographic equipment and
hormone assays, and so no direct comparisons of data were
made between breeds.

3. Results

3.1. Experiment 1

Mean serum concentrations of P₄ in the treated ewes
increased by 1.42 ± 0.36 ng/mL and 1.24 ± 0.49 ng/mL
approximately 12 h after the first and the second P₄
injection, respectively, ($P > 0.05$). Progesterone-treated
ewes (TRT) exceeded ($P < 0.05$) control animals (CTR) in
circulating P₄ concentrations from days 12 to 14 (Fig. 1). In
TRT group, serum P₄ concentrations decreased ($P < 0.05$)
from a maximum value on day 12.5 to day 14 and again
($P < 0.05$) on day 15, whereas P₄ concentrations in control
animals declined ($P < 0.05$) from day 11.5 to day 13.5. There
were significant main effects of the group and time for
mean serum concentrations of E₂ over the period of study,
but post-ANOVA analyses did not reveal significant differ-
ces over time in either group of ewes. Circulating concentrations of E₂ were greater ($P < 0.05$) in TRT than
those in CTR ewes on day 14.5 (Fig. 1). As with the E₂
concentrations, there were no significant differences in
mean serum FSH concentrations within either group of
ewes despite the significant main effect of Time. Progesterone-treated ewes exceeded ($P < 0.05$) their respective controls in the mean number of FSH peaks detected
during the 5 d after the first P₄ injection (Table 1).
The first FSH peak post-treatment occurred ~1 d later
($P < 0.05$), but the peak concentration was significantly
higher in TRT compared with CTR animals.

The mean ovulation rate at the beginning of the study
did not differ ($P > 0.05$) between animals subsequently
allocated to the treatment or control group, but the mean
ovulation rate and number of corpora lutea (CL) were lower
($P < 0.001$) in TRT compared with CTR ewes after the
treatment with exogenous P₄ (Table 2). One P₄-treated ewe
with 2 ovulations and 1 control animal with 3 ovulations
detected ultrasonographically had 1 and 2 CL, respectively,
during laparoscopic examination post-treatment; in all
remaining sheep, the numbers of ruptured follicles and
luteal structures were identical. Ovarolic follicles emerging in the penultimate wave were recorded in 25% of
TRT (2/8) and 75% of CTR (6/8) ewes ($P < 0.05$). The total
number, and the number and percentage of ovulating fol-
licles emerging in the penultimate wave of the estrous
cycle were significantly lower in TRT than those in CTR
group, but the mean growth rate of all penultimate wave
follicles and the diameter of ovulatory follicles were both
greater ($P < 0.05$) in TRT ewes (Table 2). There were no
differences ($P > 0.05$) in any other characteristic of pen-
ultimate or final wave follicles between CTR and TRT ewes
in experiment 1. There were no significant differences in
mean daily numbers of ovarian antral follicles (2, 4, and
≥5 mm in size) within either group of experimental ani-
imals or between the 2 groups of ewes studied except for
the number of 3-mm follicles that was greater ($P < 0.05$)
in TRT than that in CTR ewes from days 15 to 17 (or from 3.5 to
5.5 d after the first P₄ dose; Fig. 2A).

3.2. Experiment 2

Mean P₄ concentrations increased in the treated ewes
by 1.06 ± 0.28 ng/mL and 1.45 ± 0.40 ng/mL after the first
and the second P₄ injection, respectively ($P > 0.05$). In TRT
ewes, serum P₄ concentration decreased ($P < 0.05$) from
a maximum value on day 12.5 to day 14.5, whereas in CTR
ewes, they declined ($P < 0.05$) from day 11.5 to day 15
(Fig. 1). Neither mean serum concentrations of FSH and E₂
nor various characteristics of peaks in circulating concen-
trations of both hormones varied between the 2 subsets of
ewes except for the number of detected E₂ peaks that was
nearly 2 times greater ($P < 0.001$) in TRT than that in CTR
group (Table 1).

