Controlling the dominant follicle in beef cattle
to improve estrous synchronization and early
embryonic development

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Estrous synchronization and timed ovulation programs that permit AI at a
predetermined time (timed AI) rather than as determined by detection of
spontaneous estrus are requisite for increased adoption of AI in the beef
cattle industry. In the past two decades, significant progress has been
achieved in developing programs that synchronize ovulation to address this
need. While this progress has been driven by a multitude of fundamental
discoveries in reproductive biology, the greatest impact in the past two
decades has been the result of enhanced understanding of the pattern
of ovarian follicle growth in cattle and development of technologies to
coordinate growth and ovulation of the dominant follicle. At present,
estrous synchronization programs that result in acceptable timed AI
pregnancy rates are available for beef cattle. The capacity to control growth
of the dominant follicle and evaluate the impact of various approaches on
fertility has resulted in greater understanding of the factors that influence
maturity of ovulatory follicles. Modifications to the standard industry
breeding programs, with the aim of lengthening and/or increasing the
gonadotropin stimulus and estradiol production by preovulatory follicles,
have been shown to substantially increase timed AI pregnancy rate in beef
cattle. Associations between characteristics of follicular development
and fertility have surfaced from application of estrous and ovulation
synchronization technologies and led to investigation of the fundamental
mechanisms that underlie these relationships.

Introduction

Effective estrous synchronization technologies to facilitate the use of timed AI in beef cattle
have been developed (Larson et al., 2006; Sa Filho et al., 2009) and modifications that enhance
their efficacy (Bridges et al., 2009; Meneghetti et al., 2009) continue to be discovered (Figure
1). The current technologies reflect the culmination of relevant fundamental discoveries in
reproductive physiology that date to the early 1900s, and research in estrous synchronization
that has been ongoing for over 60 years. The rapid progress in the field of estrous synchro-
nization and timed AI over the last two decades can be directly linked to the discovery of a
wave-like pattern of follicular development in cattle (Savio et al., 1988; Sirois and Fortune,
Since that time, research has progressed through investigation of the implications of not controlling follicular development, approaches to coordinate follicle development and application of methodology to control growth and ovulation of the dominant follicle in conjunction with other components of estrous synchronization and timed AI programs. The synchronization programs currently used in cattle reflect only a portion of the valuable knowledge gained through this area of research. Through attempts to coordinate follicular development and standardize the timing of ovulation, greater understanding of the impact of variation in follicular development on fertility in cattle has evolved. Various relationships between fertility in cattle and aspects of follicular development, such as follicle size, length of proestrus, follicular estradiol production, and progesterone concentrations during follicular development to fertility in cattle have emerged, and led to fundamental investigations of the mechanisms that underlie these associations.

Fig. 1. Common estrous synchronization programs used with timed Al in postpartum beef cows. GnRH = gonadotropin releasing hormone; PGF<sub>2α</sub> = prostaglandin F<sub>2α</sub>; CIDR = Controlled Internal Drug Release Insert; EB = estradiol benzoate; ECP = estradiol cypionate; TAI = timed AI. With the GnRH-based programs in the USA, the CIDR is the only available progestin source. With Estradiol + Progestin systems, varying forms of estradiol are used at the start of the progestin treatment. Also timing of PGF<sub>2α</sub> form and timing of estradiol used to synchronize ovulation, interval to timed AI and source of progestin vary among programs.
Traditional approaches to estrous cycle control in cattle

Early attempts to synchronize estrus in cattle were based on the knowledge that progesterone prevents the occurrence of estrus and ovulation (Christian and Casida, 1948). A variety of progestins were administered in an assortment of methods to attempt to coordinate the timing of estrus in cattle (Trimberger and Hansel, 1955; Zimbelman and Smith, 1966). Progestin treatments of relatively long duration resulted in the most precise timing of estrus since this allowed for spontaneous occurrence of luteolysis during progestin treatment, but fertility was substantially less than would be expected following a spontaneous estrus (Zimbelman et al., 1970; Wishart, 1977). The inverse relationship between duration of progestin treatment and fertility led to development of short-term progestin treatments.

