Use of Estradiol Cypionate as a Substitute for GnRH in Protocols for Synchronizing Ovulation in Dairy Cattle*

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ABSTRACT

Our purpose was to determine whether estradiol cypionate (ECP) could be substituted for the second GnRH injection of the standard Ovsynch protocol (injection of GnRH given 7 d before and 48 h after PGF2α, with timed AI [TAI] 12 to 20 h after the second GnRH injection). Lactating dairy cows ranging from 61 to 82 d in milk at TAI were studied in 14 replicates. Main effects were hormone (ECP vs. GnRH) to induce ovulation and exposure to progesterone (P4) or not during the week preceding PGF2α-induced luteolysis. Four treatments were: 1) 100 μg of GnRH at 48 h after PGF2α (Ovsynch; n = 27); 2) same as Ovsynch, plus a P4-releasing intravaginal insert (CIDR) placed for 7 d beginning at the first GnRH injection (Ovsynch + CIDR, n = 20); 3) same as Ovsynch, but substituting 1 mg of ECP for GnRH, and injecting ECP at 24 h after PGF2α (Heatsynch; n = 33); or 4) Heatsynch + CIDR (n = 26). The largest follicle was identified by ultrasonography 24 h after PGF2α and was monitored every 6 h until ovulation. Incidence of estrus was less after GnRH (54%) than after ECP (87%), but more GnRH-treated cows had LH surges detected (95 vs. 65%) and ovulated (100 vs. 86%). Duration of LH surges, but not peak concentrations, was less after GnRH than after ECP (6.1 ± 0.7 vs. 12.2 ± 0.9 h). Pre-treatment with P4 reduced the incidence of LH surges but had no effects on incidence of estrus or ovulation. Intervals to the LH surge and ovulation were less after GnRH than after ECP, but intervals between onset of the LH surge and ovulation did not differ (26 ± 2 vs. 30 ± 3 h). We concluded that substituting ECP for GnRH resulted in more cows in estrus and slightly fewer ovulating.

(Key words: estrogens, progesterone, GnRH, ovulation)

INTRODUCTION

Programmed breeding makes it feasible to schedule AI of dairy cattle during the first week that follows the end of the volunteer waiting period. The Ovsynch protocol (Pursley et al., 1995) synchronizes ovulation before timed artificial insemination (TAI) with pregnancy rates in the range of 30 to 40% (Burke et al., 1996; Pursley et al., 1997; Stevenson et al., 1999). Further work in Florida (Moreira et al., 2001) and in Kansas (Cartmill et al., 2001; El-Zarkouny et al., 2004) demonstrated that presynchronizing estrous cycles of cows to d 5 to 12 of the estrous cycle (Vasconcelos et al., 1999) by either 1 or 2 injections of PGF2α preceding the initiation of the Ovsynch protocol, significantly improved pregnancy rates, by 24 to 48% beyond what was achieved with Ovsynch alone.

Use of the Ovsynch protocol produces few cows in estrus because the peak in estrogen secreted by the preovulatory follicle is prematurely abrogated by the LH surge that is induced by the second GnRH injection after PGF2α-induced luteal regression. Uterine tone and presence of mucus, which are traditional validations that cows are truly in estrus, often are limited or absent at TAI. Cervical penetration and semen placement with the breeding catheter may be more difficult when cows have not been exposed to normal estrogen concentrations secreted by mature preovulatory follicles. As few
as 20% of Ovsynch-treated cows show estrus after the PGF2α injection associated with the Ovsynch protocol (Stevenson et al., 1996).

Substituting estrogen for the second GnRH injection is a logical choice for numerous reasons, including cost and the induction of normal estrual characteristics such as mucous secretion, uterine tone, and resulting sexual behavior. This progression of events more logically simulates a natural estrus when estrogen is given at the proper time after PGF2α-induced luteal regression. The only estrogen product available in the United States was estradiol cypionate (ECP), which has multiple label indications, including one “to correct anestrus [absence of heat period] in the absence of follicular cysts” at large doses of 3 to 5 mg. Esterification of estradiol-17β to produce ECP increases the half-life of the estrogen, with release of the parent estradiol following hydrolysis (Vynckier et al., 1990). Studies using a large dose (10 mg) of ECP resulted in a peak in plasma estradiol at 20 h that remained elevated for as long as 5 d (Vynckier et al., 1990). Administration of 2 mg of ECP was sufficient to cause ovarietomized heifers to display estrus (Lefebvre and Block, 1992).

