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Expression of estrus modifies the gene expression profile in reproductive tissues on Day 19 of gestation in beef cows

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ABSTRACT

The aim of this study was to test the effect of expression of estrus at artificial insemination (AI) on endometrium, conceptus, and CL gene expression of beef cows. Thirty-six multiparous nonlactating Nelore cows were enrolled on an estradiol- and progesterone (P4)-based timed AI protocol (AI = Day 0) and then slaughtered for the endometrium, CL, and conceptus collection on Day 19. The animals were retrospectively grouped on the basis of cows that (1) showed signs of estrus near AI (n = 19; estrus) and (2) did not show any signs of estrus (n = 17; nonestrus). Body condition score, blood sampling, and ultrasound examination were performed on Days 0, 7, and 18 of the experiment followed by messenger RNA extraction and quantitative reverse transcription polymerase chain reaction analysis of 58 target genes. Data were checked for normality and analyzed by ANOVA for repeated measures using proc GLM, MIXED, and UNIVARIATE of SAS. Only pregnant cows were included in the analyses (n = 12; nonestrus, n = 11). Estrous expression had no correlation with parameters such as body condition score, preovulatory follicle and CL diameter, P4 concentration in plasma on Days 7 and 18 after AI, and interferon-tau concentration in the uterine flushing (P > 0.15); however, a significant increase was observed in conceptus size from cows that expressed estrus (P = 0.02; 38.3 ± 2.8 vs. 28.2 ± 2.9 mm). The majority of transcripts affected by estrous expression in the endometrium belong to the immune system and adhesion molecule family (MX1, MX2, MYL12A, MMP19, CXCL10, IGLL1, and SLPI; P ≤ 0.05), as well as those related with prostaglandin synthesis (OTR and COX-2; P ≤ 0.05). Genes related to apoptosis, P4 synthesis, and prostaglandin receptor were downregulated (CYP11A, BAX, and FPR; P < 0.05) in the CL tissue of cows that expressed estrus. In addition, four genes were identified as differentially expressed in the 19-day-old conceptus from cows that expressed estrus (ISG15, PLAU, BMP15, and EEF1A1; P < 0.05). There was also a significant effect of Day 7 concentration of P4 mainly affecting the immune system, adhesion molecules, and wnt signaling pathway of the endometrium (IGLL1, MX2, SLPI, TRD, APC, WNT2, GLYCAM1, and MYL12A; P < 0.05). A significant interaction between estrous expression and P4 concentration on Day 7 was more pronounced in immune system genes (MX1, MX2, TRD, SLPI, and IGLL1; P < 0.05). This study reported that estrous expression at the time of AI favorably altered the gene expression profile in reproductive tissues during the preimplantation phase toward a more receptive state to the elongating conceptus. These effects seem to be more evident in the endometrium during the time of dynamic remodeling for embryo implantation.

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1. Introduction

Early and late embryonic loss occurs mainly in the first 6 weeks of gestation and is responsible for major losses in the beef and dairy industry. A great proportion of these embryonic losses occur between Days 8 and 21 after fertilization [1,2]. The effect of changes in steroid hormone concentrations is critical as they affect the ability of the endometrium to receive and maintain the conceptus. Previous studies have reported the correlation between the concentration of estradiol (E2) in plasma and the ovulation, increased pregnancy/artificial insemination (AI), and decreased pregnancy loss in beef and dairy cattle [3,4].

Estradiol initiates crucial modifications in the endometrium environment such as increased epithelial cell height and ciliation in the fimbria [5] and ampulla [6]. Indeed, E2 concentrations during the proestrus period are positively correlated with the diameter of the preovulatory follicle, subsequent CL diameter, concentration of progesterone (P4) during diestrus [7], and conception rates in dairy cows [8,9]. Pereira et al. [10] also reported that a shorter proestrus duration decreased conception rates even when embryo transfer technology was used. Furthermore, an increase in pregnancy maintenance from Days 7 to 27 after AI was observed when serum E2 concentration on Day 0 and P4 concentration on Day 7 were greater in recipient cows [11].

Comparing transcriptome of the receptive and non-receptive endometrium has led to identifying signaling pathways involved in embryonic growth and development [12]. Before implantation, during the receptivity phase of the endometrium, specific genes related to the immune system, adhesion molecules, and developmental genes are extensively regulated [12,13]. Some of these genes are activated once the conceptus starts secreting interferon-tau (IFNT), but the timing of this activation varies considerably.