One approach to shortening the duration of progestin treatment was administration of a pharmacological dose of estrogen at the initiation of treatment to induce luteal regression; rather than relying on long-term treatments to allow spontaneous luteolysis (Wiltbank and Kasson, 1968; Mauleon, 1974). Synchrony of estrus and fertility varied with these approaches since the response to estrogen varied by stage of the estrous cycle (Lemon, 1975; Miksch et al., 1978). Development of short-term programs was also aided by identification of PGF₁₀ as a luteolysin (McCracken et al., 1972) and characterization of its efficacy to induce luteolysis (Lauderdale, 1975). However, the timing of estrus after utilizing this approach continued to vary (Wishart, 1974; Thimonier et al., 1975) due to stage of the estrous cycle at which luteal regression was induced (King et al., 1982), and fertility varied due to the stage of the estrous cycle when progestin treatment was initiated (Beal et al., 1988). Although usable estrous synchronization programs existed at this time, the efficacy of most programs was not predictable (Odde, 1990, review).

Coordinating follicular growth for estrous synchronization

Persistent ovarian follicles

Characterization of the wave-like pattern of follicular growth in cattle through the use of ovarian ultrasonography (Savio et al., 1988; Sirois and Fortune, 1988; Ginther et al., 1989) provided insight into challenges associated with earlier approaches to synchronize estrus. Although it had been known for some time that ovarian follicular development was not a random event in cattle (Rajakoski, 1960), controlling this aspect of ovarian function was not an intentional component of most programs. With new insight on the normal pattern of follicular growth and ultrasonographic capabilities, it was demonstrated that administration of a progestin in the absence of a corpus luteum extended the lifespan of the dominant ovarian follicle until the progestin was withdrawn (Sirois and Fortune, 1990). Dominant ovarian follicles with an extended lifespan ('persistent follicles') were shown to develop when practically any type of progestin was administered in the absence of a corpus luteum (Stock and Fortune, 1993; Anderson and Day, 1994), resulting in increased frequency of LH pulses and concentrations of estradiol (Kinder et al., 1996, review). Fertility was reduced if the lifespan of a dominant follicle was prolonged for four days (Mihm et al., 1994) and while fertilization of oocytes was unaffected, the embryo died shortly thereafter (Ahmad et al., 1995). Since the need of a progestin for synchronization of estrus in beef cattle has been well documented (Day, 2004, review), subsequent research focused on preventing ovulation of persistent follicles at the synchronized estrus.
Coordinating follicular development through management of persistent follicles

The discovery that atresia of persistent follicles could be induced by acutely increasing peripheral progesterone concentrations (Stock and Fortune, 1993; Savio et al., 1993), resulting in synchronous emergence of a new follicular wave 3.5 later (Anderson and Day, 1994) provided a potential tool that could be used to manipulate follicular development for estrous synchronization and avoid the infertility inherent to ovulation of persistent follicles. McDowell et al. (1998a) demonstrated that 24 hours of exposure to mid-luteal progesterone concentrations was sufficient to induce atresia of persistent follicles.

The concept of using progesterone delivered via an injection to synchronize atresia of persistent ovarian follicles and coordinate follicular development in herds of cattle was evaluated in a series of experiments (Anderson and Day, 1994, 1998; McDowell et al., 1998b). Experiments varied by progestin source and duration and the use of PGF2α to induce luteal regression. Conclusions from this series of experiments were that induced atresia of persistent follicles with progesterone resulted in a synchronous and fertile estrus and provided a means to coordinate follicular development in cows at random stages of the estrous cycle and in anestrous cows. A limitation of this approach for estrous synchronization was duration of treatment (≥ 14 days) necessary to ensure that most females initially developed persistent follicles. Synchronization programs that exceed 10 days are not acceptable to most producers. Furthermore, injectable progesterone is not commercially available in the USA.