Further, estradiol benzoate has been used to induce estrus and ovulation in lactating dairy cattle (Dailey et al., 1986). Estrogen potentiates the LH surge via actions on hypothalamic GnRH and the resulting LH surge during proestrus (Hansel and Convey, 1983) and enhances the release of GnRH in response to exogenous GnRH in the ovariectomized cow (Britt et al., 1981). Substituting ECP for GnRH in an Ovsynch-like protocol (known as Heatsynch) induced estrus, preovulatory LH surge, ovulation, and normal corpus luteum (CL) development in dairy heifers (Lopes et al., 2000), and induced estrus and ovulation in lactating dairy cows (Pancarci et al., 2002). Conception rates of ECP-treated heifers were not different from those in heifers inseminated after detected estrus (Lopes et al., 2000) and those in lactating cows after Heatsynch were similar to those after Ovsynch (Pancarci et al., 2002).

The current experiment was conducted to examine timing and variation of events critical to establishing conception (onset of estrus, LH surge, and ovulation) as useful references for future efforts to refine control of estrus and ovulation. Specifically, our objectives were to determine the incidences and characteristics of estrus, preovulatory LH surge, and ovulation in lactating dairy cattle after use of ECP as a substitute for GnRH in an ovulation synchronization protocol preceding a timed insemination.

**MATERIALS AND METHODS**

**Experimental Design**

Lactating Holstein cows (n = 119) were housed in either covered free stalls bedded with sand or in a tie-

![Figure 1](image-url) Experimental design illustrating 4 treatments in which ovulation was synchronized before timed AI (TAI). Blood was collected on d −10, −3, and −1 to determine concentrations of progesterone. In 3 replicates, blood was also collected at 0, 1, 2, 3, 4, 5, 6, 8, and 10 h after GnRH (expected LH peak at 2 h after GnRH) or at 24, 25, 26, 27, 28, 29, 30 h, then at 2-h intervals until 58 h after estradiol cypionate (ECP) (expected LH peak at 44 h after ECP) to determine incidence, timing, duration, and onset of the induced LH surge. Transrectal ultrasonography to determine incidence and timing of ovulation occurred at 6-h intervals starting 48 h after PGF2α. Onset of estrus was determined electronically by using HeatWatch in some replicates, but all cows were inseminated at one fixed time (TAI). CIDR = controlled internal drug-releasing intravaginal insert containing progesterone, P4 = progesterone.
GnRH, and injecting ECP at 24 h after PGF₂α (Heatsynch; n = 33); or 4) Heatsynch + CIDR (n = 26). Actual insemination times were 65 to 74 h after PGF₂α, (corresponding to 16 to 22 h after GnRH or 42 to 50 h after ECP). The rationale for using the CIDR inserts was to: 1) synchronize more closely proestrous events that were studied; and 2) determine any effects of P₄ on the timing of estrus, ovulation, and endocrine events studied.

Not every measurement could be assessed in each cow studied because of the necessity to have cows housed in either tie stalls or free stalls to accommodate blood collection or normal sexual behavior. Numbers of clusters and cows used for study were: timing of ovulation (11 clusters and 105 cows); timing of estral events (4 clusters and 34 cows); timing of the preovulatory LH surge (3 clusters and 42 cows); and visual detection for occurrence of estrus during the breeding week for cows housed in outside free stalls (10 clusters and 85 cows).

Blood Collection

Blood samples were collected from all cows before administering the first or only GnRH injection, before PGF₂α, and 24 h after PGF₂α, for later determination of concentrations of P₄. These samples were used to determine cycling status of cows and incidence of luteolysis after PGF₂α. Progesterone was measured in blood sera by radioimmunoassay (Skaggs et al., 1986). Inter- and intraassay CV were 9.6 and 7.9%, respectively.

In 3 replicates, blood was collected from cows housed in tie stalls at 0, 1, 2, 3, 4, 5, 6, 8, and 10 h after GnRH (expected LH peak at 2 h after GnRH) and at 24, 25, 26, 27, 28, 29, 30, 32, and 34 h, then at 2-h intervals until 58 h after ECP (expected LH peak at 44 h after ECP; Lopes et al., 2000) to determine incidence, timing, duration, and onset of the induced LH surge.