Immunologically, the embryo is an allograft for the dam and more specifically for the uterine tissue. Therefore, a complex modulation of immune cells and its signals are necessary to allow the maintenance of the conceptus. The uterus is an immunologically privileged site [14], and E2 has shown to play an important role by upregulating *SERPINA14* messenger RNA (mRNA) synthesis during estrus [15]. On the basis of studies performed in sheep [16], this serpin family member has immunomodulatory roles which include (1) blocking T cell proliferative responses [17], (2) impairing natural killer cell activity [18], and (3) decreasing antibody production [19]. A second group of genes critical for the survival of the early embryo are related to cell adhesion. Proper attachment and invasion of the embryo in the endometrium depend on adhesion-related molecules. In ruminants, the fetal tissue invades [20] the endometrium and establishes a synepitheliochorial type of placentation [21]. Apposition, adhesion, and invasion performed by the conceptus are controlled by the endometrium [22]. Studies have found upregulation of some adhesion molecules such as *SPP1* and *GLYCAM1* during the implantation phase in ruminants [23,24]. The canonical wnt signaling pathway, which is regulated by sexual steroids including E2 [25], is critical for morphogenesis and development of the preimplanted conceptus [26,27]. The wnt regulatory role in embryonic development is still unknown as previous

studies have shown that the wnt activation improves [28], reduces [29], or has no effect [30] on the proportion of embryos that can develop to the blastocyst stage.

The function of the CL and consequent P4 synthesis in the preimplantation phase is key for proper embryo elongation and IFNT synthesis. However, it is unclear whether estrous expression could further modify the transcriptome of the CL. It is reasonable to believe that a fully mature preovulatory follicle could improve the chances for a more developed CL.

The objective of this study is to test the effects of behavioral expression of estrus before AI on gene expression of target transcripts in the endometrium, CL, and conceptus on Day 19 of gestation. We hypothesized that expression of estrus is associated with a complete maturation and function of the preovulatory mechanisms, therefore improving the transcriptome profile in reproductive tissues during the preimplantation phase.

2. Materials and methods

2.1. Animals and housing

Thirty-six nonlactating multiparous Nelore cows (body condition score [BCS] = 5.5 ± 0.1) [31] were assigned to an estrous synchronization plus timed AI protocol [32] (Fig. 1). All animals were cycling and with absence of any clinical disorder. The animals were between 48 and 72 months of age. The animals were not lactating at the time of study, and previous parturition occurred 300 to 360 days before enrollment. The cows were enrolled onto a synchronization protocol that was carried out as follows: 2-mg injection of estradiol benzoate (Estrogin; Farmavet, São Paulo, SP, Brazil) and a second-use (previously used for 9 days) intravaginal P4-releasing device (CIDR, originally

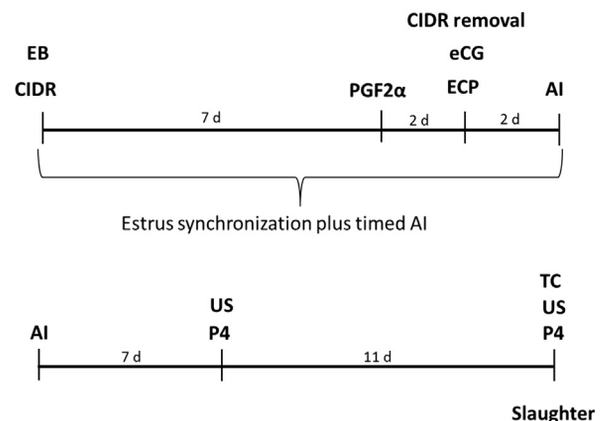


Fig. 1. Diagram of study. Cows received a 2 mg injection of estradiol benzoate (EB, Estrogin; Farmavet, São Paulo, SP, Brazil) and a second-use intravaginal progesterone-releasing device (CIDR, originally containing 1.9 g of progesterone; Zoetis, São Paulo, Brazil) on study Day -11, a 12.5-mg injection of PGF2 α (Lutalyse; Zoetis, São Paulo, Brazil) on Day -4, CIDR removal in addition to 0.6 mg of estradiol cypionate (ECP; Zoetis, São Paulo, Brazil) and 300 IU of eCG (Novormon, Schering-Plough Co., São Paulo, Brazil) on Day -2, and timed artificial insemination (AI) on Day 0. P4, blood collection for progesterone analysis, TC, tissue collection; US, ultrasonographic examination of ovaries.

