Roles of luteal and allantoic function in late embryonic/early fetal pregnancy failures in cattle

Justin D. Rhinehart
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Roles of Luteal and Allantoic Function in Late Embryonic / Early Fetal Pregnancy Failures in Cattle

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Dissertation submitted to the
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at West Virginia University
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in
Reproductive Physiology

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ABSTRACT

Roles of Luteal and Allantoic Function in Late Embryonic / Early Fetal Pregnancy Failures in Cattle

Justin D. Rhinehart

Pregnancy failures during placentation in lactating dairy cows have been associated with low concentrations of peripheral serum progesterone (P₄) and poor development of the chorioallantoic membranes. Experiment I was done to determine if pregnant cows with high or low concentrations of serum P₄ differed in luteal production of P₄ and/or clearance of injected P₄. Luteal tissue was removed from pregnant cows with ≥ 4.0 ng/mL (High) or ≤ 2.5 ng/mL (Low) serum P₄ during d 28 to 34 post-insemination. Luteal tissue was assayed for P₄ by radioimmunoassay and for expression of mRNA for preproendothelins 1 and 3, endothelin converting enzyme, endothelin receptors A and B, cyclooxygenase-2, aldoketoreductase 1B5, 15-hydroxyprostaglandin dehydrogenase, and prostaglandin E synthase by real-time RT-PCR. Dispersed luteal cells were incubated for 2 h with bovine luteinizing hormone (bLH) or arachidonic acid (AA), increasing (10⁻¹⁰ to 10⁻⁷ M) concentrations of endothelin-1 (ET-1), and combinations of ET-1 and bLH or AA. Neither luteal content of P₄ (mean 106 ± 12 μg) nor mRNAs for the endothelin or prostaglandin systems differed with serum P₄ at lutectomy. Both basal and LH-stimulated secretion of P₄ from dispersed luteal cells were inhibited (P < 0.05) by ET-1 in a dose dependent manner. Inhibition by ET-1 was greater (P < 0.05) for luteal cells from Low vs. High cows when incubated with AA. Both basal and AA-stimulated secretion of PGF₂α by luteal cells were increased by ET-1 in a dose-dependent manner. Basal and AA-stimulated 6-keto-PGF₁α were decreased by ET-1 in a dose-dependent manner. To evaluate clearance of P₄, cows were injected s.c. with 150 mg P₄ every 12 h beginning at lutectomy. In jugular blood collected every 4 h until h 48, serum P₄ was maintained at lower (P < 0.05) concentrations for Low vs. High cows. Area under the curve was less (P < 0.05) for Low (49.6 ± 6.2) vs. High (83.6 ± 12.5) cows. On the basis of these data, differences in clearance were more important than differences in luteal production to determine peripheral concentrations of P₄. However, incubation with ET-1 and AA caused a greater reduction of P₄ production for luteal cells from cows with Low than High serum P₄. Thus, CL of Low cows might be more sensitive to luteolytic influences than CL of High cows. Reduced clearance of P₄ might decrease late embryonic or early fetal mortality as efficiently as supplementation with exogenous progestogens. The objective of experiment II was to characterize the timing of detection of the allantois and diameter of the allantoic lumen, in dairy cows and heifers with embryos developed entirely in vivo, to determine whether these variables might be used to investigate late embryonic / early fetal loss. Reproductive tracts of dairy heifers (n = 33) and lactating dairy cows (n = 30) were examined daily, via transrectal ultrasonography (Aloka 900), beginning on d 21 post-insemination (PI). Variables included: first day of detection of the allantois, diameter of the allantoic lumen, length of the embryo at first detection of the allantois, and pregnancy retention on d 60 PI. Range and mean of first day of detection of the allantois (d 21 to 26, mean = 23 ± 0.18 PI) agreed with previous reports. The allantois was detected earlier (P < 0.05) in heifers (22.4 ± 0.2 d) than in cows (23.6 ± 0.2 d).
Diameter of the allantoic lumen at first detection (4.6 ± 0.3 mm) did not differ with day of detection and was not affected by age of dam. In contrast, length of embryo (range 2.6 to 7.9 mm) varied with day of first detection of the allantois (P < 0.05), but not with age of dam. Pregnancy failure by day 60 PI did not differ with age of the dam. Differences in failure for pregnancies in which the allantois was detected earlier (≤ d 23 PI) or later (≥ d 24 PI) approached significance (P = 0.08) with those detected earlier failing more frequently than those detected later. Differences in day of first ultrasonographic detection of placental membranes, between heifers and cows, might reflect differing developmental rates and, based on present data, asynchrony in development of the embryo and the placental membranes might occur in some animals. Therefore, day of detection of the allantois might be useful for predicting survival or loss of pregnancy during placentation.
# TABLE OF CONTENTS

ABSTRACT........................................................................................................................................ ii

TABLE OF CONTENTS...................................................................................................................... iv

LIST OF TABLES ........................................................................................................................... vi

LIST OF FIGURES......................................................................................................................... vii

ACKNOWLEDGEMENTS ................................................................................................................ viii

REVIEW OF LITERATURE .............................................................................................................. 1

Introduction ...................................................................................................................................... 1

Economic Impact of Pregnancy Loss .............................................................................................. 1

Fertilization ...................................................................................................................................... 4

Pregnancy Loss ................................................................................................................................ 5

   Early Embryonic Loss ................................................................................................................ 5
   Late Embryonic / Early Fetal Loss ............................................................................................ 7
   Placental Development .............................................................................................................. 8

Frequency of Late Embryonic / Early Fetal Loss ....................................................................... 13

Effects of Low Progesterone on Pregnancy Loss ........................................................................ 14

Pregnancy Loss Associated with Assisted Reproductive Technology ............................................ 16

   Allantoic Development for IVF and SCNT embryos ............................................................... 18

   Ultrasonographic Evaluation of Embryonic Development .................................................... 19

STATEMENT OF THE PROBLEM ................................................................................................... 22

LUTEAL FUNCTION AT DAY 30 OF PREGNANCY IN RELATION TO SERUM PROGESTERONE IN DAIRY COWS AT RISK FOR LATE EMBRYONIC OR EARLY FETAL MORTALITY ........................................................................................................... 24

Abstract .......................................................................................................................................... 24

Introduction ...................................................................................................................................... 25

Materials and Methods .................................................................................................................. 29

Results ............................................................................................................................................. 33

   Luteal Progesterone and Gene Expression .............................................................................. 33
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Luteal Cell Incubation</td>
<td>34</td>
</tr>
<tr>
<td>Serum Progesterone Post-Lutectomy</td>
<td>35</td>
</tr>
<tr>
<td>Discussion</td>
<td>35</td>
</tr>
<tr>
<td>ASSOCIATION OF TIMING OF CHORIOALLANTOIC MEMBRANE DEVELOPMENT WITH AGE IN DAIRY CATTLE</td>
<td>49</td>
</tr>
<tr>
<td>Abstract</td>
<td>49</td>
</tr>
<tr>
<td>Introduction</td>
<td>50</td>
</tr>
<tr>
<td>Materials and Methods</td>
<td>51</td>
</tr>
<tr>
<td>Results</td>
<td>52</td>
</tr>
<tr>
<td>Discussion</td>
<td>53</td>
</tr>
<tr>
<td>LITERATURE CITED</td>
<td>60</td>
</tr>
</tbody>
</table>
LIST OF TABLES
Table 1.1 Primer sequences and PCR conditions ............................................................. 41
Table 2.1 Allantois Results Relative to Age of Dam........................................................ 58
# LIST OF FIGURES

<table>
<thead>
<tr>
<th>Figure</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Figure 1.1</td>
<td>Luteal Progesterone</td>
<td>42</td>
</tr>
<tr>
<td>Figure 1.2</td>
<td>Endothelin Gene Expression</td>
<td>43</td>
</tr>
<tr>
<td>Figure 1.3</td>
<td>Prostaglandin Gene Expression</td>
<td>44</td>
</tr>
<tr>
<td>Figure 1.4</td>
<td>Post Luteectomy Serum Progesterone</td>
<td>45</td>
</tr>
<tr>
<td>Figure 1.5</td>
<td>Luteal Cell Progesterone Production</td>
<td>46</td>
</tr>
<tr>
<td>Figure 1.6</td>
<td>Luteal Cell PGF₂α Production</td>
<td>47</td>
</tr>
<tr>
<td>Figure 1.7</td>
<td>Luteal Cell 6-keto-PGF₃α Production</td>
<td>48</td>
</tr>
<tr>
<td>Figure 2.1</td>
<td>Representative Ultrasonographic Image</td>
<td>59</td>
</tr>
</tbody>
</table>
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REVIEW OF LITERATURE

Introduction

Failure of lactating dairy cows to establish and maintain pregnancy is the most significant impediment to production efficiency in the United States dairy industry. Over the past four decades, conception rates for dairy cows that are artificially inseminated (AI) have declined from 65% to as little as 35% (Santos et al., 2004). Several factors have been investigated as possible causes for the decline in reproductive efficiency of lactating dairy cows. These include the effects of semen quality (Dejarnette, 2005), selection pressure for increased production (Ferguson, 2005) and changes in nutritional management (Lucy, 2003). The magnitude of the impact of poor reproductive efficiency on the dairy industry can be illustrated by review of the economic consequences.

Economic Impact of Pregnancy Loss

Reproductive performance is one of the most important factors regulating profitability in U.S. dairy herds. The significance of poor reproductive efficiency is broad in scope; reproduction impacts many aspects of overall herd management. One of the most apparent influences of deficient fertility is increased calving intervals. Longer calving intervals are less profitable because of the associated decline in milk yield with duration of lactation. This becomes even more important when considering the timing of pregnancy loss. Early embryonic death extends the calving interval by one estrous cycle length if the return to estrus is detected. Late embryonic or early fetal loss increases the calving interval by the time required for resumption of cyclicity in addition to the number
of days at which loss occurs (usually 28 to 60 days). Long calving intervals increase days in milk per lactation but reduce days in early lactation, the period of highest production per day, over a cow’s productive life. Overall, extended calving intervals decrease the cow’s lifetime milk production.

To estimate the economic benefits of improved reproductive performance through increasing pregnancy rates, De Vries and coworkers (2005) developed “The Florida Dairy Computer Program.” This advanced economic modeling program was used to estimate the economic benefits of changed management practices. The model included variables such as lactation curves, body weights, efficiency of estrous detection, conception rates, risk of voluntary culling, labor, and feed intake. Costs considered included the prices of milk, feed, labor, veterinary services, semen, and estrous synchronization drugs. By using this comprehensive model, the authors estimated that the cost per cow per extra day open (beyond 80 days postpartum) varied from $0.81 to $13.33 and increased as days open increased (De Vries et al., 2005).

Other estimates support the concept that increased cost per day open is a compounding factor as calving intervals increase (Plaizier et al., 1997). These authors used a computer simulation program to estimate the impact of poor reproductive efficiency on profit for Canadian dairy herds. By their estimation, each day increase in adjusted calving interval reduced net revenue by a mean of $4.70 (Canadian) per cow. Additionally, adjusted calving interval was more accurate in predicting economic impact because it included the cost due to reproductive culling rather than only the impact of calving interval on milk production. French and Nebel (2003), using a spreadsheet model to predict the economic impact of extended calving interval, estimated that each extra day
open at 100 days in milk cost an additional $0.50 and at 175 days cost an additional $4.52 per day per cow.