Serum LH concentrations were measured by a validated radioimmunoassay. Serum (200 μL) was incubated with 200 μL of anti-oLH (TEA #35; J. J. Reeves, Washington State University, Pullman) diluted 1:50,000 in assay buffer (0.1% gelatin, 0.01% thimerosal, 0.01 M PO₄, 0.9% NaCl, pH 7.2, with normal rabbit serum [1:300] and 0.01 M EDTA) at 4°C for 24 h. On d 2, 100 μL of assay buffer containing approximately 20,000 cpm [¹²⁵I]-oLH (LER 1374a; gift from L. E. Reichert, Albany Medical College, NY) were added, and the incubation continued for an additional 24 h at 4°C. Precipitation of antibody complexes began on d 3 with the addition of 100 μL of goat anti-rabbit antiserum (1:50 dilution in assay buffer). On d 6, 3 mL of PBS was added to each tube, and tubes were centrifuged at 3000 × g for 30 min at 4°C. Supernatant was decanted, and the pellet was counted for 1 min. Concentrations of LH in unknown samples were estimated from a standard curve (0.02, 0.04, 0.08, 0.16, 0.31, 0.63, 1.25, 2.50, 5.0, and 10 ng/tube) using bLH (USDA-bLH-B6). Increasing volumes of bovine plasma (50, 100, and 200 μL) resulted in a displacement curve that was parallel to the standard curve. The addition of different masses of bLH to the assay (0.25, 0.5, and 1.0 ng/tube) resulted in an average recovery of 101%. Intra- and interassay CV were 8 and 12%, respectively.

Ovarian Ultrasonography

Ovarian structures were monitored in 11 replicates of cows by using transrectal ultrasonography (real time, B-Mode, linear array, diagnostic, ultrasound scanner equipped with a 5-MHz transducer, Aloka 500V, Wallingford, CT). Diameter of the largest ovarian follicle identified 24 and 72 h after PGF₂α, was measured by using electronic calipers (average of vertical and horizontal measures) and then monitored at 6-h intervals until it disappeared (ovulation) or until 102 h after PGF₂α. Ovaries of cows in most replicates were reexamined by ultrasonography to validate ovulation and location of the CL on d 13 after TAI. Pregnancy was diagnosed 30 to 32 d after TAI by using ultrasonography (presence of uterine fluid and conceptus).

Estrous Behavior

In 4 replicates of cows housed in free stalls, HeatWatch devices (DDx, Inc., Denver, CO) were affixed to cows to monitor characteristics of estrus (duration, number of standing events, and total duration of standing time per estrus), and intervals from PGF₂α to estrus and ovulation. Estrual activity was monitored electronically, during and after treatments, until pregnancy diagnosis. Cows having at least 2 or more standing events, for at least 2 s each were defined to be in estrus; otherwise, those HeatWatch data were counted as false-positive events.

Statistical Analyses

Characteristics of estrus (incidence [all cows, including those fitted with detection devices] and traits associated with estrus [duration, number of standing events, and total standing time]) were analyzed by ANOVA (general linear models procedure; SAS Inst. Inc., Cary, NC) according to the following model: hormone (ECP vs. GnRH), CIDR (CIDR vs. no CIDR), lactation number (1 vs. 2+), all 2-way interactions among the previous independent variables, and season (fall, winter, and spring).
All characteristics of ovulation (incidence and intervals between PGF$_{2\alpha}$, ECP, and GnRH and ovulation) were analyzed as just described according to a similar model. Characteristics of the LH surge (incidence, duration, and peak) and intervals from PGF$_{2\alpha}$, ECP, and GnRH to onset of the LH surge were analyzed as just described without the season effect (these studies were conducted during a 35-d period). In instances for which limited numbers of observations were available for analyses (hours to estrus after PGF$_{2\alpha}$, ECP, and GnRH) and hours from estrus to ovulation, a reduced model was employed that excluded 2-way interactions of the main effects with lactation number. To determine differences in diameter of the largest preovulatory follicle (measured 24 h after PGF$_{2\alpha}$), the model only included CIDR, lactation number, their interaction, and season because this measurement occurred before administration of either ECP or GnRH after PGF$_{2\alpha}$-induced luteolysis. In fewer numbers of cows, the putative ovulatory follicle was measured 72 h after PGF$_{2\alpha}$, and the effect of GnRH or ECP and their interaction with CIDR was added to the previous model.