Two of 7 TRT ewes (29%) and 6 of 7 CTR animals (86%) had an ovulatory follicle(s) that emerged in the penulti-
mate wave of the estrous cycle ($P < 0.05$). The number and
percentage of ovulatory follicles emerging in the penulti-
mate wave of the estrous cycle in Olkuska sheep were
significantly greater in CTR than those in TRT ewes. In
addition, the mean growth and regression rate of all folli-
cles ≥5 mm in diameter in the penultimate wave were
greater ($P < 0.05$) in TRT compared with CTR ewes (Table 2).
The remaining characteristics of follicles emerging in the
penultimate and final wave of the interovulatory interval
did not vary ($P > 0.05$) between CTR and TRT Olkuska
sheep. Mean daily numbers of ovulatory-sized follicles
(≥5 mm in diameter) decreased sharply ($P < 0.05$) from
day 15 to day 17 in P₄-treated ewes (Fig. 2B). Mean daily
numbers of follicles in other size categories did not vary
($P > 0.05$) either over time or between CTR and TRT ewes in
experiment 2.

4. Discussion

In the present study, exogenous P₄ injections given
before the end of diestrus in prolific ewes were intended to
delay a decline in serum P₄ to basal levels by 1 to 2 d to
resemble the situation seen in less prolific, meat breeds of
sheep [10–12]. Both injections were followed by an
increase in serum P₄ concentrations, but 12 h after the
administration stopped, serum P₄ levels began to decline.
In the first experiment, mean serum P₄ concentrations
remained higher in the treated ewes until day 14 post-
ovulation. Quite unexpectedly, however, circulating concentrations of P₄ were numerically higher in control
compared with treated Olkuska sheep (experiment 2) from
1 to 3 d post-treatment. These differences between the 2
experiments can be attributed to variable endogenous P₄
concentrations in control Olkuska ewes (experiment 2), as
the pattern of changes in P\textsubscript{4} concentrations in the treated ewes was strikingly similar in both genotypes of sheep studied. The latter suggests that elevated P\textsubscript{4} levels at the end of diestrus may dictate and synchronize the rate of functional luteal regression in cyclic ewes. In both experiments, a decline in circulating P\textsubscript{4} concentrations to basal levels (regarded as a period encapsulating all days on which P\textsubscript{4} concentrations did not vary statistically from a minimal value recorded on day 16.5) occurred earlier in controls than in the treated ewes (~2.5 d earlier in experiment 1 and ~1.5 d earlier in experiment 2). Induction of a prolonged luteal phase allowed us to determine the changes that P\textsubscript{4} inflicted on circulating FSH and E\textsubscript{2} concentrations and antral follicular development.

In experiment 1, 2 injections of exogenous P\textsubscript{4} on days 11 and 12 after ovulation were followed by FSH peaks on days 12.5, 14.5, and 16.5, whereas untreated controls had 2 waves on days 11.7 and 15.8 postovulation. In an earlier study, administration of exogenous P\textsubscript{4} twice daily from day 0 (ovulation) to day 4 was associated with the occurrence of...

Fig. 1. Mean (±standard error of the mean) serum concentrations of progesterone, FSH, and estradiol from days 11 to 16.5 (day 0 = first ovulation of the estrous cycle studied) in moderately prolific Rideau Arcott × Poled Dorset ewes (experiment 1) and highly prolific Olkuska sheep (experiment 2) that received progesterone (progesterone-treated) or vehicle injections (control) on day 11 at PM and day 12 at AM. Syringe symbols along the x-axis denote the timing of progesterone injections. Asterisks indicate the differences (P < 0.05) between groups, and different letters denote significant differences over time.
### Table 1

Characteristics of peaks in serial FSH and estradiol (E2) concentrations detected from days 11.5 to 16.5 in individual Rideau Arcott × Polled Dorset ewes (experiment 1) or Olkuska sheep (experiment 2) that received intramuscular progesterone (treatment) or vehicle (control) injections on day 11 at PM and day 12 at AM.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Experiment 1 (Rideau Arcott × Polled Dorset)</th>
<th></th>
<th>Experiment 2 (Olkuska)</th>
</tr>
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<tr>
<td></td>
<td>Treatment (n = 8)</td>
<td>Control (n = 8)</td>
<td>Treatment (n = 7)</td>
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<td><strong>Follicle-stimulating hormone peak characteristics</strong></td>
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<td>No. of peaks (days 11.5–16.5)</td>
<td>2.7 ± 0.2ac</td>
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<td>Time of occurrence</td>
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<td>14.4 ± 0.4a</td>
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<td>Peak 3</td>
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<td>Peak 4</td>
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<td>No. of peaks (days 11.5–16.5)</td>
<td>3.2 ± 0.2</td>
<td>2.7 ± 0.3</td>
<td>2.9 ± 0.3ac</td>
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<tr>
<td>Time of occurrence</td>
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<tr>
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<td>[4.08]</td>
<td>[6.33]</td>
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Abbreviation: ND, not detected.