The average value of a new pregnancy for an individual cow has been estimated by several authors. “The value of a pregnancy for an individual cow can be defined as the difference in discounted future cash flows when she is pregnant compared with when she is not pregnant” (De Vries, 2006). For timed insemination protocols, the estimated value of a new pregnancy was between $253 and $274 (Stevenson, 2001). Similarly, using a modified version of the previously described “Florida Dairy Computer Program,” the average value of a new pregnancy for cows that differed in lactation number, stage of lactation and milk yield was $278 (De Vries, 2006). The values reported by De Vries (2006) increased with days-in-milk early in lactation, but decreased with days-in-milk late in lactation.

Alternatively, the cost of “mid-term” abortion (days 42 to 260 days gestation) has been estimated to range from $600 to $1,200 (Piedrahita et al., 2002). The cost for mid-term abortion in Canadian production systems, including reproductive loss and reduced milk yield, was estimated to be $1,286 (US$, Weersink et al., 2002). Although these reports were intended to illustrate the economic impact of pregnancy loss resulting from infectious diseases, the average cost of pregnancy loss at the transition to fetal development is similar for spontaneous pregnancy loss. These calculations illustrate the costs associated with reproductive failure after recognition of pregnancy.

Factors that interfere with the timely establishment of pregnancy and subsequent calving in lactating dairy cows reduce economic gain. Clearly, the amount of revenue potential lost by pregnancy wastage in dairy herds justifies research with the objective to
identify its sources. The two points at which these factors might change the underlying biology of reproductive efficiency are: 1) fertilization and 2) pregnancy maintenance.

**Fertilization**

Reports of fertilization rates have varied but, on average, were similar for lactating and non-lactating dairy cows (76.2% and 78.1% respectively; reviewed by Santos et al., 2004). However, the wide range of successful fertilization in that review (55.3 to 98.0%) indicates that environmental impacts and management strategies likely affect the ability of the male and female gametes to establish a two-cell embryo. Fertilization rates tend to be less variable and more successful for nonlactating beef cows and heifers. Sreenan and Diskin (1983) reported a 90% fertilization rate for beef heifers following a single AI, while Maurer and Chenault (1983) reported 100% fertilization rate for parous beef cows pen-mated at the onset of estrus. However, fertilization rates for a series of studies of early postpartum lactating beef cows averaged 75%, with a range from 61 to 100% (Breuel et al., 1993). These findings are similar to those in lactating dairy cows and indicate an impact of lactation on fertilization success.

In each of the reports cited above, fertilization rate was considerably greater than calving rate. For the dairy herds reviewed by Santos et al. (2004), calving rates to first service were generalized to be 28%. For beef heifers and cows, calving rates to single inseminations were reported to be only 53% (Austin et al., 1999). Regardless of relative fertilization rate and breed type, the disparity between fertilization rate and number of calves born to a single insemination indicates an extremely high occurrence of pregnancy wastage from the time a zygote is created to parturition.
Pregnancy Loss

Pregnancy loss can be defined broadly as death of the conceptus at any point from syngamy to parturition. However, as investigation of this phenomenon increased, terminology has been applied that delineates developmental periods during pregnancy. According to the Committee on Bovine Reproductive Nomenclature (1972), embryonic mortality is considered to be “the death or loss of conceptus during the embryonic period”. This same committee recommended that the definition for the embryonic period set forth by Dennis (1969) be accepted: “the period from conception to the end of the stage of differentiation”.

The embryonic period has been divided further into more concise periods of development. However, unlike much of the nomenclature used to describe periods of gestation, these terms have not yet been recognized formally but have advanced into common use in the literature. Early embryonic development refers to the period from fertilization to approximately day 20 of pregnancy. Late embryonic development indicates the period from days 20 to 45 post-fertilization. Fetal development is specified as the period of development from day 45 to term. More specifically, early fetal development refers to the period of gestation from completion of differentiation to day 60 post-fertilization (Committee on Bovine Reproductive Nomenclature, 1972).

Early Embryonic Loss

As reviewed by Aylon (1978), much of the early data collected for investigation of embryonic loss relied on determination of whether the post-insemination estrous cycle was longer than the previous or longer than would be considered average (17 – 25 days;
Erb and Holtz, 1958). More precise determination of actual embryonic loss, independent of fertilization failure, came about with the implementation of timed slaughter of inseminated cows. Tanabe and Casida (1949) slaughtered repeat-breeding (anatomically normal with a minimum of four infertile services) cows at 3 and 34 days post-AI and found that 66.1% presented a fertilized ovum at day 3 while only 23.1% had a normal embryo at day 34. Normal cows exhibited less fertilization failure and less embryonic loss at day 35 (16 and 14.5%, respectively; Ayalon, 1978).

Inskeep and Dailey (2005) reviewed the distribution of pregnancy failure from fertilization through transition to the fetal period. These authors concluded that the majority (57%) of failed pregnancies occurred during early embryonic development, within one estrous cycle after insemination. In a review of the literature compiled upon request by the Commission of the European Communities, Sreenan and Diskin (1986) suggested that, for normal cows, “while embryo losses occur gradually from fertilization onwards, the greatest increment of losses would seem to occur between about days 15 and 18”. They reached this conclusion by summarizing published fertilization rates for cows and heifers as “the number of normal cleaved ova or embryos as a proportion of all ova (fertilized or unfertilized) recovered” up to and including day 8 post-insemination. The fertilization rates for heifers and cows to natural mating or AI were 88% and 90%, respectively (Sreenan and Diskin, 1986). They then compiled data from several other authors who employed oocyte and embryo recovery at various times after insemination. When considered together, these reports supported the authors’ conclusion that most of the embryonic loss occurred between days 15 and 18 (Sreenan and Diskin, 1986).
**Late Embryonic / Early Fetal Loss**

The period of development between approximately days 25 and 60 post-insemination is often referred to as late embryonic / early fetal development (Inskeep, 2004). Pregnancy loss during this developmental period constitutes a lower percentage of the total than earlier loss. However, the economic impact is significant because it results in an extended inter-estrous interval and more days in milk before a successful pregnancy is established (reviewed under heading “Economic Impact of Pregnancy Loss”). The frequency of late embryonic / early fetal pregnancy failure is easier to establish than early embryonic loss because the pregnancy can be observed by transrectal ultrasonography on gestation day 25 or detected by manual palpation on day 35.

During the late embryonic / early fetal period of development, one of the most critical processes is placentation. Formation of an efficient system of nutrient and waste transfer between dam and fetus is required to take over support of fetal growth as the demand for nutrition becomes more than can be supplied by histotrophic maintenance from the uterus (Roberts and Bazer, 1988). Failure to form a proper and efficient placental transfer has been suggested as the factor leading to a majority of late embryonic loss that occurs between days 30 and 45 of gestation (Dailey et al., 2002). This concept was supported by Starbuck and coworkers (2004) who reported that most pregnancy loss after day 30 in dairy cows and heifers had occurred before day 42 post mating. To understand pregnancy wastage during placentation, placental development should be considered.
Placental Development

The mammalian placenta is composed of three membranes: the chorion, allantois, and amnion. Each membrane has differing degrees of functional importance depending on species. For mammals, the chorion directly apposes the uterine endometrium and is responsible for generating the areas of nutrient and waste transfer between fetal and maternal circulation. The allantois produces the vasculature and hematopoetic potential that will create blood flow for placental function. The amnion develops primarily as a fluid-filled protective barrier against environmental insult to the fetus.

One of the first major steps in placental development is formation of the endoderm, mesoderm and ectoderm. This process, referred to as gastrulation, transforms the embryo into three primary germ layers that will give rise to all tissues and organs that comprise the mature mammal (Kaufman, 1995). Gastrulation begins with formation of the primitive streak, a group of cells that arise from the epiblast layer. The cells of the primitive streak then begin to migrate into the space between the epiblast and hypoblast. Cells that will become endoderm move into the hypoblast and displace it into the extraembryonic space. Cells that will become mesoderm populate the area between the epiblast and hypoblast. The ectoderm is composed of the remaining cells of the epiblast. The embryo becomes elongated as these cell layers begin to migrate and differentiate. Contact between the germ layers is responsible for stimulating organogenesis, the next major progression in embryogenesis (Kaufman, 1995). Relatively small changes in the local environment or disregulation of genetic control for this process can impact development of the embryo and its extraembryonic membranes.
**Chorion:** Two layers of differing cellular origin comprise the chorion: the trophoblast and mesoderm. The trophoblast is the outer layer of the chorion that contacts uterine endometrium, and the inner mesodermal layer contacts the amnion. The trophoblast includes two distinct cellular types; an internal layer of cuboidal cells called cytotrophoblast and an external layer of cells called syncytiotrophoblast. By definition (syncytium), these cells become multinucleated by either karyokinesis without cell division or by the fusion of several mononucleated cells. For mammalian embryos that implant into the endometrium, this layer is highly invasive near the inner cell mass. For the horse and human being, the syncytiotrophoblast produces placental gonadotropins (equine chorionic gonadotropin, eCG and human chorionic gonadotropin, hCG; respectively) as luteotropins. Binucleate giant cells, also of chorionic origin, are unique to the ruminant and produce pregnancy specific protein B. Cells originating from the cytotrophoblast are responsible for placental steroid production.

Classification of placental relationship with the uterus is based on the number of tissue layers that separate the fetal and maternal blood. The epitheliochorial placenta presents the highest degree of maternal and fetal separation. In this situation, both the endometrial epithelium and the epithelium of the chorionic villi remain intact. This type of placentation is found in ruminants as well as in the mare and sow. Lack of intimate apposition of blood flow is offset by a large placental surface area in the sow and mare and by increased efficiency of transfer in ruminants. The endotheliochorial placenta, as for dogs and cats, lacks endometrial epithelium such that maternal capillaries are directly exposed to chorionic epithelium. Direct exposure of chorionic epithelium to pools of maternal blood is referred to as a hemochorial placenta (primates). Finally,
hemoendothelial placenta (rat, rabbit and guinea pig) is the most intimate type as the placental vasculature is immersed in pools of the maternal blood.

**Allantois:** In Knobil and Neill’s *Encyclopedia of Reproduction*, the allantois is defined as “a membrane that develops as an evagination of the hindgut to form a sac filled with water, urine, and nutrients” (Bazer, 1998). It begins as a diverticulum of endoderm covered by mesoderm. For cattle, the allantois begins to develop on approximately day 20 post-fertilization (Flechon and Renard, 1978; Greenstein et al., 1958). It continues to grow and fill with allantoic fluid until it occupies the extra-embryonic space and contacts the chorion.

Greenstein and coworkers (1958) used serial slaughter of heifers and cows to characterize gross anatomical and histological features of the bovine preattachment placenta from days 16 to 33 of gestation. They stated that, on days 21 and 22, “the allantoic diverticuli have become a constant feature of embryos at this age and the enlarging distal portions approach a centimeter in length.” During days 23 to 25, the allantois continues to expand rapidly, is becoming highly vascularized and ranges from 0.8 to 5.5 cm in length. On days 26 and 27, the allantoic vesicle ranges in length from 5.5 cm to 11.0 cm. During days 28 to 30 of gestation, the allantois has begun to fuse with the “overlying trophoderm” and contains an extensive network of vessels filled with nucleated fetal red blood cells (Greenstein et al., 1958).