Infertility associated with long term progestin use and development of persistent follicles can be avoided if cattle are not inseminated at the first estrus after progestin withdrawal but are inseminated at a subsequent synchronized estrus. Initial investigation of this approach involved feeding of an oral progestin for 14 days and induction of luteolysis with PGF2α 17 days later (Brown et al., 1988). This presynchronization with a progestin takes advantage of the beneficial effects of progestins and passively coordinates follicular development since a majority of females are in a similar stage of the estrous cycle at the time of PGF2α-induced luteal regression. This approach has been further refined (Lamb et al., 2000; Mallory et al., 2010) and effectively synchronizes estrus in cattle. The interval from initiation of progestin treatment to AI with this approach ranges from 33 to 36 days.

Coordination of follicular development with estradiol

An alternative to managing persistent follicles to coordinate growth of the dominant follicle would be to 'reset' follicular development at the outset of an estrous synchronization program and avoid development of persistent follicles altogether. Early evidence that estrogens may function in this manner was that injection of estradiol valerate induced atresia of large ovarian follicles (Engelhardt et al., 1989). It was subsequently demonstrated that injection of estradiol 17β induced atresia of the dominant follicle of the first follicular wave resulting in emergence of a new follicular wave approximately four days later in all females (Bo et al., 1995) thus providing a valuable tool to control growth of the dominant follicle for estrous synchronization. Several studies have focused on understanding the variation and mechanisms associated with the use of estradiol to reset follicular development using estradiol benzoate (EB; see Day and Burke, 2002 for review). Intramuscular injection of EB consistently induced atresia of dominant follicles and resulted in emergence of a new follicular wave (Burke et al., 2000, 2001), interval from EB to emergence of the new follicular wave increased with dose and EB was effective across stages of the estrous cycle (Day and Burke, 2002). The conception rate of cows administered EB at the initiation of progestin treatment was similar to that of cows exhibiting a spontaneous estrus (Day et al., 2000). Further, it was demonstrated that sustained
Concentrations of progesterone, for a minimum of 48 h, were necessary for EB to consistently reset follicular development (Burke et al., 1998).

In intact cows, administration of EB during the luteal phase suppressed LH and/or FSH in the periphery and induced an immediate decline in estradiol production by dominant follicles through suppression of aromatase (Burke et al., 2005; 2007). Timing of emergence of the new wave of follicular development was dependent upon the duration of FSH suppression by the exogenously administered estradiol and was dose dependent (Burke et al., 2003). These findings, and others (not cited) clearly established that the administration of exogenous estradiol represents a predictable and highly effective method to coordinate growth of the dominant follicle for estrous synchronization and timed AI in cattle.

**Coordination of follicular development with GnRH**

A second method to reset follicular development at the outset of an estrous synchronization program is the use of GnRH or its agonists. Injection of GnRH induces an LH surge within 2 hours (Kaltenbach et al., 1974) and, within 1 to 2 days, there is a reduction in the number of large follicles (Thatcher et al., 1989) due to ovulation and subsequent formation of a new corpus luteum (Twagiramungu et al., 1994). Synchronized emergence of a new follicular wave after GnRH (Macmillan and Thatcher, 1991) occurred within 1 to 2 days after GnRH injection (Twagiramungu et al., 1994, 1995).

A challenge of using GnRH for estrous synchronization is that ovulation in response to the first GnRH is induced in only 66% of beef cows (Geary et al., 2000). In heifers, response rates approaching 50% have been reported (Pursley et al., 1995). Variation in response to GnRH has been attributed to day of the estrous cycle (Moreira et al., 2000) and the influence of day of the estrous cycle on ovulatory response and growth of newly emerged dominant follicles in cows and heifers has recently been investigated (Atkins et al., 2008, 2010). Estrous synchronization programs that use GnRH should be structured to minimize the negative impact of uncoordinated follicular development that occurs in 20 to 50% of females treated.

**Estrous synchronization programs for timed AI in beef cows**

The capacity to coordinate growth of the dominant follicle stimulated a period of intense investigation that led to development of programs currently used for estrous synchronization in beef cattle. The importance of these contributions is recognized, but the focus here is on the current programs used for timed AI in beef cows, their benefits and shortcomings, and ongoing research to improve the current methodology.