In all models, treatment differences were determined by resulting F-tests in the ANOVA, whereas all other differences were determined by using the least-significant difference test (PDIF test in SAS plus the Tukey adjustment) when protected by a significant ($P \leq 0.05$) F-test in the ANOVA.

RESULTS

Characteristics of Estrus

Estrus was detected in 34 of 44 (77.3%) cows. A greater ($P < 0.05$) proportion of cows were detected in estrus (also includes visually detected estrus in cows without electronic devices) after ECP than after GnRH during the week that followed the injection of PGF$_{2\alpha}$ (87% vs. 54%). Only 4 Ovsynch cows (2 with and 2 without previous P4 exposure) equipped with electronic estrus-detection devices were detected in estrus. The addition of the P4 via the CIDR insert had no effect on the incidence of estrus. Duration of estrus among Heatsynch cows tended ($P = 0.08$) to be greater in those previously treated with P4 via the CIDR insert (7.4 ± 1.3 h [n = 10]) than in those not given P4 (4.4 ± 1 h [n = 13]). The number of standing events per female in estrus (31 ± 11 vs. 11 ± 13) and total standing time (7.4 ± 6 vs. 13.5 ± 5 s) did not differ between Heatsynch cows treated with P4 and those without CIDR inserts, respectively.

Intervals from PGF$_{2\alpha}$ to onset of estrus were only numerically less ($P = 0.11$) in Heatsynch cows after previous exposure to P4 (46 ± 7 h) than in those not given P4 (60 ± 5 h). Heatsynch cows treated with P4 tended ($P = 0.06$) to have shorter intervals to estrus after ECP (19 ± 5 h) than did herdmates not given P4 (33 ± 4 h).

Follicle Diameters

Diameter of the ovulatory follicle 24 h after PGF$_{2\alpha}$ was greater ($P < 0.01$) in cows previously treated with P4 (14.6 ± 0.4 vs. 13.1 ± 0.4 mm). The difference tended to be greater, however, in first lactation (15.2 ± 0.8 [n = 14] vs. 12.5 ± 0.7 mm [n = 17]) than in multiple-lactation cows (14.0 ± 0.5 [n = 29] vs. 13.6 ± 0.5 mm [n = 41]; CIDR × lactation number interaction; $P = 0.06$). By 72 h after PGF$_{2\alpha}$ (24 h after GnRH and 48 h after ECP), no differences in diameter of the ovulatory follicle were detected.

Characteristics of Ovulation and Induced LH Surges

Mean responses and ranges in values for each characteristic are summarized for 42 cows studied to determine the onset of the preovulatory LH surge (Table 1). Proportionally more ($P < 0.05$) LH surges were detected during the sampling period in Ovsynch than in Heatsynch cows (95 vs. 65%). In contrast, among cows treated with a CIDR insert, fewer ($P < 0.05$) LH surges were detected (68 vs. 90%), particularly among Heatsynch cows previously treated with P4.

As expected, onset of the LH surge occurred earlier ($P < 0.001$) after administration of GnRH than of ECP (1 ± 2 vs. 35 ± 2 h). Among Heatsynch cows, however, onset occurred much sooner in those previously treated with P4 (CIDR × hormone interaction; $P < 0.05$). Average LH response curves among GnRH-treated cows treated with P4 were superimposable upon those of GnRH-treated cows not exposed previously with P4. Duration of the LH surge was greater ($P < 0.001$) after ECP than after GnRH (12.2 ± 0.9 vs. 6.1 ± 0.7 h), but the maximum concentration (peak) associated with the LH surge did not differ among treatments.

Intervals from ECP to ovulation were greater ($P < 0.001$) than those from GnRH to ovulation (59 ± 2 vs. 26 ± 2 h). But intervals from onset of the LH surge to ovulation were consistent among hormones and did not differ, whether induced by GnRH or ECP (26 ± 2 vs. 30 ± 2 h). Exposure of cows to P4 did not influence timing of ovulation after either hormone (GnRH vs. ECP) or influence onset of the induced LH surge (Table 1).