The ultrastructural characterization of post-hatching development for the bovine embryo was reported recently (Maddox-Hyttel et al., 2003). For seven, day-21 embryos, gestated together as a result of superovulation and breeding on day 0, the length of the embryo ranged from 3.14 to 5.79 mm while the allantois ranged in width from 1.15 to
4.41 mm. The authors noted “a clear relationship between the embryo length and width of the allantois”, but provided no statistical analysis to support the comment. The allantois had not yet developed for two embryos that were 3.14 and 3.77 mm long (Maddox-Hyttel et al., 2003).

Recently several genes have been identified as being expressed differentially during allantoic emergence in bovine and ovine conceptuses (Ledgard et al., 2006). These authors used suppressive subtractive hybridization to compare mRNA expression in preimplantation conceptuses prior to emergence of the allantois and in allantoic samples after emergence. Nine genes were identified as differentially expressed. All were associated with extracellular matrix and vasculogenesis. Similarly, Ushizawa et al. (2004) used cDNA microarray analysis to determine gene expression profiles in the bovine embryo during the preattachment period. They found differential expression of more than 1,000 genes on days 7, 14, 21 and 28. Comparison of extra-embryonic membranes and the embryo proper on day 28 revealed 119 down-regulated genes and 74 up-regulated genes in the membranes. The up-regulated genes included placental lactogen, prolactin-related proteins, pregnancy associated glycoproteins, and major histocompatibility complex molecules.

A great deal of focus has been placed on development of the murine allantois (Downs et al., 2004). Results from several of these experiments indicated that development of the allantois requires both intrinsic mechanisms and communication with the primitive streak. Recall that the primitive streak is a thickening at the midline of the epiblast from which the endoderm and mesoderm differentiate and move to appropriate sites in the conceptus (Poelmann, 1981). The posterior primitive streak is the site of
origin of the allantoic bud. Extraembryonic mesoderm accumulates between the amnion and yolk sac and enlarges into the exocoelomic cavity by proliferation, continuous deposition of mesoderm from the streak, and distal cavitation (Ellington, 1987).

Vascularization of the murine allantois arises from an outer layer of mesothelium and an inner layer of mesoderm (Downs, 1998). Formation of endothelium occurs de novo in the allantois. Endothelial tubules begin to form at the distal region, proceed proximally to the base of the allantois, and join with the yolk sac to create a continuous vasculature throughout the conceptus (Downs, 1998). Eventually, the plexus of vasculature in the allantois is transformed into the umbilical vein and artery (Kaufman, 1995).

Unlike the yolk sac, in which development of blood islands depends on contact with endoderm (Belaoussoff et al., 1998), the allantoic mesoderm grows into the exocoelomic cavity as an independent structure that does not contact other tissue until after the allantoic endothelium and mesothelium have developed (Downs, 1998). Therefore, it would appear that development of the allantois is guided either by intrinsic factors or by the primitive streak. When detached portions of allantoic membrane were inserted into the exocoelomic cavity of synchronous hosts, they attached to the host chorion at the appropriate time (Downs and Gardner, 1995). Furthermore, when murine allantoises were cultured apart from the conceptus, they developed normal vascularization (Downs and Harmann, 1997). Both of these observations support the concept that allantoic development is directed by intrinsic factors.

Differentiation of the allantois into endothelial and chorio-adhesive cells begins in the region furthest from the primitive streak (Downs and Harmann, 1997). Downs and
coworkers (2004) hypothesized that the primitive streak controls cellular differentiation because the expression of vascular cell adhesion molecule (VCAM1; an adhesion molecule specific to chorio-adhesive cells) was limited to the distal regions of the allantois (Gurtner et al., 1995). When mesoderm from the primitive streak (precursor of allantoic tissue) was transplanted into the distal region of recipients, it exhibited limited development (Dunwoodie and Beddington, 2002). Similarly, when newly budded mesoderm of the primitive streak was transplanted to distal regions of hosts, it contributed only to blood vessels and not to somatic mesoderm (Downs and Harmann, 1997). Taken together, the results of these studies indicated that allantoic endothelial cells are specified within the primitive streak.

From the results discussed in the two previous paragraphs, it appears that differentiation of cell types found in the murine allantois is determined in the primitive streak before it begins to bud from the embryo proper. After differentiation, development is guided by intrinsic factors until the allantois contacts the chorion. However, it is not completely independent of the embryo because, as it continues to develop and grow into the exocoelomic cavity, the primitive streak maintains contribution of mesoderm to the membrane. The hypothesized intrinsic factors seem to be regulated by age of the cells, due to the pattern of development beginning at the portion of the allantois distal to the primitive streak.

**Frequency of Late Embryonic / Early Fetal Loss**

Several reviewers have compiled data from field trials that describe the rate of pregnancy failure from pregnancy diagnosis (gestation days 25 or 30) until the early fetal period (gestation days 45, 60, 80 and/or 100). Smith and Stevenson (1995) reported an
average “late embryonal loss” of 12.4% for cows and heifers bred after several different estrous synchronization protocols. Pregnancy survival was not altered by method of synchronization but tended to be higher in heifers than in multiparous cows (Smith and Stevenson, 1995). Vasconcelos et al. (2003) reported 10.5% pregnancy loss between days 28 and 42 of gestation for lactating dairy cows.

Milk progesterone has been used to identify luteal phases presumed to have been extended (> 24 days) by early pregnancy. Using this method, several authors have reported the frequency of pregnancy loss up to day 100 post-insemination (Kummerfeld et al., 1978; Bulman and Lamming, 1979; Grimard et al., 2006). These estimates ranged from 7% to 25.2% and agreed with the ranges reported in the reviews by Dailey et al. (2002) and Inskeep (2004). Humbolt (2001) reported that late embryonic mortality ranged between 8 and 17% with a mean of 14%. More recently, Grimard et al. (2006) reported that season and body condition score (BCS) affected pregnancy loss between pregnancy days 21 and 100 despite not affecting conception rates.

**Effects of Low Progesterone on Pregnancy Loss**

The effect of progesterone on embryonic survival in the cow has been reviewed (Inskeep, 2004). In part, that review focused on the effect of progesterone on growth dynamics of the preovulatory follicle and how those changes affect early embryonic development. When circulating concentrations of progesterone or progestagen were low, follicles were maintained in a static growth phase while ovulation is inhibited (Imwalle et al., 2002). When follicles were forced to persist, fertility at the resulting ovulation was compromised (Stock and Fortune, 1993; Smith and Stevenson, 1995). This decrease in fertility can be traced to poor oocyte quality resulting from follicular persistence (Austin
et al., 1999; Revah and Butler, 1996). The oocyte can be fertilized, but development of the resulting embryo is interrupted (Ahmad et al., 1995).

Late embryonic / early fetal losses in dairy cows have been associated with low circulating concentrations of progesterone (Starbuck et al., 2004). In this study, the authors evaluated cows bred to either spontaneous or synchronized estrus and by AI or natural service. Pregnancy was diagnosed on days 28 to 37, and blood samples were collected for quantification of progesterone and estradiol. Pregnant cows were examined again between days 45 to 51 and between days 59 and 66. Overall pregnancy loss between days 28 to 37 and 59 to 66 was 11.4% (Starbuck et al., 2004). This result agreed with the averages reported by reviewers of late embryonic / early fetal mortality (Kummerfeld et al., 1978; Dailey et al., 2002; Inskeep and Dailey, 2005). Pregnancy loss before day 45 was greater for cows in the lowest 25th percentile of circulating progesterone concentrations (0.4 to 3.7 ng/mL; 20%) on days 28 to 37 than for cows in the middle 50% (3.9 to 5.9 ng/mL; 4%) or upper 25% (6.0 to 17.0 ng/mL; 4%; Starbuck et al., 2004).

The cause of low concentrations of progesterone in peripheral circulation is not yet certain. It could be due to low luteal progesterone production, increased progesterone metabolism or both. Sangsritavong et al. (2002) reported an acute increase in liver blood flow and metabolic clearance rate for progesterone and estradiol 17β in lactating and non-lactating dairy cows fed above maintenance. They indicated that this difference in metabolic clearance of infused steroids could be responsible for reduced fertility in high-producing dairy cows that had increased feed intake (Sangsritavong et al., 2002).
Lower circulating concentrations of progesterone and estradiol for lactating cows compared to heifers, in spite of the fact that follicular and luteal volumes were greater in cows, might indicate a difference in metabolic clearance (Sartori et al., 2004). This same group of authors compared dry and lactating dairy cows. They found that follicles and luteal structures were larger for lactating cows, but peripheral concentrations of estradiol and progesterone did not differ (Sangsritavong et al., 2002). This might indicate that the added metabolic demand of lactation leads to compensatory growth and function of steroidogenic ovarian structures.

**Pregnancy Loss Associated with Assisted Reproductive Technology**

While the timing of embryonic and fetal death has been well characterized and the frequency of pregnancy failure in cattle has been associated with several hormones, changes in the molecular biology of conceptuses that lead to late embryonic or early fetal death have not been determined. Other situations that lead to poor reproductive efficiency might prove to be useful tools for developing hypotheses to be tested in lactating dairy cows. One such situation is *in vitro* production of bovine embryos.

Calving rates from transfer of *in vitro*-produced bovine embryos to apparently normal recipient cows or heifers are poor. The degree to which reproductive success is altered depends on the method of fertilization and type of culture system used. Even though the amount of loss is substantially more, the timing of pregnancy failure for these situations often resembles the timing of both early embryonic and late embryonic / early fetal losses for AI-derived pregnancies in dairy cattle. Therefore, studying pregnancy failure from embryos produced by *in vitro* fertilization (IVF) and somatic cell nuclear
transfer (SCNT) cloning might provide new insight into pregnancy loss for AI pregnancies in dairy cows and heifers.

Embryos produced by IVF and SCNT cloning exhibit differences in morphology, gene expression and developmental ability (Crosier et al., 2001; Farin et al., 2001; Hansen and Block, 2004; Hasler, 2000) when compared to in vivo-produced embryos. Embryos produced by IVF have higher rates of embryonic mortality than those produced by AI (Hansen and Block, 2004; Hasler, 2000). Farin (2001) reported that the majority of pregnancy loss after transfer of IFV-produced embryos occurred before day 21 of gestation. In this study, 19% of the IVF embryos recovered on day 17 were degenerating compared to none of the in vivo produced and transferred embryos (Farin et al., 2001).

Survival rates to term for SCNT-cloned bovine embryos ranged from 0% to 80% (Kato et al., 1998; Piedrahita et al., 2002). Hill and coworkers (2000) reported survival rates for embryos derived in vivo or by SCNT. At day 30 of gestation, there was no difference in pregnancy rates of recipients receiving in vivo- or SCNT-produced embryos. However, the in vivo group received one embryo per recipient while the other group received two cloned embryos. By day 90 of gestation, 82% of the SCNT pregnancies were terminated while all the in vivo-derived pregnancies survived. Pregnancy loss for the SCNT group was distributed evenly over the 60-day period with 35% dead on day 40, an additional 32% dead on day 60 and 15% more dead on day 90 (Hill et al., 2000). In a similar study, pregnancy rates for recipients that received only one embryo produced by IVF or SCNT cloning were similar to each other and not different from AI pregnancies, on day 50 of gestation (De Vries et al., 2005). No pregnancy loss occurred for AI and
IVF embryos, but the SCNT embryos continued to die until only 40% of those fetuses remained on gestation day 150 (De Vries et al., 2005).