*abc* P < 0.05; *ABc* P < 0.01.

Here, day 0 = first ovulation of the estrous cycle studied. Data obtained for only 1 or 2 animals per group are shown in brackets. Data are expressed as mean ± standard error of the mean.

*ac* Mann–Whitney rank-sum test; day 0 = ovulation.

### 2 FSH peaks on days 1.5 and 3.9, whereas the control group of cyclic ewes only exhibited 1 FSH peak during the treatment period (on day 4.2; [5]). Therefore, creation of mid-cycle P4 concentrations early or late in the luteal phase of ewes increases the number of fluctuations in serum FSH concentrations. In the aforementioned study by Baby et al [5], serial P4 injections given early in metestrus advanced follicle wave emergence in treated compared with control ewes; as a result, P4-treated ewes had 1 more follicular wave compared with control ewes during the same time frame (days 0 to 6 from ovulation) [5]. Cyclic ewes with 4 follicular waves per estrous cycle (emerging on ~ days 0, 4, 8, and 12 after ovulation) also have higher serum P4 concentrations than their counterparts with 3 waves (emerging on ~ days 0, 5.5, and 10) [4]. Injections of exogenous ovine FSH resulting in an additional FSH peak midway through the endogenous interpeak intervals in anestrous ewes consistently induced wave emergence ~ 1 d after FSH peaks [6]. The characteristics of induced follicular waves did not differ from those of endogenous follicular waves (eg, mean duration of the growth phase spans ~3 d [6]). In the present study, an additional endogenous FSH peak detected approximately 2 d before the preovulatory discharge of gonadotropins in P4-treated ewes occurred too late to stimulate the growth of ovarian antral follicles to ovulatory sizes (follicle wave emergence) before the end of the interovulatory period studied. This notwithstanding, a decline in numbers of ovulating penultimate wave follicles in the treated ewes in this study does not seem to be caused by diminished FSH secretion. It is equally unlikely that changes in LH pulsatility affected the numbers of emerging antral follicles, as they normally do not vary among waves developing at different stages of the estrous cycle [2]. However, it is attractive to speculate that these ovarian effects of P4 late in diestrus are direct [27] and may involve transient alterations in follicular responsiveness to gonadotrophic stimuli and/or biosynthesis of various locally acting promoters of antral follicle growth [3,28]; the underlying molecular mechanisms remain to be elucidated.

In the first experiment, P4-treated ewes exceeded control animals in circulating E2 concentrations 2.5 d after the second P4 dose (day 14.5). In cyclic ewes, the largest antral follicles of waves begin to produce E2 at the time of emergence (ie, recruitment from the pool of 2 to 3-mm follicles), and peak E2 concentrations occur when the follicles reach their maximum diameters (ie, end of the growth phase; [1,2]). In both groups of animals, the end of follicular growth phase for the final wave of the cycle was observed between days 14 and 15. There were no differences in mean numbers of large antral during that period, but treated ewes had more small antral follicles (3 mm in diameter) than controls. Therefore, the difference in serum E2 concentrations in experiment 1 was most likely due to the difference in total numbers of ovarian antral follicles or transiently increased follicular estrogenicity in P4-treated ewes. In the second experiment, there were no significant differences in the numbers of ovarian follicles or growth
Table 2
Comparisons of ovulation rates, estrous cycle duration, and follicle wave characteristics (penultimate and final wave of the interovulatory interval studied) between progesterone (P4)-treated ewes and their respective controls in experiment 1 (moderately prolific cross-bred ewes) and experiment 2 (highly prolific Olkuska sheep).