Incidences of ovulation among cows in which we attempted to detect the LH surge tended ($P = 0.09$) to be greater in Ovsynch than Heatsynch cows, with previous treatment with P4 having no effect (Table 1). Some discrepancy in the number of cows in each category in Table 1 is explained by the summary of outcomes listed.
Table 1. Timing and characteristics of preovulatory LH surges induced by either GnRH or estradiol cypionate (ECP) after Ovsynch or Heatsynch with or without previous exposure to progesterone via a progesterone-releasing intravaginal (CIDR) insert.1

<table>
<thead>
<tr>
<th>Item</th>
<th>GnRH (Ovsynch)</th>
<th>ECP (Heatsynch)</th>
<th>P-values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No CIDR</td>
<td>CIDR</td>
<td>No CIDR</td>
</tr>
<tr>
<td>Incidence of LH surge, %</td>
<td>100 (9)</td>
<td>90 (10)</td>
<td>82 (11)</td>
</tr>
<tr>
<td>Time from GnRH or ECP to LH surge, h</td>
<td>1 ± 2 (9)</td>
<td>1 ± 2 (9)</td>
<td>42 ± 2 (9)</td>
</tr>
<tr>
<td>(0–1)</td>
<td>0–1</td>
<td>11 ± 1 (9)</td>
<td>13 ± 1 (6)</td>
</tr>
<tr>
<td>Duration of LH surge, h</td>
<td>7 ± 1 (9)</td>
<td>6 ± 1 (9)</td>
<td>4–111</td>
</tr>
<tr>
<td>Peak magnitude of LH surge, ng/mL</td>
<td>4.8 ± 1.2 (9)</td>
<td>4.7 ± 1.2 (9)</td>
<td>3.3 ± 1.1 (9)</td>
</tr>
<tr>
<td>Incidence of ovulation, %</td>
<td>100 (9)</td>
<td>100 (10)</td>
<td>915 (11)</td>
</tr>
<tr>
<td>Time from GnRH or ECP to ovulation, h</td>
<td>28 ± 3 (9)</td>
<td>25 ± 3 (8)</td>
<td>61 ± 3 (7)</td>
</tr>
<tr>
<td>(19–321)</td>
<td>20–31</td>
<td>37–76</td>
<td>43–73</td>
</tr>
<tr>
<td>Time from LH surge to ovulation, h</td>
<td>26 ± 2 (9)</td>
<td>26 ± 3 (7)</td>
<td>29 ± 3 (6)</td>
</tr>
<tr>
<td>(18–322)</td>
<td>19–31</td>
<td>11–36</td>
<td>28–36</td>
</tr>
</tbody>
</table>

1Cows were administered GnRH 7 d before PGF2α. Ovsynch cows were administered GnRH 48 h after PGF2α, whereas Heatsynch cows were given ECP 24 h after PGF2α. Cows in both treatments either were exposed to progesterone via a CIDR insert (CIDR) for 7 d preceding PGF2α, or served as controls (no CIDR). A total of 42 cows were studied to determine these characteristics.

2LH surges were not detected in 9 cows during respective sampling periods (additional details in footnotes below).

3Range in values.

4In 1 cow in which an LH surge was not detected, premature luteolysis occurred before CIDR removal and early ovulation occurred before GnRH was administered.

5In 2 cows in which an LH surge was not detected, 1 cow ovulated early (37 h after ECP) and the other cow did not ovulate during the observational period probably because of delayed luteolysis (serum P4 = 3.1 ng/mL 24 h after PGF2α).

6In 6 cows in which an LH surge was not detected, 3 cows ovulated early (43, 50, and 64 h after ECP), 1 cow did not ovulate during the observational period (4 h after ECP), and 2 cows had either a CL or cyst detected 13 d after TAI.

Below and in footnotes to Table 1. Concentrations of P4 on the day of PGF2α, and 24 h later, ovarian scans to detect ovulation beginning 48 h after PGF2α, and ovarian scans in most cows 13 d after TAI were used to account for these outcomes in each cow sampled. As indicated in Table 1, of all cows in which the LH surge was detected, only 1 cow failed to ovulate in the ECP + CIDR treatment. For at least 5 of the 6 cows that ovulated in which no LH surge was detected, the LH surge likely occurred before the blood sampling began (1 cow ovulated before GnRH and 4 cows ovulated after ECP during the early scanning period). The sixth cow had delayed luteolysis, and the LH surge likely occurred after the sampling period ended. Of the 2 cows in which neither ovulation nor the LH surge was detected, 1 had a CL on d 13 after TAI and likely ovulated late and the second failed to ovulate (a cystic structure was detected on d 13 after TAI).