**Allantoic Development for IVF and SCNT embryos**

A majority of the embryonic mortality between days 30 and 60 of gestation for IVF- and SCNT-derived pregnancies can be attributed to improper development of the allantois (Hill et al., 2000). The allantois is the critical link for developing an efficient placenta that will be able to support fetal development beyond the transition away from histotrophic support. Using serial slaughter of recipient cows that received an IVF embryo or were artificially inseminated, it was determined that the degree of malformation of the allantois ranged from aplasia to normal growth and vascularization without haematopoiesis (Piedrahita et al., 2002).

Size of the follicle donating an oocyte for IFV affected placental development of the resulting embryo (Piedrahita et al., 2002). The length and width of the allantois on day 27 of gestation were smaller for embryos derived from small than from large follicles. However, there was no difference in length and width of the embryo proper or the degree of vascularization of the allantois in relation to follicular size. In addition to differences in growth rate of the allantois, these authors reported a difference in the ability of the allantois to fuse with the chorion. For all 17 conceptuses derived from small follicles, the allantois and chorion detached during collection. In contrast, 14 of 18 conceptuses derived from large follicles presented fully fused chorioallantoic membranes that could not be detached manually after collection (Piedrahita et al., 2002). These findings hold particular interest in relation to the reports of increased embryonic loss for
cows bred to ovulation of either immature (Perry et al., 2005) or persistent (Ahmad et al., 1995) follicles.

As reviewed here, and considered in light of the information gained from recent studies that characterize the genes controlling allantois development, new hypotheses can be tested to determine particular alterations caused by IVF and SCNT cloning. Moreover, when the genetic aberrations that lead to poor allantoic development are elucidated, their patterns of expression can be compared in AI pregnancies that are destined to succeed or fail during placentation. However, at this time, methods to determine the pattern of allantoic development without termination of the pregnancy have not been reported.

**Ultrasonographic Evaluation of Embryonic Development**

Transrectal ultrasonography holds promise as a non-invasive tool to observe extraembryonic membrane development. Use of ultrasonography to observe reproductive anatomy in cattle has allowed investigators to characterize changes in the gross anatomical features of the ovary, uterus and conceptus. More recently, Doppler ultrasonography has been used to measure changes in blood flow to and from tissues. Texture analysis or quantification of pixel intensity can now be used to make inferences about tissue ultrastructure.

Researchers at the University of Wisconsin were among the first to monitor embryonic and fetal development in the cow (Pierson and Ginther, 1984; Curran et al., 1986a, 1986b; Kastelic et al., 1989, 1991). Average first detection of the embryonic vesicle in heifers occurred on day 19 post-insemination when a 5 MHz transducer was used (Curran et al., 1986a). Using a transducer with greater resolution (7.5 MHz) to image the reproductive tract of cows, the embryonic vesicle was reported to have been
observed as early as day 9 post-insemination (Boyd et al., 1988) but those results have not been repeated.

Average first day of detection of the embryo proper, as a small echoic spot in a bulged area of the vesicle, was day 20 for Curran and colleagues (1986a). Boyd et al. (1988) reported to have found the embryo proper on day 13. On average, the heartbeat was detected on days 21 (Boyd et al., 1988) or 22 (Curran et al., 1986b). Curran and others (1986a) reported average heart rate of 188 beats-per-minute on the first day of detection. In a more recent study, the average first days of detection of the embryo and embryonic heartbeat were 24 and 28, respectively (Kolour et al., 2005). These authors used an 8 MHz probe. It is interesting to note the variability for day of first detection of the vesicle and embryo proper. However, the first day of detection for structures developing later in gestation becomes more consistent.

Each of the authors cited above reported the first day of detection of the allantois. On average, the allantois was detected first by Curran and colleagues (1986b) on day 23 post-insemination. Similarly, Boyd and others (1988) reported first observing the allantois on day 23. Though reporting a similar range, parity of the dam affected the timing of detection of the allantois by the Iranian group (Kolour et al., 2005). They first observed the allantois on days 22, 23 and 26 in heifers, primiparous and multiparous cows, respectively. The day of detection of the allantois was similar for heifers and primiparous cows and each was statistically different than multiparous cows. In their discussion, Kolour et al. (2005) suggested that the relatively large size of the uterine horns affected their ability to detect structures earlier in multiparous cows. However, the possibility that embryonic development occurs at a faster rate in heifers and younger
cows cannot be excluded. This concept would be supported by the proposal that increased circulating progesterone, as a result of lower average milk production, maintains more histotrophic support, similar to that demonstrated in the ewe (Satterfield et al., 2006).
STATEMENT OF THE PROBLEM

Currently, reproductive efficiency is poor for lactating dairy cows. Embryonic mortality occurs more frequently during early embryonic development (days 1 to 20 post-insemination) than late embryonic / early fetal development (days 20 to 45 post-insemination Sreenan and Diskin, 1986). However, when pregnancy failure occurs during late embryonic / early fetal development, the inter-estrous interval, calving interval and days in milk are all increased beyond the length of one estrous cycle. Increased calving interval decreases the time a cow is in peak production over her lifetime and decreases profit.

Pregnancy wastage during late embryonic / early fetal development averages 10 to 15% for dairy cattle. For contemporary production scenarios, loss of a pregnancy at this time constitutes a revenue loss of $600 (De Vries, 2006). For the US dairy herd (approximately 9 million cows; US Agriculture Census, 2002) an average of 12% pregnancy wastage at placentation would mean a revenue loss of $6.5 million. Clearly, finding management techniques or treatments that reduce the amount of late embryonic / early fetal loss would have a significant impact on the profitability of dairy production.

The main objective of these experiments was to determine causes of pregnancy loss during placentation in dairy cattle. One specific objective was to determine if decreased circulating concentration of progesterone at this time, with which pregnancy loss is known to be associated, is due to decreased luteal secretion of progesterone or to increased metabolism of progesterone. To address this objective, corpora lutea were removed from cows determined to fall in the low and high classifications of circulating progesterone as determined by the work of Starbuck et al. (2004). Luteal function was
evaluated while the fate of supplemental progesterone was monitored for cows after CL removal.

A second objective was to determine if growth characteristics of the allantois could be observed by transrectal ultrasonography and if pregnancy loss up to day 60 post-insemination was associated with these characteristics. To evaluate these questions, late embryonic development in dairy cows and heifers was monitored daily by transrectal ultrasonography. Allantoic growth traits were compared between cows and heifers and with regard to development of the embryo proper.
LUTEAL FUNCTION AT DAY 30 OF PREGNANCY IN RELATION TO SERUM PROGESTERONE IN DAIRY COWS AT RISK FOR LATE EMBRYONIC OR EARLY FETAL MORTALITY

Abstract
Pregnancy failures during placentation in lactating dairy cows have been associated with low concentrations of peripheral serum progesterone (P₄). Experiments were done to determine if pregnant cows with high or low concentrations of serum P₄ differed in luteal production of P₄ and/or clearance of injected P₄. Luteal tissue was removed from pregnant cows with ≥ 4.0 ng/mL (High) or ≤ 2.5 ng/mL (Low) serum P₄ during d 28 to 34 post-insemination. Luteal tissue was assayed for P₄ by radioimmunoassay and for expression of mRNA for preproendothelins 1 and 3, endothelin converting enzyme, endothelin receptors A and B, cyclooxygenase-2, aldoketoreductase 1B5, 15-hydroxyprostaglandin dehydrogenase, and prostaglandin E synthase by real-time RT-PCR. Dispersed luteal cells were incubated for 2 h with bovine luteinizing hormone (bLH) or arachidonic acid (AA), increasing (10⁻¹⁰ to 10⁻⁷ M) concentrations of endothelin-1 (ET-1), and combinations of ET-1 and bLH or AA. Neither luteal content of P₄ (mean 106 ± 12 μg) nor mRNAs for the endothelin or prostaglandin systems differed with serum P₄ at lutectomy. Both basal and LH-stimulated secretion of P₄ from dispersed luteal cells were inhibited (P < 0.05) by ET-1 in a dose dependent manner. Inhibition by ET-1 was greater (P < 0.05) for luteal cells from Low vs. High cows when incubated with AA. Both basal and AA-stimulated secretion of PGF₂α by luteal cell culture were increased by ET-1 in a dose-dependent manner. Basal and AA-stimulated 6-keto-PGF₁α were decreased by ET-1 in a dose-dependent manner. To evaluate clearance of P₄, cows were injected s.c. with 150 mg P₄ every 12 h beginning at lutectomy. In jugular blood
collected every 4 h until h 48, serum P₄ was maintained at lower (P < 0.05) concentrations for Low vs. High cows. Area under the curve was less (P < 0.05) for Low (49.6 ± 6.2) than for High (83.6 ± 3.3) cows. On the basis of these data, differences in clearance were more important than differences in luteal production to determine peripheral concentrations of P₄. However, culture with ET-1 and AA caused a greater reduction of P₄ production for luteal cells from cows with Low than High serum P₄. Thus, CL of Low cows might be more sensitive to luteolytic influences than CL of High cows. Reduced clearance of P₄ might decrease late embryonic or early fetal mortality as efficiently as supplementation with exogenous progestogens.

**Introduction**

Failure of lactating cows to establish and maintain pregnancy is a significant impediment to production efficiency. Over the past four decades, conception rates to a single service for artificially inseminated (AI) dairy cows have declined from around 65% to as little as 35% (Santos et al., 2004). Several factors have been investigated as possible causes for the decline, including semen quality (Dejarnette, 2005), selection pressure for increased production (Ferguson, 2005), and changes in nutritional management (Lucy, 2003).

During the late embryonic / early fetal period of development (d 30 to 60 post-insemination) formation of an efficient system for transfer of nutrients and wastes between dam and fetus is required to support fetal growth as the demand for nutrition becomes more than can be supplied by histotroph (Roberts and Bazer, 1988). Failure to form an appropriate placental transfer system has been suggested as the factor leading to a majority of the embryonic losses that occur between d 30 and 45 of gestation (Dailey et
Starbuck and coworkers (2004) reported that most pregnancy loss after d 30 in dairy cows and heifers had occurred before d 42 post mating, the time at which embryonic development can no longer be supported by uterine histotroph alone. In addition, losses were greater in association with low circulating concentrations of progesterone around d 30 of pregnancy. Low concentrations of progesterone in peripheral circulation of pregnant cows could be due to low luteal production of progesterone, increased metabolism of progesterone or both. Much of the current research supports the hypothesis that lower progesterone during the estrous cycle is due to increased metabolism (Wiltbank et al., 2006). Sangsritavong et al. (2002) reported acute increases in liver blood flow and metabolic clearance rates for progesterone and estradiol-17β in lactating and non-lactating dairy cows fed above maintenance. A series of studies using Holstein-Friesian dairy cows established that greater feed intake reduced serum concentrations of progesterone during the luteal phase or during treatment with exogenous progesterone via subcutaneous injection or a controlled internal drug releasing device (Rabiee et al., 2001ab, 2002ac).