**Strategy of current timed AI programs**

In beef cows, timed AI programs include a progestin source to suppress estrus and for induction of estrous cycles in anestrous females, and PGF_2_ to induce luteal regression (Figure 1). In the USA, GnRH or its agonist is used to reset follicular development at the initiation of the synchronization program and at the end of the program to synchronize ovulation for timed AI. In many other countries estradiol (either 17β-estradiol or estradiol benzoate) is used in timed-AI to reset follicular development and estradiol (including estradiol cypionate) is again used at the end of the program to synchronize ovulation.
Regardless of whether programs are GnRH- or estradiol-based, the rationale for the design of current programs is similar. Treatments begin with insertion of an intravaginal progestin device that is accompanied by GnRH or estradiol to reset follicular development. Emergence of a new follicular wave occurs 1 to 2 (GnRH) or 3 to 4 (estradiol) days later. When GnRH is used, emergence of the new follicular wave is accompanied by formation of the GnRH-induced accessory corpus luteum. Due to the difference in time of emergence between GnRH and estradiol, the progestin is withdrawn and PGF$_{2\alpha}$ is given either 7 (GnRH) or 8 to 9 (estradiol) days later. Hence, progesterone is withdrawn 5 to 6 days after emergence of a dominant follicle. The final step is to synchronize ovulation with either GnRH, given 60-66 hours after PGF$_{2\alpha}$, estradiol cypionate which is given at progestin withdrawal, or 17β-estradiol/EB given 24 to 48 hours after PGF$_{2\alpha}$.

Estrous synchronization and timed AI with GnRH

The standard GnRH-based timed AI program used in the USA is referred to as the “CO-Synch + CIDR” program (Figure 1). Some of the key research that led to development of this program (Macmillan and Thatcher, 1991; Twagiramungu et al., 1995; Pursely et al., 1995; Geary et al., 2001; Lamb et al., 2001) culminated in a large multistate project (Larson et al., 2006) which reported that timed-AI pregnancy rate with the CO-Synch + CIDR program was 54%. This program is a widely used estrous synchronization system for timed AI in beef cows in the USA. While the CO-Synch + CIDR program has been successful and yielded acceptable timed-AI pregnancy rates, limitations to fertility have been linked to the lack of coordination of follicular development in some females. Failure of the first GnRH to reset follicular development leads to aberrant follicular dynamics during the synchronization program (Geary et al., 2000) and the induced ovulation of small follicles in some animals by the second GnRH (Perry et al., 2005). Cattle induced to ovulate follicles smaller than observed at a spontaneous estrus with the second GnRH have decreased fertility (Lamb et al., 2001; Perry et al., 2005). The capacity of the first GnRH to reset follicular development as well as stage of the estrous cycle at treatment has been linked to size of the dominant follicle at the time of the second GnRH (Atkins et al., 2008, 2010). Hence, pregnancy rate to timed AI with the standard CO-Synch + CIDR program is limited by induced ovulation of compromised follicles.