Of the 98 cows that ovulated during the study (regardless of whether they were sampled to detect the LH surge), 11.2% ovulated less than 48 h after PGF2α (before the first ovarian scans that continued every 6 h), and 7.1% ovulated more than 102 h after PGF2α (based on ovarian scans made on d 13 after TAI). As expected, concentrations of P4 at the time of PGF2α injection were less (P < 0.01) in those cows having early ovulation (1.3 ± 0.7 ng/mL) than in cows that ovulated between 48 and 102 h after PGF2α (4.0 ± 0.3 ng/mL) and those ovulating thereafter (4.5 ± 0.9 ng/mL). Of the 8 cows that failed to ovulate during the study, 6 cows had elevated P4 (> 1 ng/mL) at 24 h after PGF2α, although concentrations of P4 had declined in all 8 cows by >50% by 24 h after PGF2α.

Of the various characteristics associated with ovulation (Table 2) among cows in which blood was not sampled to detect LH surges, none was influenced by previous exposure of P4. This was consistent with findings for cows in which we attempted to detect LH surges (Table 1). Frequency of ovulation was greater (P < 0.05) in cows treated with GnRH than in those treated with ECP (100 vs. 86%). Intervals to ovulation after the onset of estrus were not different among Heatsynch cows, regardless of previous P4 exposure. Intervals from PGF2α to ovulation were greater (P < 0.001) in ECP-treated than GnRH-treated cows (86 ± 2 vs. 77 ± 3 h).

Pregnancy rates after Heatsynch were 28.4% (n = 67) and after Ovsynch were 27% (n = 52). Those after treatment with the CIDR were 30.6% (n = 72) and no CIDR were 21.4% (n = 47).

**DISCUSSION**

Although few cows generally are detected in estrus during the breeding week after the PGF2α injection of the standard Ovsynch protocol (Stevenson et al., 1996), just more than 50% were detected in estrus in the cur-
Table 2. Incidence of ovulation and intervals to ovulation after estrus or after injections of PGF$_{2_a}$, GnRH, and estradiol cypionate (ECP) in all cows not sampled to determine preovulatory LH surges.$^1$

<table>
<thead>
<tr>
<th>Item</th>
<th>No CIDR</th>
<th>CIDR</th>
<th>No CIDR</th>
<th>CIDR</th>
<th>P-values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incidence of ovulation, %</td>
<td>100 (18)</td>
<td>100 (11)</td>
<td>91 (22)</td>
<td>79 (14)</td>
<td>0.03</td>
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<td></td>
<td>21 (1)</td>
<td>21 (2)</td>
<td>28 ± 3 (8)</td>
<td>30 ± 3 (7)</td>
<td>0.52</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>24–34$^3$</td>
<td>24–45</td>
<td></td>
</tr>
<tr>
<td>Time from estrus to ovulation, h</td>
<td>27 ± 3 (17)</td>
<td>29 ± 4 (11)</td>
<td>64 ± 3 (20)</td>
<td>61 ± 4 (11)</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td>15–35$^3$</td>
<td>21–34</td>
<td>42–99</td>
<td>24–45</td>
<td></td>
</tr>
<tr>
<td>Time from PGF$_{2_a}$ to ovulation, h</td>
<td>76 ± 2 (17)</td>
<td>74 ± 3 (11)</td>
<td>88 ± 2 (20)</td>
<td>84 ± 3 (11)</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td>64–85$^3$</td>
<td>69–84</td>
<td>66–123</td>
<td>65–115</td>
<td>0.52</td>
</tr>
</tbody>
</table>

$^1$Cows were administered GnRH 7 d before PGF$_{2_a}$. Ovsynch cows were administered GnRH 48 h after PGF$_{2_a}$, whereas Heatsynch cows were given ECP 24 h after PGF$_{2_a}$. Cows in both treatments either were exposed to progesterone via a CIDR insert (CIDR) for 7 d preceding PGF$_{2_a}$, or served as controls (no CIDR).

$^3$Range in values.

It is possible that our collection period was not entirely appropriate for all cows relative to hormonal milieu and timing of luteolysis. In cows previously treated with P4, it is clear that proportions of LH surges detected were reduced compared with those not treated with P4. Despite fewer detected LH surges in ECP-treated cows, durations of their LH surges were about twice as long as those detected after GnRH, suggesting that the LH surge mimics characteristics of the natural LH surge more closely in proestrous cows treated with estrogen (Chenault et al., 1975), which is consistent with observations in heifers (Lopes et al., 2000).