Other studies have been focused on luteal secretion of progesterone. One area of investigation has been interactions of the prostaglandin (PG) and endothelin systems. The inhibitory effect of PGF$_2$α on luteal progesterone production *in vitro* was observed in luteal slices and large steroidogenic luteal cells only when co-cultured with endothelial cells (Girsh et al., 1995) indicating that PGF$_2$α might influence steroidogenic cells indirectly through endothelial cells. The endothelin system includes a group of at least 6 genes that encode a family of signaling peptides, receptors, and converting enzymes (Meidan and Levy, 2002). In bovine CL, synthesis of ET-1 has been demonstrated by the
presence of ET-1 protein and mRNA (Girsh et al., 1996a). Addition of ET-1 reduced in vitro production of progesterone by dispersed luteal cells from midcycle CL (Girsh et al., 1995; Hinckley and Milvae, 2001). Furthermore, in the midluteal phase of the bovine estrous cycle, PGF$_2\alpha$ stimulated expression of preproendothelin-1 (Girsh et al., 1996b). Therefore, ET-1 might mediate the luteolytic effects of PGF$_2\alpha$. Both PGF$_2\alpha$ and ET-1 appear to be required for functional luteolysis (Hayashi et al., 2003).

Choudhary et al. (2004) suggested that expression of the endothelin system is regulated by both PGF$_2\alpha$-independent and dependent pathways. They measured luteal expression of mRNA for prepro-ET-1 on d 1, 4, 10 and 17 of the bovine estrous cycle. It increased from d 1 to d 4 and 10 and was not changed by administration of PGF$_2\alpha$ (PGF$_2\alpha$-independent phase). On d 17, ET-1 mRNA expression was increased by exogenous PGF$_2\alpha$ compared to saline injection (PGF$_2\alpha$-dependent phase). In a subsequent experiment, ET-1, but not PGF$_2\alpha$, inhibited tonic secretion of progesterone from d 4 luteal cells (Choudhary et al., 2005). These findings led to the hypothesis that the ability of PGF$_2\alpha$ to decrease production of progesterone develops as its signal transduction cascade matures, at which point PGF$_2\alpha$ and ET-1 can act together to facilitate luteolysis. By testing this hypothesis, Sen and coworkers (2006) found that conventional protein kinase C (PKC) isozymes mediate inhibitory effects of ET-1 on progesterone secretion in the d 4 CL. A specific isozyme, PKC$\varepsilon$, was detected only in steroidogenic cells, increased from d 4 to 10, and likely modulates the ability of PGF$_2\alpha$ to reduce progesterone secretion (Sen et al., 2005, 2006).

The uterus traditionally has been considered the primary source of luteolytic PGF$_2\alpha$ in the cow. However, early reports demonstrated that corpora lutea of domestic
animals produced significant amounts of PGF$_2$$\alpha$ (Shemesh and Hansel, 1975; Patek and Watson, 1976). Watson and Patek (1979) reported reduced secretion of PGF$_2$$\alpha$ by luteal tissue of pregnant pigs compared to pigs that were late in the estrous cycle. Subsequently, Rexroad and Guthrie (1979) and Guthrie et al. (1979) showed that corpora lutea of ewes and pigs, respectively, secreted more PGF$_2$$\alpha$ after treatment with exogenous PGF$_2$$\alpha$ to induce luteolysis. Wade and Lewis (1996) confirmed that exogenous PGF$_2$$\alpha$ stimulated utero-ovarian secretion of PGF$_2$$\alpha$, as shown by an increase in PGF$_2$$\alpha$ in vena caval plasma in intact, but not in hysterectomized / ovariectomized ewes.

Similar studies in the cow revealed that synthesis of PGI$_2$ (measured as its metabolite 6-keto-PGF$_1$$\alpha$) and PGF$_2$$\alpha$ by luteal cells in vitro varied with stage of the estrous cycle (Milvae and Hansel, 1983). Tsai and Wiltbank (1997, 1998) proposed that secretion of PGF$_2$$\alpha$ from the corpus luteum, in an autocrine or paracrine manner, amplifies the endocrine luteolytic signal from the uterus. A single injection of PGF$_2$$\alpha$ upregulated mRNA encoding prostaglandin G/H synthase 2 (PGHS-2, COX-2) in mid- and late cycle ovine and bovine corpora lutea, but was not effective in animals on d 4 of the estrous cycle. Arosh et al. (2004) proposed that increased luteal prostaglandins mediate luteolysis because the enzymatic activity has shifted to favor synthesis of PGF$_2$$\alpha$ rather than PGE$_2$.

Whether low circulating concentrations of progesterone in pregnant cows are the result of decreased luteal production, increased clearance, or both, has yet to be directly and simultaneously investigated during pregnancy in the lactating dairy cow. Therefore, the first objective of this study was to compare: 1) steroidogenic function, 2) expression
of enzymes involved in synthesis and degradation of prostaglandins, and 3) expression of components of the luteal endothelin system of CL collected from cows with low or high peripheral progesterone on d 30 of pregnancy. The second objective was to compare concentrations of progesterone in peripheral blood of these cows when a standard dosage of progesterone was injected after lutectomy.

Materials and Methods
Lactating cows in the West Virginia University herd were examined by transrectal ultrasonography for the presence of a viable embryo with a heartbeat on d 28 to 34 of gestation. Days postpartum at initial blood sample averaged 102 ± 8.2. A blood sample was collected from the jugular vein, allowed to clot for 6 h, and serum was assayed immediately for progesterone so that the results were available the next morning. Only cows with concentrations of progesterone ≤ 2.5 ng/mL (Low) or ≥ 4.0 ng/mL (High) were selected for the study (n = 7 Holstein and 5 Ayrshire). These criteria were chosen based on the report by Starbuck et al. (2004) and intended to provide a comparison of cows that have an increased risk of pregnancy loss (Low) with cows that would be expected to have a greater pregnancy retention rate (High). All procedures were approved by the Animal Care and Use Committee of West Virginia University (ACUC # 03-0402).

Immediately prior to lutectomy, size of the corpus luteum (vertical by horizontal diameter) and embryonic viability were determined via transrectal ultrasonography. Corpora lutea were removed via supravaginal incision as described by Casida (1959), under epidural anesthesia with 2% lidocaine hydrochloride, the day after initial sampling for circulating progesterone concentration. Luteal tissue was placed in cold physiological saline and immediately transported to the laboratory for further processing. A portion of
the tissue was minced with a sterile scalpel in cold Medium 199 (M 199; GIBCO, Carlsbad, CA) and shipped, on ice, overnight to the University of Connecticut for luteal cell incubation. The remaining luteal tissue was cubed, snap-frozen in liquid nitrogen, and stored at -80°C.

Jugular blood samples were taken every 4 h beginning at luteectomy (h 0) and continuing through h 48. Samples were allowed to clot for 12 h, centrifuged at 1,500 x g for 30 min, and two aliquots of serum were stored at -20°C. Beginning immediately after blood sampling at h 0, progesterone (150 mg) was injected subcutaneously every 12 h until the last blood sample was taken. This dosage was used in earlier studies (Bridges et al., 2000; Starbuck et al., 2004) to successfully maintain pregnancy in cows lutectomized at various times from day 5 to 35. Milk was discarded during and after treatment with exogenous progesterone as directed by the United States Food and Drug Administration.

Real-Time RT-PCR. Expression of messenger ribonucleic acid (mRNA) for preproendothelin-1, preproendothelin-3, endothelin receptor-A, endothelin receptor-B, endothelin converting enzyme-1, cyclooxygenase-2, aldoketoreductase 1B5, prostaglandin 15-dehydrogenase, and prostaglandin E synthase in luteal tissue was measured by quantitative real-time PCR. Total RNA was isolated from frozen luteal tissue using Trizol reagent according to the manufacturer's instructions (GIBCO BRL, Gaithersburg, MD). RNA quality was determined by separation and visualization on a 1.5% agarose gel stained with ethidium bromide and quantified spectrophotometrically at 260 nm. For each sample, 2 μg of DNase-treated total RNA was reverse transcribed to first-strand cDNA using oligo (dT)18 primer and Superscript II reverse transcriptase (Invitrogen, Carlsbad, CA). Negative control reverse transcription reactions without the
enzyme were carried out to confirm that there was no contamination with genomic DNA. Real-time PCR primers for the genes listed above and the control gene glyceraldehyde-3-phosphate dehydrogenase (GAPDH) were designed based on each gene sequence using Primer3 software (Table 1.1; http://frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi). Quantitative PCR was performed in duplicate for each cDNA sample on a Bio-Rad iCycler iQ Real-Time PCR Detection System using iQTM SYBR1 Green Supermix (Bio-Rad, Hercules, CA) in 25-ml reaction volumes containing 300 nM of each primer and 2 μl diluted cDNA. Standard curves for each gene and the control were constructed using 10-fold serial dilutions of cDNA from reverse transcription of primers yielding a larger product (≈ 350 bp) than cDNA from samples (≈ 150 bp). Threshold lines were adjusted to intersect amplification lines in the linear portion of the amplification curve and cycles to threshold (Ct) were recorded. For each sample, the quantities of mRNA for the genes of interest and the control gene (GAPDH) were determined from the appropriate standard curves. The quantity of each mRNA was then divided by the quantity of mRNA for GAPDH to obtain a normalized value.

**Incubation of Luteal Cells.** Dispersed luteal cells were obtained as described by Milvae et al. (1986). For analysis of net biosynthesis of progesterone, 1 X 10⁵ cells were incubated in quadruplicate with treatments for 2 h and frozen at the end of incubation. Treatments included: control, 5 ng bovine LH (USDA bLH B-5, USDA Animal Hormone Program, Beltsville, MD), 1 μg arachidonic acid (AA; Sigma Chemical Co., St. Louis, MO), 10⁻¹⁰ to 10⁻⁷M endothelin-1 (ET-1; Sigma Chemical Co., St. Louis, MO), and combinations of each dosage of ET-1 with LH or AA. One set of quadruplicate tubes for each CL was frozen immediately after cell dispersion for determination of pre-incubation
concentrations of progesterone. Data were analyzed and reported as net biosynthesis (progesterone after 2 h culture - progesterone pre-incubation). For analysis of synthesis of PGF$_2\alpha$ and 6-keto-PGF$_1\alpha$, 1 X $10^6$ cells were incubated in duplicate with treatments for 2 h and frozen at the end of incubation. Treatments included: control, 1 $\mu$g AA, $10^{-10}$ to $10^{-7}$M ET-1, and combinations of each dosage of ET-1 with AA. One set of quadruplicate tubes for each CL was frozen immediately after cell dispersion for determination of pre-incubation concentrations of each PG. Data were analyzed and reported as the difference between pre- and post-incubation values for each PG.

*Assays for Progesterone, PGF$_2\alpha$, and 6-keto-PGF$_1\alpha*$. Luteal and serum concentrations of progesterone were determined by radioimmunoassay as described by Sheffel et al. (1982). The inter- and intra-assay coefficients of variation were 7.1% and 5.5%, and sensitivity was 0.12 ng/mL. Luteal tissue was homogenized in 1 mL phosphate buffered saline per mg tissue. The homogenate was centrifuged at 1,500 x $g$ for 15 min., the supernatant was diluted 1 to 800 in PBS, and 100 $\mu$l diluted supernatant was used in the progesterone assay. Frozen serum was allowed to thaw at 4$^\circ$C and 100 $\mu$l was used in the progesterone assay. Concentrations of progesterone, PGF$_2\alpha$, and 6-keto-PGF$_1\alpha$ from incubation of luteal cells were determined by radioimmunoassays (Beal et al., 1980 and Milvae and Hansel, 1980, 1983; respectively). The inter- and intra-assay coefficients of variation were: progesterone = 8.4% and 9.3%; PGF$_2\alpha$ = 9.8% and 10.4%; and 6-keto-PGF$_1\alpha$ = 11.1% and 9.7%.