Follicular maturity and fertility in cattle

The influence of ovulatory follicle maturity on fertility in beef cattle has been investigated (Mussard et al., 2003; 2007; Bridges et al., 2010; Perry et al., 2005). One hypothesis was that diameter of ovulatory follicles was the most appropriate indicator of follicle “maturity” and that cows induced to ovulate small follicles would have decreased fertility compared to cows which are induced to ovulate large dominant follicles. Within each of three experiments (Table 1; Mussard et al., 2003; 2007) this hypothesis was supported, but as data accumulated from multiple experiments, the relationship of follicle diameter to pregnancy rate appeared inconsistent. In fact, across experiments, the more consistent predictor of pregnancy rate appeared to be the interval from initiation of luteal regression with PGF$_{2\alpha}$ to either a spontaneous or GnRH-induced LH surge (Table 1). Since the stage of a spontaneous estrous cycle between luteal regression and initiation of estrus and the LH surge is typically referred to as proestrus, we, in turn, refer to the interval from PGF$_{2\alpha}$ (and progestin withdrawal when applicable) to GnRH treatment as ‘proestrus’ in our research. The observed relationship between length of proestrus and conception rate led to an additional experiment (Table 1; Bridges et al., 2010) designed
to hold follicle diameter constant and only vary length of proestrus. It was demonstrated that at a constant ovulatory follicle diameter, length of proestrus had a substantial influence on conception rate. Taken together, data from this series of studies suggested a strong positive relationship of duration of proestrus with follicle maturity and fertility, and furthermore, suggested that diameter of the ovulatory follicle, in itself, was not a consistent predictor of follicle maturity. The effect of ovulatory follicle size at GnRH-induced ovulation or at spontaneous ovulation on conception rate has also been evaluated by Perry et al. (2005, 2007). It was reported that diameter of the ovulatory follicle influenced conception rate after detection of estrus in heifers, but not in postpartum cows. In postpartum cows that did not exhibit estrus, diameter of the ovulatory follicle was positively associated with conception rate when ovulation was induced with GnRH. Thus, if a ‘complete’ spontaneous proestrus occurred in cows (confirmed by exhibition of estrus), diameter of the ovulatory follicle did not impact fertility. The impact of follicle diameter on conception rate was evident when ovulation was induced with GnRH; at a constant duration of proestrus. Since findings suggested that maturity of the ovulatory follicle and probability of conception is perhaps best defined by length of proestrus, we applied this knowledge towards optimizing the existing CO-Synch + CIDR program to take advantage of this relationship.

Table 1. Conception rate, diameter and age of the ovulatory follicle, duration of proestrus, and number of cows included from a series of experiments investigating the effect of follicle maturity on fertility.

| Conception rate (%) | Follicle diameter at ovulation (mm) | Duration of proestrus (days) | n | Experiment |
|---------------------|------------------------------------|----------------------------|---|------------|
| 4                   | 11.1 ± 0.2                         | 1.0 ± 0.1                  | 45 | Mussard et al., 2003a |
| 8                   | 11.1 ± 0.2                         | 1.0 ± 0.1                  | 12 | Mussard et al., 2003b |
| 10                  | 12.6 ± 0.2                         | 1.25                       | 10 | Bridges et al., 2010a |
| 57                  | 13.6 ± 0.2                         | 2.2 ± 0.1                  | 54 | Mussard et al., 2003a |
| 67                  | 13.7 ± 0.2                         | 2.0 ± 0.1                  | 12 | Mussard et al., 2003b |
| 71                  | 12.9 ± 0.2                         | 2.25                       | 28 | Bridges et al., 2010a |
| 76                  | 10.7 ± 0.1                         | 3.3 ± 0.1                  | 29 | Mussard et al., 2007a |
| 100                 | 12.0 ± 0.3                         | 4.7 ± 0.2                  | 24 | Mussard et al., 2007a |

* Percentage of animals determined to be pregnant following insemination. Pregnancy determination was conducted via ultrasonography at approximately 30 days post-insemination.
* Diameter of the largest ovulatory follicle as determined by ultrasonography conducted either at GnRH administration or estrus.
* Interval from PGF to GnRH administration.
* Cows were either induced with GnRH to ovulate a small (~11 mm) follicle or allowed to spontaneously exhibit estrus. Cows were inseminated 12 hours following estrus or GnRH.
* Cows were induced to ovulate either a small (~11 mm) or large (~13 mm) ovarian follicle with GnRH. Animals were inseminated 12 h following GnRH administration.
* Cows were induced to ovulate either a small (~11 mm) or large (~13 mm) ovarian follicle with GnRH. Embryo from non-treated cows were then transferred 7 days after GnRH. Cows were induced to ovulate an ovarian follicle of similar diameter with GnRH either 1.25 or 2.25 days following PGF administration. Animals were inseminated 12 h following GnRH administration. Includes only cows with a luteal phase of normal length.