| Variables | Experiment 1 (Rideau Arcott × Polled Dorset) | | | Experiment 2 (Olkuska) |
|-----------|------------------------------------------------|-------------------|-------------------|
|           | Treatment (n = 8) | Control (n = 8) | Treatment (n = 7) | Control (n = 7) |
| Ovulation rate #1 | 2.8 ± 0.3 | 2.8 ± 0.2 | | 4.0 ± 1.0 |
| Ovulation rate #2 (post-treatment) | 1.6 ± 0.2a | 3.2 ± 0.4b | | 3.2 ± 0.8c |
| No. of luteal structures post-treatment | 1.5 ± 0.2a | 3.1 ± 0.3b | | 3.2 ± 0.8c |
| Interovulatory interval (d) | 17.3 ± 0.4 | 16.6 ± 0.1 | | 16.9 ± 0.6 |
| Penultimate wave | | | | |
| Day of emergence (Day 0 = first ovulation of the estrous cycle studied) | 9.6 ± 0.4 | 9.9 ± 0.3 | | 10.1 ± 0.3 |
| No. of follicles | 1.4 ± 0.2ac | 2.1 ± 0.3bc | | 3.2 ± 0.2 |
| % of ovulating follicles | 0.25 ± 0.16b | 1.75 ± 0.45bc | | 0.5 ± 0.3bc |
| Duration of the growth phase (d) | 2.9 ± 0.6 | 3.6 ± 0.3 | | 2.4 ± 0.2bc |
| Growth rate (mm/d) | 1.1 ± 0.2a | 0.78 ± 0.05bc | | 1.4 ± 0.2a |
| Duration of the static phase (d) | 2.0 ± 0.6 | 2.9 ± 0.4 | | 2.3 ± 0.4 |
| Duration of the regressing phase (d) | 2.0 ± 0.3 | 1.7 ± 0.2 | | 1.8 ± 0.6 |
| Regression rate (mm/d) | 1.2 ± 0.2 | 1.1 ± 0.1 | | 1.4 ± 0.2a |
| Maximum follicle diameter (MaxFD) (mm) | 5.7 ± 0.3b | 5.4 ± 0.1b | | 5.6 ± 0.2 |
| Mean day achieving MaxFD | 11.9 ± 0.6 | 13.6 ± 0.6 | | 12.5 ± 0.5 |
| Ovulatory follicle diameter (mm) | 6.5 ± 0.5bc | 5.3 ± 0.1b | | 6.0 ± 1.0 |
| Final wave | | | | |
| Day of emergence | 12.4 ± 0.4 | 12.6 ± 0.4 | | 11.8 ± 0.6 |
| No. of follicles | 1.5 ± 0.3 | 1.7 ± 0.3 | | 3.7 ± 0.6 |
| % of ovulating follicles | 95.9 ± 4.1 | 93.8 ± 6.3 | | 73.8 ± 9.6 |
| Duration of the growth phase (d) | 3.3 ± 0.4 | 2.5 ± 0.3 | | 2.7 ± 0.4 |
| Growth rate (mm/d) | 1.0 ± 0.1 | 1.1 ± 0.1 | | 1.3 ± 0.1bc |
| Duration of the static phase (d) | 2.7 ± 0.3 | 2.4 ± 0.3 | | 2.1 ± 0.1 |
| Duration of the regressing phase (d) | [2.0] | [2.0] | | [1.0, 1.0, 2.0] |
| Regression rate (mm/d) | [1.0] | [1.0] | | [2.0, 2.5, 1.0] |
| Maximum follicle diameter (MaxFD) (mm) | 5.8 ± 0.2 | 5.7 ± 0.2 | | 6.0 ± 0.3 |
| Mean day achieving MaxFD | 15.1 ± 0.3b | 14.8 ± 0.3b | | 14.7 ± 0.6 |
| Ovulatory follicle diameter (mm) | 5.8 ± 0.2 | 5.7 ± 0.3 | | 6.0 ± 0.3 |
| Inter-wave interval (d) | 3.1 ± 0.3 | 3.1 ± 0.3 | | 2.4 ± 0.7 |