*Statistical Analysis*. All data were distributed normally (Shapiro and Wilk, 1965). Total luteal progesterone, luteal progesterone concentration, and luteal concentration of mRNA were examined for relationship to progesterone in serum at h 0 by linear
regression with h 0 concentrations of progesterone in serum. Data for progesterone, 
PGF₂α, and 6-keto- PGF₁α were examined by analysis of variance, using pre-incubation 
congencentrations as covariates, for effects of classification of the cow based upon 
concentration of progesterone in serum at h 0 above or below the overall mean (High or Low, respectively), treatments during incubation and interaction of classification and 
treatment. Each breed was represented in both classes: High, n = 3 Ayrshire, 3 Holstein; 
Low, n = 2 Ayrshire, 4 Holstein. Fogwell et al. (1978) demonstrated that concentrations 
of progesterone in serum decreased approximately 80% during the first hour, and reached 
their lowest concentration by 6 h, after removal of the CL from heifers. Therefore, h 0 
values were not included for examination of serum progesterone after lutectomy so that 
the biological response to exogenous progesterone would not be biased by the initial 
endogenous progesterone. Patterns of progesterone in serum were compared by analysis 
of variance with progesterone at h 0 (High vs. Low) in the main plot and time by 
classification in the sub-plot. Areas under the curve for serum progesterone after 
lutectomy were compared with concentration of progesterone at h 0 as a covariate. 
Analyses were performed by the SAS® software (SAS Institute; Cary, NC).

Results

Luteal Progesterone and Gene Expression

Size of the CL at ultrasound (mean = 22.4 mm) and luteal wet weight (mean = 
8.55 g) were similar for each treatment. Total luteal progesterone content (mean = 106.0 
± 3.3 μg) and concentration (mean = 12.7 ± 0.5 μg/g) did not differ between High and 
Low cows (Figure 1.1) and neither was associated with h 0 concentrations of 
progesterone in serum (b = 0.36 and 0.11, respectively). Total luteal progesterone was not
associated with luteal wet weight or size at ultrasonography (b = 0.24). Luteal contents of mRNA for enzymes from the endothelin and prostaglandin pathways were not associated with h0 progesterone and did not differ between classes (Figures 1.2 and 1.3, respectively).

**Luteal Cell Incubation**

Hour 0 progesterone and net progesterone biosynthesis during 2 h incubation without treatment (control) did not differ for luteal cells from High and Low cows (Figure 1.5a). Incubation with 5 ng bovine LH increased (P < 0.05) net biosynthesis of progesterone above control with no difference between classes (Figure 1.5a). Incubation with 1 μg AA alone did not alter biosynthesis of progesterone (Figure 1.5a). Both basal and LH-stimulated secretion of progesterone from luteal cells were inhibited by ET-1 in a dose-dependent manner (P < 0.0001; Figures 1.5b and c, respectively). A class effect was detected when ET-1 was added in the presence of AA; in that situation, reduction of progesterone production by ET-1 was greater (P < 0.05) for luteal cells of Low cows (Figure 1.5d).

Initial, control, and AA-stimulated concentrations of PGF$_2$α (Figure 1.6a) and 6-keto-PGF$_1$α (Figure 1.7a) did not differ for luteal cells from High and Low cows. Concentrations of PGF$_2$α were increased (P < 0.0001) in a dose-dependent manner by addition of ET-1 or ET-1 and AA to incubation media with no difference between classes (Figures 1.6b and c, respectively). In contrast, concentrations of 6-keto-PGF$_1$α were decreased (P < 0.0001) in a dose-dependent manner by addition of ET-1 or ET-1 and AA to incubation media, and again, there was no difference between High and Low cows (Figure 1.7b and c, respectively).
**Serum Progesterone Post-Luteectomy**

Patterns of concentrations of progesterone in serum after CL removal differed (P < 0.0001). Concentrations were greater and increased more after 36 h for cows in the High than in the Low group during twice-daily injections of exogenous progesterone (150 mg s.c.; Figure 1.4). Area under the curve for serum progesterone, not including h 0 values, ranged from 125 to 36 (mean = 66 arbitrary units) for individual cows and differed between treatment groups (P = 0.03) Area under the curve was correlated positively (b = 0.74; P = 0.006) with h 0 progesterone in serum. Plasma concentrations of progesterone, either before or after lutectomy, were not associated with milk production.

**Discussion**

Luteal content and concentration of progesterone did not differ for cows with High versus Low circulating concentrations of progesterone. Stormshak et al. (1963) measured the association of luteal progesterone content and concentration with ovarian veinous drainage throughout the estrous cycle of ewes. They found that luteal progesterone content increased along with total luteal weight. Concentration of progesterone in ovarian effluent blood was correlated positively with total luteal content of progesterone. By sampling over time, they calculated secretory rate of luteal progesterone and demonstrated that total luteal progesterone content was indicative of luteal secretion of hormone into the blood (Stormshak et al., 1963). Gomes et al. (1963) found that jugular blood progesterone was not predictive of the quantity of progesterone in the CL of non-pregnant dairy cows during the estrous cycle. These reports demonstrate that luteal progesterone content at a single time point indicates luteal function more specifically than concentrations of progesterone in peripheral serum, and that was confirmed in the present study.
Expression of mRNAs for genes involved in the endothelin and prostaglandin systems was similar for luteal tissue from High and Low cows. The mRNA for preproendothelin-3 was less abundant than preproendothelin-1 and endothelin receptor-A was more abundant than endothelin receptor-B. These data correspond well with previous reports of relative expression of genes from the endothelin family in cow CL (Choudhary et al., 2004). Interestingly, the expression of aldoketoreductase 1B5 (AKR1B5), the prostaglandin F synthase (PGFS) considered important for synthesis of PGF$_2\alpha$ in the cow CL during luteal regression (Arosh et al., 2004), was relatively low compared to other genes of the prostaglandin system measured in this experiment (Figure 1.3). Costine et al. (2007) reported that the expression of AKR1B5 was similar in the CL of pregnant and non-pregnant ewes before PGF$_2\alpha$ injection on d 12 post-estrus, but that its expression decreased in non-pregnant, but not in pregnant ewes 4 h after injection of PGF$_2\alpha$.

Abundance of AKR1B5 mRNA in CL of cows was greater on d 16 than d 5 and greater in d 5 short-lived CL than in d 5 normal CL (Costine, 2004). Arosh et al. (2004) found a shift in expression of AKR1B5 relative to expression of PGES that favored PGES in cow CL during d 10 to 13 of the estrous cycle. The actual quantity that could be expected in CL of pregnant cows is not clear. However, the relatively low abundance reported here might be specific to CL of pregnancy and could indicate an important self-protective mechanism.

Endothelin-1, either alone or in combination with AA, decreased basal and LH-stimulated progesterone production by dispersed luteal cells in a dose-dependent manner. This result corresponds with similar in vitro experiments conducted with luteal cells from the mid-luteal phase of the bovine estrous cycle (Girsh et al., 1996a; Sen et al., 2005; Sen
et al., 2006). Moreover, luteal cells from Low cows incubated with ET-1 and AA produced less progesterone than luteal cells from High cows. Addition of AA to incubated luteal cells in culture increased secretion of anti-steroidogenic PGF$_2\alpha$, which might have occurred by shifting the enzymatic conversion away from the luteotrophic prostaglandin, PGI$_2$, as indicated by decreased accumulation of its metabolite 6-keto-PGF$_{1\alpha}$ (Figures 1.6 and 1.7). However, there was no significant effect of class on stimulation of PGF$_2\alpha$ or inhibition of PGI$_2$ by ET-1.

Alternatively, luteal cells from Low cows might have less capacity to catabolize PGF$_2\alpha$ into 13,14 dihydro 15-keto PGF$_2$ (PGFM) compared to cows with high serum progesterone. Corpora lutea from pregnant ewes on d 14 had a greater capacity to catabolize PGF$_2\alpha$ into PGFM than CL from non-pregnant ewes in spite of the fact that mRNA expression for PGDH did not differ (Costine et al., 2007). However, the hypothesis that increased PGDH activity is responsible for the class difference in progesterone in response to ET-1 and AA in this study was not supported because there was not a class difference in either mRNA for PGDH or luteal secretion of PGF$_2\alpha$. The data presented here indicate that luteal secretion of progesterone, and the systems known to alter it, function similarly for cows with high and low peripheral serum concentrations of progesterone during gestation. Changes in factors that modulate luteal function throughout the estrous cycle do not appear to explain lower serum concentrations of progesterone in some pregnant cows around d 30 of gestation.

Concentrations of serum progesterone during replacement with a dosage of exogenous progesterone known to maintain pregnancy after lutectomy (Bridges et al., 2000; Starbuck et al., 2004) were greater for cows in the High than in the Low class.
Earlier reports have indicated that steroid clearance differs with physiological status. For example, although ovulatory follicles and corpora lutea were smaller in heifers, circulating concentrations of progesterone and estradiol were greater in heifers than in lactating cows (Wiltbank et al., 2006). Luteal size and follicular diameter also were smaller for non-lactating versus lactating dairy cows without differences in serum concentrations of progesterone or estradiol (Sangsritavong et al., 2002). In their review, Wiltbank and coauthors (2006) calculated an “index of circulating progesterone concentration divided by luteal volume” and found that heifers had twice the calculated value of lactating cows. This ratio was related more closely to milk production than circulating steroid concentration (Wiltbank et al., 2006).

The effects of dry matter or energy intake on steroid metabolism have varied with method of investigation. By calculating energy balance as daily net energy intake minus net energy required for maintenance and lactation, Villa-Godoy et al. (1988) determined that milk progesterone was reduced by caloric deficit in lactating dairy cows. Moreover, the degree of this reduction was modulated by the timing and magnitude of maximal caloric deficit. In contrast, Sangsritavong et al. (2002) reported an acute increase in liver blood flow and metabolic clearance rate for infused progesterone and estradiol-17β in lactating and non-lactating dairy cows fed above maintenance. Rabiee et al. (2002c) found greater serum concentrations of progesterone for feed-restricted compared to ad libitum-fed, non-lactating, ovariectomized cows when exogenous progesterone was supplied by either 1 or 2 simultaneous CIDRs. Furthermore, concentrations of serum progesterone in non-lactating, ovariectomized cows receiving subcutaneous injections of 200 mg progesterone daily and fed at 1/2-maintenance were greater than in similarly
treated cows fed at maintenance (Rabiee et al., 2001c). However, serum, milk and fecal concentrations of progesterone, in lactating dairy cows treated with CIDRs, were not affected by intakes of dry matter and metabolizable energy (Rabiee et al., 2002b). The bulk of evidence indicates that increased dry matter and metabolizable energy intake is negatively associated with steroid metabolism. Whether this effect is solely the result of increased liver blood flow is yet to be determined. Greater feed intake decreased concentrations of progesterone in the serum of ewes during the estrus cycle (Parr et al., 1993; Kiyma et al, 2004), increased liver blood flow (Parr et al., 1993), and increased the concentration of mixed function oxidase enzymes in liver biopsies (Thomas et al., 1987). Smith et al. (2006) hypothesized that fatty acids could alter metabolism of progesterone by the liver. They showed a transient difference in serum concentrations of progesterone between acetate or propionate gavaged ewes receiving intramuscular injections of progesterone. However, their results were not confirmed by Lemley et al. (2007) who fed acetate or propionate and injected progesterone intravenously.