Lengthening proestrus in the CO-Synch + CIDR program

The length of proestrus (interval from PGF to second GnRH and timed AI) with the traditional 7-day CO-Synch + CIDR program was varied from 56 to 72 hours in mature cows without
influencing timed AI pregnancy rate, but in younger cows (≤ 3 years of age), greatest pregnancy rates were achieved with timed AI at 56 hours (Dobbins et al., 2009). Others (Busch et al., 2008) have reported that timed AI pregnancy rates were greater when proestrus was 66 than 54 hours. In practice, the second GnRH is given and timed AI is performed in most herds between 54 and 66 hours after PGF2α, and there is no evidence that extending this interval beyond 66 hours will increase timed AI pregnancy rate. It was theorized that if the interval from the first GnRH and insertion of the CIDR to PGF2α was shortened to 5 days, proestrus could be lengthened and timed-AI pregnancy rate would increase. In the 5-day CO-Synch + CIDR program (Figure 1), two rather than one injection of PGF2α are administered at CIDR withdrawal, which is 5 days after CIDR insertion and the first GnRH injection. With this approach, the new follicular wave emerges 3 to 4 days before PGF2α is given, which is earlier relative to emergence of the ovulatory follicle (5 to 6 days) in the 7-day CO-Synch + CIDR program. Thus, the dominant follicle enters proestrus at a follicular age when follicular fluid concentrations and capacity of granulosa cells to produce estradiol are greater than with the 7 day program (Valdez et al., 2005). Bridges et al. (2008) determined that the appropriate interval from PGF2α to the second GnRH and timed-AI with the 5 day program was 72 hours. In comparison with the traditional 7 day CO-Synch + CIDR program, the 5-day CO-Synch + CIDR program increased timed-AI pregnancy rate from 59.9 to 70.4% (Bridges et al., 2008). Similarly, a 14% increase in timed-AI pregnancy rate was detected in yearling heifers with the 5-day as compared to 7-day program (Wilson et al., 2007). This substantial increase in timed-AI pregnancy rate with the 5-day program has been achieved using two doses of PGF2α (or cloprostenol sodium, CLP) spaced at 12 hours apart. Pregnancy rate is decreased with a single dose of PGF2α or CLP (Kasimanickam et al., 2009; Souto et al., 2009) due to failure of a single dose to induce luteal regression in approximately 1/3 of cows (Souto et al., 2009). Cruppe et al. (2010) recently reported that giving two simultaneous doses of PGF2α together at the time of CIDR withdrawal in the 5-day program resulted in pregnancy rates similar to those achieved with two injections, spaced 8 hours apart. Collectively, experiments that have included over 1700 cows across 23 herds and four years, have achieved a mean pregnancy rate of 68.4% with the 5-day CO-Synch + CIDR program using two doses of PGF2α given simultaneously or spaced 2, 7, 8 or 12 hours apart, depending upon individual experimental objectives.

Hormonal impact of lengthened proestrus

Proestrus starts with the diappearance of progesterone and terminates with either a spontaneous or GnRH-induced LH surge. An immediate response to declining progesterone is an increase in the frequency of LH pulses (Kinder et al., 1996), providing the primary stimulus for final development of preovulatory follicles (Ireland and Roche, 1983) and the preovulatory increase in estradiol (Kaneko et al., 1991). Concentrations of estradiol, magnitude of the LH surge, and progesterone concentrations during the ensuing luteal phase were compared in cattle experiencing either a short (SPE) or long (LPE) proestrus (Figure 2; Bridges et al., 2010). Ovulatory follicle size was similar and the magnitude of the GnRH-induced LH surge did not differ between treatments. There tended to be lower concentrations of progesterone in the SPE treatment, however, the most striking difference was the greater concentrations of estradiol in the LPE treatment during the 32 hours preceding GnRH administration (Figure 3). In another model, when proestrus was lengthened in the 5- vs. the 7-day CO-Synch + CIDR program, diameter of the ovulatory follicle did not differ but peak estradiol concentration tended to be greater with the 5-day program (Bridges et al., 2009). It has been concluded that increasing the length of proestrus escalates preovulatory concentrations of estradiol, presumably through
prolonged gonadotropic stimulation of the dominant follicle, resulting in increased timed-Al pregnancy rates in the 5-day CO-Sync + CIDR program.