Data are expressed as mean ± standard error of the mean. 

\[ \text{ap} < 0.05; \text{ABp} < 0.01. \]

\[ c \] Mann–Whitney rank-sum test.

kinetics of the last follicular wave of the cycle between the 2 groups of ewes; consequently, serum E2 concentrations were similar in both subsets of ewes although the treatment group exhibited more frequent peaks in mean E2 levels. The latter could be attributed to an occurrence of an additional preovulatory peak in blood E2 concentrations (detected on ~ day 15.5) in a majority of P4-treated Olkuska sheep. The specific reason for these alterations in E2 release before estrus and ovulations in the treated animals in experiment 2 remains uncertain.

In both experiments, the treatment with exogenous P4 significantly increased the growth rate of large antral follicles and reduced the number and percentage of ovulatory follicles emerging in the penultimate wave of the estrous cycle studied. However, the total number of follicles attaining ≥5 mm in size in the penultimate wave of the cycle was decreased in P4-treated Rideau Arcott × Polled Dorset ewes but not in Olkuska sheep. Albeit the difference was not statistically significant, due mainly to relatively high individual variability, P4-treated Olkuska ewes had on the average 1 follicle more in the final wave of the cycle than controls. Consequently, the mean ovulation rate was only decreased in moderately prolific cross-bred ewes receiving P4 injections at the end of diestrus. Clearly, differences do exist in the endocrine control of antral follicle development between highly and moderately prolific breeds of sheep. The latter observation is further supported by dissimilar effects of P4 treatment on the regression rate of large antral follicles (increased in P4-treated Olkuska sheep but not in cross-bred ewes) and maximum diameter of ovulating follicles from the penultimate wave (greater in P4-treated Rideau Arcott Polled Dorset sheep) as well as the number of 3-mm follicles post-treatment (increased in P4-treated Rideau Arcott × Polled Dorset sheep but not in Olkuska breed) as well as the number of 3-mm follicles post-treatment (increased in P4-treated Rideau Arcott × Polled Dorset sheep from days 15 to 17). The reasons for these differences remain equivocal. Apart from a rise in mean daily numbers of 3-mm follicles in experiment 1, there was no apparent effect of P4 treatment on antral follicle populations in the ewes of the present study.

The present results may pave the way to the development of simple, practical methods to control sheep fertility, using protocols that mimic a physiological luteal mechanism. Treatment with exogenous P4 can be employed to temporarily reduce the incidence of twinning or triplet pregnancies in moderately prolific strains of sheep; this would help control the incidence of twin-lamb disease.
(pregnancy toxemia) and underfed suckling lambs. Alternatively, manipulating circulating concentrations of P₄ using prostaglandins or oxytocin, with or without P₄-releasing pessaries, or application of P₄ antagonists could be used to improve fertility in nonprolific ewes. If administering P₄ late in the cycle reduces ovulation rates in moderately prolific breeds of sheep, it is very likely that reducing P₄ concentrations or blocking its effects at the end
of diestru may increase ovulation rates in nonprolific ewes. Future studies are required to determine the optimal timing and doses of exogenous hormones or chemicals.

In summary, the present study demonstrates that prolif er ewes treated with P4 at the end of diestru had a significantly lower percentage of follicles ovulating from the penultimate wave of the estrous cycle. Moreover, exogenous P4 given to moderately prolific R~deau Arcott × Polled Dorset ewes reduced the number of ovulatory-sized antral follicles in the wave in which the treatment was applied. As a result, the moderately prolific sheep receiving P4 injection on days 11 and 12 postovulation had a reduced ovulation rate compared with untreated controls. These observations can be interpreted to suggest that P4 is an important endocrine signal governing antral follicular development and ovulation rate in sheep, and that differences exist in the endocrine control of antral follicle development between prolific and nonprolific genotypes as well as between moderately and highly prolific strains of sheep. Our results also show that ovulation rates in cyclic ewes can be artificially manipulated. Finally, the present observations in cyclic ewes provide the basis for further investigations into whether similar biological mechanisms governing the ovulation rate are conserved among other polyovulatory species or if manipulating the luteal function can lead to multiple ovulations and twinning in mono-ovulatory species.

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