Indirect indicators of increased steroid metabolism for lactating dairy cows have been reported. Duration of estrus was reduced for cows producing above the herd average milk yield as compared to those producing below herd average (Lopez et al., 2004). Additionally, high-producing cows had a larger ovulatory follicle, but lower serum estradiol than low-producing cows. Fricke and Wiltbank (1999) reported that occurrence of double ovulation was 13 percentage points greater for cows with above average milk production compared to below average contemporaries. In a retrospective study, Lopez et al. (2005) demonstrated that development of multiple dominant follicles was associated with greater LH and FSH during the 24-h period prior to follicular selection. Circulating
concentration of progesterone was lower for these cows, and the authors hypothesized that this decrease caused increased gonadotropin secretion, consistent with the data of Kinder et al. (1996) and Taft et al. (1996).

On the basis of the present data, clearance was a more important determinant of peripheral concentrations of progesterone than luteal production. This finding supports the hypothesis that low circulating concentrations of progesterone, known to affect fertility among lactating dairy cows, result from altered nutritional demands. It stands to reason that the factors controlling rate of steroid clearance for lactating dairy cows would remain similar during the estrous cycle and early to mid gestation. Dry matter and energy intake were not controlled or measured in this experiment. The fact that serum concentration of progesterone before and after lutectomy was not correlated with milk production in this study might reflect the degree of variation and the relatively small number of cows studied. However, the finding that injected progesterone supported greater circulating concentrations of progesterone in cows with greater serum progesterone prior to lutectomy compared to those with lower concentrations, is strong evidence that metabolic clearance is responsible for lower peripheral concentrations of progesterone in some cows.
<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer Sequence</th>
<th>Accession number</th>
<th>Product Size (bp)</th>
<th>Annealing temperature (°C)</th>
<th>Intra-assay CV&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
</table>
| GAPDH                             | Forward 5' AATATCATCCCTGCTTCTACTGG 3'  
Reverse 5' CATACTTGAGGGTTTTCACA 3' | gi:77404272      | 154               | 57                         | 6.1<sup>b</sup>             |
| Cyclooxygenase-2                  | Forward 5' ATGTATCCTCCCACGTCAAAG 3'  
Reverse 5' TGGTCCCAGCAGCAAATG 3' | gi:1703495       | 146               | 53                         | 10.1                        |
| Aldoketoreductase 1B5             | Forward 5' AAGTGGAAGCCTGAGG 3'  
Reverse 5' CAGTGGAGGTAGAGGGTC 3' | gi:265403        | 138               | 57                         | 11.3                        |
| Prostaglandin 15-dehydrogenase    | Forward 5' ACCTACCTGGAAGCACGTAACG 3'  
Reverse 5' CTGCGCAGCGTGTCGAATCG 3' | gi:4033852       | 150               | 64                         | 9.3                         |
| Prostaglandin E synthase          | Forward 5' CAACTGAGGGCGAGGGTAGG 3'  
Reverse 5' CCAGGAAACAGGAGGGGTAG 3' | gi:31341986      | 150               | 62                         | 9.8                         |
| Preproendothelin-1                | Forward 5' ATCATCTGGTCAAACGCTTCT 3'  
Reverse 5' TAGCAGACTGGCATCTCTTG 3' | gi:31341389      | 129               | 51                         | 16.5                        |
| Preproendothelin-3                | Forward 5' TTGCGGAGGGAGGAGTTG 3'  
Reverse 5' GCCCTAAGGAACACAAGGCTG 3' | gi:119905653     | 155               | 59                         | 6.5                         |
| Endothelin converting enzyme-1    | Forward 5' GCATCCAATAAGCAGCACGAAGAC 3'  
Reverse 5' ACAGGACAGAGAAGAAGTGC 3' | gi:31341344      | 127               | 58                         | 16.3                        |
| Endothelin receptor-A             | Forward 5' TGTCCCTCTGGGGACTCG 3'  
Reverse 5' TTGTGGGGGATGCTGGTG 3' | gi:27805816      | 144               | 54                         | 6.9                         |
| Endothelin receptor-B             | Forward 5' GCAGGATTTGAGCTACTTCTC 3'  
Reverse 5' TTTTGCTACCAATAAGATCTACAG 3' | gi:31342360      | 150               | 57                         | 9.7                         |

<sup>a</sup> CVs were determined following linearization of C<sub>T</sub> values and were calculated for each sample.  
<sup>b</sup> Value represents the average of GAPDH CV for all samples.
Figure 1.1 Luteal Progesterone
Luteal concentration (μg / gram tissue) and content (μg) of progesterone in CL of cows with High (open bars) or Low (hashed bars) circulating concentrations of progesterone. Means ± SEM.
Figure 1.2 Endothelin Gene Expression
Abundance of mRNA for preproendothelin-1 (ET-1), preproendothelin-3 (ET-3), endothelin converting enzyme (ECE), endothelin receptor A (ET-A), and endothelin receptor B (ET-B) in CL of cows with High (open bars) or Low (hashed bars) circulating concentrations of progesterone. Means ± SEM.
Figure 1.3 Prostaglandin Gene Expression
Abundance of mRNA for cyclooxygenase-2 (COX-2), aldoketoreductase 1B5 (AKR1B5), prostaglandin 15-dehydrogenase (PGDH), and prostaglandin E synthase (PGES) in CL of cows with High (open bars) or Low (hashed bars) circulating concentrations of progesterone. Means ± SEM.
Figure 1.4 Post Luteectomy Serum Progesterone

Patterns of progesterone (ng / mL) in serum, after luteectomy, for cows with High (open circles) or Low (filled circles) circulating concentrations of progesterone prior to luteectomy. Samples were taken every 4 h for 48 h. Progesterone (150 mg) was injected s.c. every 12 h. Means ± SEM.
Figure 1.5 Luteal Cell Progesterone Production
Net biosynthesis of progesterone (ng/10^5 cells/2 h) by dispersed luteal cells, from CL of cows with High (open bars) or Low (hashed bars) circulating concentrations of progesterone, incubated without treatment (a, Control) or with 1 μg arachidonic acid (a, AA), 5 ng bovine luteinizing hormone (a, LH), LH and AA (a) increasing concentrations of endothelin-1 (b, 10^{-7} M ET to 10^{-10} M ET), or combinations of LH or AA and increasing concentrations of ET (c, ET+LH and d, ET+AA; respectively). Means ± SEM.
Figure 1.6 Luteal Cell PGF$_2\alpha$ Production

Net production of prostaglandin F$_2\alpha$ (PGF$_2\alpha$; ng / 10$^6$ cells / 2 h) by dispersed luteal cells, from CL of cows with High (open bars) or Low (hashed bars) circulating concentrations of progesterone, incubated without treatment (a, Control) or with 1 $\mu$g arachidonic acid (a, AA), increasing concentrations of endothelin-1 (b, $10^{-10}$ M ET to $10^{-7}$ M ET), or the combination of AA and increasing concentrations of ET (c, ET+AA). Means ± SEM.
Figure 1.7 Luteal Cell 6-keto-PGF$_1\alpha$ Production

Net production of 6-keto-prostaglandin F$_1\alpha$ (6-keto-PGF$_1\alpha$; ng / $10^6$ cells / 2 h) by dispersed luteal cells, from CL of cows with High (open bars) or Low (hashed bars) circulating concentrations of progesterone, incubated without treatment (a, Control) or with 1 µg arachidonic acid (a, AA), increasing concentrations of endothelin-1 (b, $10^{-10} M$ ET to $10^{-7} M$ ET), and combination of AA and increasing concentrations of ET (c, ET+AA). Means ± SEM.
ASSOCIATION OF TIMING OF CHORIOALLANTOIC MEMBRANE DEVELOPMENT WITH AGE IN DAIRY CATTLE

Abstract

About 12% of lactating dairy cows experience late embryonic or early fetal death (d 30 to 60 of gestation). These losses are economically important because lengthened interestrous intervals lead to longer calving intervals. Abnormal growth of the allantois and timing of its fusion with the chorion have preceded high rates of late embryonic / early fetal loss of manipulated embryos. This study characterized timing of detection of the allantois and diameter of the allantoic lumen, in dairy cows and heifers with embryos developed entirely in vivo, to determine whether these variables might be used to investigate late embryonic / early fetal loss. Reproductive tracts of dairy heifers (n = 33) and lactating dairy cows (n = 30) were examined daily, via transrectal ultrasonography (Aloka 900), beginning on d 21 post-insemination (PI). Variables included: first day of detection of the allantois, diameter of the allantoic lumen, length of the embryo at first detection of the allantois, and pregnancy retention on d 60 PI. Range and mean of first day of detection of the allantois (d 21 to 26, mean = 23 ± 0.18 PI) agreed with previous reports. The allantois was detected earlier (P < 0.05) in heifers (22.4 ± 0.2 d) than in cows (23.6 ± 0.2 d). Diameter of the allantoic lumen at first detection (4.6 ± 0.3 mm) did not differ with day of detection and was not affected by age of dam. In contrast, length of embryo (range 2.6 to 7.9 mm) varied with day of first detection of the allantois (P < 0.05), but not with age of dam. Pregnancy failure by day 60 PI did not differ with age of the dam. Differences in failure for pregnancies in which the allantois was detected earlier (≤ d 23 PI) or later (≥ d 24 PI) approached significance (P = 0.08) with those detected earlier failing more frequently than those detected later. Differences in day of first
ultrasonographic detection of placental membranes, between heifers and cows, might reflect differing developmental rates and, based on present data, asynchrony in development of the embryo and the placental membranes might occur in some animals. Therefore, day of detection of the allantois might be useful for predicting survival or loss of pregnancy during placentation and analysis of molecular characteristics of the developing allantois could help to identify mechanisms that compromise late embryonic / early fetal survival.

**Introduction**

Late embryonic / early fetal pregnancy failure (d 25 to 60 of gestation) in dairy cows leads to lengthened interestrous intervals and more days in milk before a successful pregnancy is established. The frequency of pregnancy loss during this period of gestation has been reviewed thoroughly (Kummerfield et al., 1978; Sreenan and Diskin, 1986; Dailey et al., 2002; Inskeep 2004; Inskeep and Dailey, 2005; and Diskin et al., 2006). Estimates ranged from 7 to 25% and are generally considered to occur at an approximate rate of 12%.

Abnormal development of the allantois has been reported in situations associated with very high pregnancy loss for *in vitro*-produced embryos and somatic-cell-nuclear-transfer, cloned pregnancies (Thompson and Peterson, 2000, Piedrahita et al., 2002). However, association of pregnancy survival or loss with allantoic development is not known for embryos developed entirely *in vivo*. When genetic aberrations that lead to poor allantoic development are elucidated, their patterns of expression can be compared in pregnancies derived by artificial insemination that are more likely to either succeed or fail during placentation. However, at this time, methods to determine the pattern of
allantoic development without termination of the pregnancy have not been reported. This type of investigation might be facilitated by using ultrasonography to evaluate allantoic development.