Fig. 2. Animal model used in a series of experiments cited to result in treatments that varied in length of proestrus and prevovulatory estradiol concentrations. Animals were synchronized to a common day of estrus prior to ovarian follicular aspiration. Ovarian follicular aspiration resulted in emergence of a new wave of follicles either 1 or 2 days after aspiration. In the Long Proestrus treatment, PGF$_{20}$ (PGF) was administered approximately 4 days after aspiration and GnRH given 2.5 days later. In the Short Proestrus treatment, PGF was given 5 days after emergence and GnRH administered 1.5 days later.

*Increased gonadotropic stimulus in estradiol-based synchronization programs*

Scientists that work with estradiol-based programs (Figure 1) in *Bos indicus* influenced cattle have investigated alternative approaches to enhance gonadotropic stimulation of the dominant follicle during synchronization. The capacity of progesterone to regulate frequency of LH pulses occurs along a continuum of concentrations. If concentrations of progesterone are reduced but not to basal levels, a measurable increase in LH pulse frequency that is capable of stimulating growth of the dominant follicle occurs (Kinder et al., 1996). Therefore, it is conceivable that if progesterone concentrations are less during the period of progestin treatment during estrous synchronization, maturity of the dominant follicle at initiation of proestrus would be advanced and thereby result in increased estradiol concentrations and fertility.

One approach is to induce luteal regression before the end of progestin treatment using PGF$_{20}$ in order to remove contributions to peripheral progesterone concentrations from the corpus luteum, leaving the exogenous treatment as the only progestin source. When PGF$_{20}$ was given on the fourth day of an 8-day estradiol based program, pregnancy rate to embryo transfer was enhanced (Moreno et al., 2002). In cyclic cows that received PGF$_{20}$ on either day 7 or day 9
Fig. 3. Preovulatory plasma estradiol concentrations of beef cows that were induced to ovulate a similar sized follicle following either a long (2.2 days; LPE; ▲) or short (1.2 days; SPE; ■) proestrus. Asterisks indicate times that differed (P < 0.05) between treatments.

of a 9-day estradiol-based synchronization program (Peres et al., 2009), treatment with PGF$_{2\alpha}$ on day 7 reduced progesterone concentrations on day 9 (just before progestin withdrawal) and increased diameter of the ovulatory follicle and timed-Al pregnancy rate. Alternatively, a device that releases less progesterone, such as a CIDR that has been used previously to synchronize estrus (Martinez et al., 2003), could be inserted to lessen progesterone concentrations during treatment. Diameter of the ovulatory follicle and timed Al pregnancy rate were greater in Nelore heifers that received a twice-used CIDR compared to a new CIDR (Dias et al., 2009). While neither of these approaches technically increases the length of proestrus, the reduction in progesterone concentrations before the onset of proestrus increased follicular diameter, suggesting that the endpoint of increasing maturity of the dominant follicle and timed Al pregnancy rate may have been achieved.

Rather than using the approaches designed to increase endogenous LH listed above, administration of exogenous gonadotropins has also been used enhance follicular maturity. Treatment of beef cows with eCG on the fifth day of an 8-day estradiol based program (Tribulo et al., 2002) increased pregnancy rate of embryo transfer recipients. Likewise, treatment with exogenous eCG at the start of proestrus increased diameter of the ovulatory follicle and timed-Al pregnancy rate in estradiol-based programs (Baruselli et al., 2004; Peres et al., 2009; Dias et al., 2009). With this approach the gonadotropic stimulus is provided exogenously, but a similar net outcome of enhanced follicular growth and fertility is achieved.