Incidence of ovulation was greater after GnRH than after ECP in the present study, in which ovulation was monitored between 48 and 102 h after PGF$_{2_a}$. Equal frequencies of early (<24 h after ECP or <24 h after GnRH) and late (>78 h after ECP or >54 h after GnRH)
ovulation occurred independent of main effects. Intervals from ECP to estrus, to onset of the LH surge, and to ovulation were greater than those after GnRH, but intervals from either hormone to the onset of those events were proportional. More importantly, interval from the LH surge to ovulation was about 26 to 32 h after either hormone. Intervals to ovulation are consistent with other studies in which intervals were monitored in cows subject to the Ovsynch protocol (28 to 30 h between GnRH and ovulation; Pursley et al., 1995) or in cows spontaneously coming to estrus or in response to PGF$_2$$_\alpha$ (28 h between onset of estrus and ovulation; Dransfield et al., 1998). Intervals to ovulation after ECP in the current study (60 to 65 h) corroborate results in replacement heifers (Lopes et al., 2000), in which ovulation occurred an average 60 to 62 h after ECP.

Because of the difficulty of determining onset of estrus and onset of the LH surge in the same cows, we could not determine whether those 2 events were concurrent. Although average intervals from estrus to ovulation were about 5 h less in GnRH-treated cows than average intervals from the onset of the LH surge to ovulation, those in ECP-treated cows were nearly simultaneous, at 31 to 34 h and 29 to 32 h, respectively. It seems that estrus and onset of the LH surge are nearly simultaneous after ECP and are not too different from what occurs in untreated cows that spontaneously come to estrus (Chenault et al., 1975) or in those in which estrus is induced by PGF$_2$$_\alpha$ (Chenault et al., 1976).

Field trials in Texas and Florida designed to compare pregnancy rates after the Ovsynch and Heatsynch protocols reported no differences in fertility (Pancarci et al., 2002). Our unpublished studies confirm the previous report. In a concurrent study to the one reported herein, in which cows in the same herd were diagnosed not pregnant and then assigned to either the Ovsynch or Heatsynch protocols (without CIDR inserts), pregnancy rates did not differ (64/219 or 29.2% vs. 63/229 or 27.5%; Stevenson and Tiffany, unpublished data). In another experiment (Blevins et al., 2002) conducted concurrently with the present report, in which 141 cows were assigned to the Heatsynch protocol for insemination at their first service, pregnancy rates averaged 40% when assessed at 30 to 35 d after TAI.

Substituting estrogen for the second GnRH injection in an Ovsynch protocol is a logical choice because of cost. Cost of an ovulatory dose of estrogen, where market available, is less than 10% of the cost of an ovulatory dose of GnRH (100 μg). Unfortunately, since the present study was completed, ECP has been withdrawn from the U.S. market. Other positive benefits of an estrogen include induction of normal estrual characteristics, such as mucous secretion, uterine tone, and resulting sexual behavior. These traits are positive indicators of estrus for inseminators because these characteristics validate the likelihood that the cow is in estrus (Pancarci et al., 2002).

CONCLUSIONS

Substituting ECP for GnRH resulted in more cows in estrus after PGF$_2$$_\alpha$, than cows treated with the standard Ovsynch protocol. Although we have somewhat limited information, no evidence exists to suggest that injection of ECP (1 mg) produced characteristics of estrus that were outside of normal physiological ranges. Of those cows monitored for incidence of ovulation, fewer ovulated after ECP than after GnRH. Pretreatment with P4 had no effects on the incidence of estrus or ovulation, but fewer cows had a detected LH surge. Intervals from estrus to ovulation and from onset of the LH surge to ovulation were similar after either GnRH or ECP, suggesting that onset of estrus and the LH surge are nearly concurrent events. Pregnancy rates to timed AI after ECP in the current trial and others are similar to those to timed AI after GnRH.

ACKNOWLEDGMENTS

We express appreciation to: Pharmacia Animal Health (Kalamazoo, MI) for donation of Lutalyse and ECP used in these studies; Merial (Iselin, NJ) for supplying the Cystorelin; InterAg (Hamilton, NZ) for donation of the CIDR inserts; L. E. Reichert, Albany Medical College, NY, for donation of oLH for iodination; J. J. Reeves, Washington State University for donation of LH antiserum; and Betty Hensley for assistance in conducting laboratory work.

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