Transrectal ultrasonography has been used successfully to determine the first day of appearance of the allantois in cows and heifers. On average, the allantois was first detected on day 23 post-insemination (PI) by Curran and colleagues (1986b) and by Boyd et al. (1988). Parity of the dam affected the timing of detection of the allantois by Kolour et al. (2005). They first observed the allantois on days 22, 23 and 26 in heifers, primiparous and multiparous cows, respectively. The first day of detection of the allantois was similar for heifers and primiparous cows and each was statistically different than multiparous cows.

Development of the allantois can influence pregnancy maintenance during placentation and can be monitored by transrectal ultrasonography in cows and heifers. Furthermore, the range of days PI that the allantois is first detected by ultrasound, and evidence that age of the dam affects this timing, indicate inherent variability in allantoic development that might be associated with its ability to supply adequate vasculature to the placenta. Therefore, the objective of this study was to characterize the timing of detection of the allantois and diameter of the allantoic lumen, in dairy cows and heifers, to determine whether these variables might be used to investigate effects on late embryonic / early fetal loss.

**Materials and Methods**

Beginning on d 21 PI, the reproductive tracts of 33 nulliparous heifers and 30 parous cows (Holstein and Ayrshire) were examined daily with a real-time, B-mode
ultrasound console equipped with a linear array, 7.5 MHz transducer (ALOKA SSD-900; Aloka Co., Ltd.; Tokyo, Japan). Day 0 was designated as the day of insemination. The first day of detection of the allantois, diameter of the allantoic lumen, length of the embryo proper at first detection of the allantois, and pregnancy retention on d 60 PI were recorded. Detection of the allantois occurred when a circular line of echo-producing tissue appeared in the uterine lumen, sharing a field of view with the embryo (Fig. 2.1). All anatomical measurements were taken with electronic calipers at the time of ultrasonography. Diameter of the allantoic lumen was recorded as the average of the vertical and horizontal inside diameters at the widest point sharing a field of view with the embryo proper. Statistical analyses were performed by the SAS® software (SAS Institute; Cary, NC). Analysis of variance was used to examine the effect of age on day of detection of the allantois, diameter of the allantois, and length of the embryo proper. Linear regression was used to examine association of length of the embryo or diameter of the allantoic lumen with day the allantois was detected for cows and heifers combined. Differences in pregnancy retention on d 60 PI between cows and heifers and between pregnancies for which the allantois was detected early (≤ d 23 PI) or late (≥ d 24 PI) were examined by Chi square frequency analysis.

**Results**

Range and mean of first day of detection of the allantois (d 21 to 26, 23 ± 0.18 PI) agreed with previous reports (Curran et al., 1986, Boyd et al., 1988 and Kolour et al., 2005) and were compatible with descriptions of embryonic gross anatomical development (Greenstein et al., 1958). The allantois was detected earlier (P < 0.05) in heifers (22.4 ± 0.2 d) than in cows (23.6 ± 0.2 d). Diameter of the allantoic lumen at first
detection (4.6 ± 0.31 mm) did not differ with day of detection and was not affected by age of dam (Table 2.1). In contrast, length of embryo (range 2.6 to 7.9 mm) increased 0.37 mm with each additional day to first detection of the allantois (b = 0.37, P < 0.05), but was not affected by age of dam. Overall pregnancy loss by d 60 PI averaged 11% (7 of 63) and did not differ with age of the dam (cows = 3 of 30; heifers = 4 of 33). The difference in pregnancy loss by d 60 PI between pregnancies for which the allantois expanded earlier (≤ d 23 PI; 16%, 7 of 45) and those for which the allantois expanded later (≥ d 24 PI; 0%, 0 of 18) approached significance (P = 0.08).

**Discussion**

The first day of detection of the allantois was earlier for heifers than cows. Kolour and coworkers (2005) also imaged the allantois earlier in heifers and primiparous cows than in multiparous cows. They postulated that earlier detection in heifers and primiparous cows was easier because of the relatively small size of the uterus (not quantified). However, that hypothesis excludes the possibility that extraembryonic membranes develop sooner for embryos in younger dams. In the present data, regardless of age, size of its lumen was not associated with earlier detection of the allantois. This might indicate that, because the younger allantois was not smaller at first detection, earlier detection is a function of more rapid embryonic development for pregnancies gestated in younger dams.

Differences in concentration of serum progesterone affect early embryonic development. Administration of progesterone to ewes early after estrus (d 0 – 3) advanced the development of the uterus such that, on d 6 after estrus, it provided a suitable environment for continued development of a d 10 embryo (Lawson and Cahill,
1983 and Vincent et al., 1986) but not for a d 6 embryo. Administration of supplemental progesterone during d 1 to 4 and 5 to 9 post-insemination increased length of trophoblast on d 14 and 16 in cyclic beef and lactating dairy cows (Garrett et al., 1988 and Mann et al., 2006; respectively). Embryonic development was retarded on d 16 in non-lactating dairy cows with a delayed increase in serum progesterone compared to those that reached a concentration of 1 ng/mL by d 4 post-insemination (Mann and Lamming, 2006). The mediator for the effect of advanced embryonic development by increased progesterone concentration is likely histotrophic support from the uterine endometrium. Satterfield et al. (2006) demonstrated that augmenting serum concentrations of progesterone beginning 36 h after mating through d 9 or 12 increased the rate of blastocyst development and the concentration of uterine histotroph in ewes. The tendency for pregnancy failure to occur more frequently for pregnancies in which the allantois developed early might be explained by a discrepancy in the timing of attachment. If the conceptus begins attachment before the uterine endometrium is prepared to accept it, placentation could fail. The fact that the allantois was detected earlier in heifers than cows while pregnancy failure did not differ with age, might either be due to a wider window of uterine receptivity for heifers or simply an inadequate sample size to test this variable.

Regardless of age of dam, length of the embryo continued to increase with each additional day to detection of the allantois, despite the fact that the allantois was similar in size regardless of the day that it was detected. This might indicate that some pregnancies exhibit an asynchrony in development of the embryo and placental membranes. None of the studies reviewed here, using transrectal ultrasonography to characterize embryonic development, noted a relationship between the size of the
allantois and length of the embryo. Maddox-Hyttel et al. (2003) examined seven embryos collected from two superovulated heifers slaughtered 21 d PI. Length of the embryo ranged from 3.14 to 5.79 mm while the allantois ranged in width from 1.15 to 4.41 mm. The authors noted “a clear relationship between the embryo length and width of the allantois,” but provided no statistical analysis to support the comment. The allantois had not yet developed for two embryos that were 3.14 and 3.77 mm long (Maddox-Hyttel et al., 2003).

Placental development was altered for embryos derived by assisted reproductive technologies such as in vitro fertilization (IVF) and somatic-cell-nuclear-transfer (SCNT) cloning. In fact, a majority of the embryonic mortality between days 30 and 60 of gestation for IVF- and SCNT-derived pregnancies was attributed to improper development of the allantois (Hill et al., 2000). Using serial slaughter of recipient cows that received an IVF embryo or were inseminated artificially, it was determined that the degree of malformation of the allantois ranged from aplasia to normal growth and vascularization without haematopoesis (Piedrahita et al., 2002). Furthermore, size of the follicle donating an oocyte for IVF affected placental development of the resulting embryo (Piedrahita et al., 2002). The length and width of the allantois on day 27 of gestation were smaller for embryos derived from small follicles. However, length and width of the embryo proper or the degree of vascularization of the allantois did not differ in relation to follicular size. In addition to differences in growth rate of the allantois, these authors reported a difference in the ability of the allantois to fuse with the chorion. For all 17 conceptuses derived from small follicles, the allantois and chorion detached during collection. In contrast, 14 of 18 conceptuses derived from large follicles presented
fully fused chorioallantoic membranes that could not be detached manually after collection (Piedrahita et al., 2002). These findings hold particular interest in relation to the reports of increased embryonic loss for cows bred to ovulation of immature (Perry et al., 2005) and persistent (Ahmad et al., 1995) follicles.

Whether the degree of allantoic or chorionic vascularization could be assessed by color Doppler ultrasound is yet to be determined. However, if growth characteristics of the allantois, as determined by transrectal ultrasonography, are indicative of other distinct functions of allantoic development, it might prove to be a beneficial tool for investigating pregnancy failure during placentation. Monitoring the development of embryos derived by IVF and SCNT cloning by ultrasound could provide a means to determine these characteristics without terminating the pregnancy and allow testing for an association of the eventual sustainability of that specific conceptus with the developmental traits. It might also provide a method to select tissues that exhibit an abnormal growth pattern for molecular investigations.

Several genes have been identified as differentially expressed during allantoic emergence in bovine and ovine conceptuses (Ledgard et al., 2006). These authors used suppressive subtractive hybridization to compare mRNA expression in preimplantation conceptuses prior to emergence of the allantois and in allantoic samples after emergence. Nine genes were identified as differentially expressed. All were associated with extracellular matrix and vasculogenesis. Similarly, Ushizawa et al. (2004) used microarray analysis of cDNA to determine gene expression profiles in the bovine embryo during the preattachment period. They found differential expression of more than 1,000 genes on days 7, 14, 21 and 28. Comparison of extra-embryonic membranes and the
embryo proper on day 28 revealed 119 down-regulated genes and 74 up-regulated genes in the membranes. The up-regulated genes included placental lactogen, prolactin-related proteins, pregnancy-associated glycoproteins and major histocompatibility complex molecules.

Collectively, results reported here illustrate that earlier ultrasonographic detection of the allantois in heifers than cows might represent an increased rate of embryonic development rather than detection of smaller structures. The relatively wide range for first day of detection of the allantois indicates that the factors controlling its development might be influenced by differing physiological states associated with age or lactational status. One factor that could mediate the effects of age and lactation on allantois development is concentration of serum progesterone. Low circulating concentrations of progesterone before breeding, during the first several days post-estrus, and during placentation have been identified as causes of embryonic loss during and after these respective periods and occur frequently in lactating dairy cows. Moreover, the use of transrectal ultrasonography to monitor development of the allantois between days 21 and 26 post-insemination could prove to be a valuable tool for investigating pregnancy failure at placentation.
### Table 2.1 Allantois Results Relative to Age of Dam

<table>
<thead>
<tr>
<th>Age</th>
<th>Day of Detection</th>
<th>Diameter of Allantois</th>
<th>Length of Embryo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heifer</td>
<td>22.4 ± 0.2&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4.79 ± 0.43 mm&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4.96 ± 0.27 mm&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Cow</td>
<td>23.6 ± 0.3&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.23 ± 0.49 mm&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4.75 ± 0.23 mm&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Differing superscripts within columns indicate significance at P < 0.05.
Figure 2.1 Representative Ultrasonographic Image
Representative image of the allantois (white arrow) and embryo proper (black arrow) on the first day of detection of the allantois at d 23 PI.


Smith, M. W., and J. S. Stevenson. 1995. Fate of the dominant follicle, embryonal survival, and pregnancy rates in dairy cattle treated with prostaglandin F2α and