Impact of length of proestrus on embryonic mortality

The animal model described in Figure 2 has been used to investigate the impact of length of proestrus and associated changes in preovulatory estradiol concentrations on uterine function and embryo development in cattle (Bridges et al., 2005; 2006a; 2006b; 2010). Due to the impact on estradiol concentrations, hereafter LoE refers to the SPE treatment and HiE refers the LPE treatment. In the first experiment gene expression for oxytocin receptor and cyclooxygenase-2 in the uterus on day 5 of the estrous cycle were greater in LoE than HiE treatment (Bridges et al., 2005) indicating that treatments influence expression of genes known to regulate luteal function. The objective of the second experiment was to determine the effect of decreased preovulatory estradiol concentrations
on embryonic development to day 6 of pregnancy (Bridges et al., 2006a). Embryos were collected on day 6 of gestation from cows that were AI following the HiE and LoE treatments. Fertilization rate, embryo grade, total cells per embryo, and accessory sperm cell number were similar between treatments. Although the number of recovered embryos was somewhat limiting, findings indicated that catastrophic defects in early embryo development were not associated with the LoE treatment. In a third experiment, aspects of conceptus and uterine function were compared on day 15.5 of gestation between cows that received the HiE and LoE treatments and were implanted with embryos from untreated cattle on d 15.5 of gestation (Bridges et al., 2006b, Bridges and Day, unpublished). Content of IFN-γ in the uterine lumen and uterine concentrations of mRNA for ISG-15 and MX-1 from pregnant animals did not differ between treatments. Nuclear progesterone receptors, predominantly localized in the deep glandular epithelium were more abundant (P < 0.05; Figure 4) in the HiE than LoE treatment. Results of this experiment suggest that up to day 15.5 of pregnancy, decreased preovulatory estradiol concentrations do not impair conceptus development or its ability to induce mRNA for interferon stimulated genes in the uterus. However, alterations in uterine progesterone receptors existed at this stage of pregnancy. Previous research (Table 1) has demonstrated a substantial negative impact of shortened proestrus on pregnancy rate by day 30 after ovulation (either with AI or embryo transfer) and in Experiment 2 (Mussard et al., unpublished) return to estrus following AI for non-pregnant cows with a shortened proestrus was approximately 25 days, suggesting that embryos survived through maternal recognition of pregnancy in some females but were lost shortly thereafter. It is therefore proposed that pregnancy loss in animals that have suboptimal preovulatory estradiol concentrations occurs after maternal recognition of pregnancy during the pre-implantation period but before day 30 of gestation. The hypothesis of ongoing experiments is that decreased preovulatory estradiol concentrations lead to inappropriate gene expression and protein synthesis by the uterus, which results in the inability of the uterus to sustain the conceptus through the pre-implantation period.

![Fig. 4. Nuclear progesterone receptor in the deep uterine glandular epithelium in cattle on day 15.5 of the estrous cycle in pregnant (P) and nonpregnant (NP) heifers that were induced to ovulate a similar sized follicle following either a long proestrus (2.5 days) that increased preovulatory estradiol concentrations (HiE) or a shortened proestrus (1.5 days) that resulted in reduced preovulatory concentrations (LoE). Representative sections from females in each treatment are shown above the respective treatment. Staining intensity differed between treatment, P < 0.05.](image-url)
Significant discoveries in the past two decades in reproductive biology regarding the manner in which ovarian follicles grow, the impact of exogenous intervention on this growth and description of methods to coordinate this process have led to development of estrous synchronization programs that consistently result in timed-Al pregnancy rates of 50% or greater in beef cattle. Recent advancements have pushed this proportion to almost 70% with some programs and emphasize that adequate gonadotropic stimulation of follicular function before ovulation is requisite to optimize fertility. The mechanisms responsible for variation in fertility related to follicular maturity may include influences on oocyte quality, embryonic growth, oviductal function, luteal function or uterine environment, and a combination of some or all of these critical aspects of pregnancy establishment is likely. The findings suggest that the elevated estradiol concentrations that result from increased stimulation of the dominant follicle may be responsible for increased fertility through actions to create a uterine environment capable of sustaining the embryo through to attachment.

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