containing 1.9 g of P4; Zoetis, São Paulo, Brazil) on study Day –11, a 12.5-mg injection of PGF2 α (Lutalyse; Zoetis) on Day –4, CIDR removal in addition to 0.6 mg of estradiol cypionate (Zoetis) and 300 IU of eCG (Novormon; Schering-Plough Co., São Paulo, Brazil) on Day –2, and timed AI on Day 0. All cows were inseminated on Day 0 by the same technician, using semen from the same bull and batch. The cows were maintained in a single *Brachiaria brizantha* pasture (10 ha) with ad libitum access to forage and water. All animals received a 100 g of a protein–mineral mix + 100 g of ground corn per cow daily (on an as-fed basis).

The cows were observed for behavioral expression of estrus by visual observation twice a day for 30 minutes each from the administration of the PGF2 α injection until timed AI. The cows were visually observed for mounting activity and secondary signs of estrus (e.g., chin rest, following, vaginal mucus, swollen vulva) and then clustered in two different groups (1) estrus (n = 19), when cows expressed evident signs of estrus the day before (PM) and/or the day of AI (AM), and (2) nonestrus (n = 17), for cows that did not show any signs of estrus. To clearly define the subgroups, only animals that were positive (estrus) or negative (nonestrus) for mounting activity and secondary signs of estrus were considered for this study (n = 36), whereas animals that were positive for only primary or secondary signs were removed from the project. Only pregnant cows were then included in the analyses (estrus, n = 12; nonestrus, n = 11).

2.2. Blood samples and ultrasound examinations

Blood samples were collected immediately before AI (Day 0) and on Days 7 and 18 of the experiment via jugular venipuncture into commercial blood collection tubes (Vacutainer, 10 mL; Becton Dickinson, Franklin Lakes, NJ, USA) containing 158 USP units of freeze-dried sodium heparin. After collection, the blood samples were placed immediately on ice, centrifuged (2500 \times g for 30 minutes, 4 °C) for plasma harvest, and stored at –20 °C on the same day of collection for further analysis of P4 using an ELISA procedure according to manufacturer's guidelines (Ovu-Check Plasma Elisa Kit; Biovet Inc., Saint Hyacinthe, Québec, Canada). Transrectum ultrasonography (7.5-MHz transducer, 500 V; Aloka, Wallingford, CT, USA) was performed concurrently with blood sampling on Days 0, 7, and 18 to verify ovulation and CL development. Corpus luteum volume was calculated using the formula for volume of a sphere: volume = $4/3\pi \times (D/2)^3$, where D is the maximum luteal diameter. All animals analyzed had a preovulatory follicle with the absence of a CL on Day 0, confirmed ovulation on Day 7 (presence of a CL in the ipsilateral ovary of the preovulatory follicle observed on Day 0), and a CL greater than 0.38 cm³ in volume on Days 7 and 18.

2.3. Slaughter and tissue collection

The cows were slaughtered on Day 19 after timed AI, and reproductive tracts were immediately collected, placed on ice, and processed for collection of the conceptus, uterine luminal flushing, and tissue samples

from the CL and endometrium on the basis of the procedures described by Bilby et al. [33]. More specifically, the uterine horn ipsilateral to the CL was isolated from the reproductive tract, and the ovary containing the CL was removed. The CL was incised with a scalpel for collection of luteal tissue. Subsequently, 20 mL of saline were injected into the uterotubal junction of the selected uterine horn, massaged gently, and exited through an incision at the tip of the uterine horn. Uterine luminal flushing media and the conceptus were recovered in a sterile 100 by 15-mm Petri dish. The conceptus was measured for length and weight, whereas the uterine luminal flushing was stored in a 15-mL sterile conical tube (Corning Life Sciences, Tewksbury, MA, USA) for further analysis of IFNT concentrations using a bovine-specific commercial ELISA kit (MyBioSource LLC, San Diego, CA, USA). The selected uterine horn was then cut along the mesometrial border, and samples of the endometrium were collected. After collection, the conceptus, as well as luteal and endometrial samples, were stored in 5-mL sterile cryogenic tubes (CRAL Artigos para Laboratórios, Cotia, São Paulo, Brazil) containing 2 mL of RNA stabilization solution (RNAlater; Ambion Inc., Austin, TX, USA), maintained at 4 °C for 24 hours, and stored at –20 °C until further processing.

2.4. RNA extraction

Total RNA was extracted from samples using the TRIzol Plus RNA Purification Kit (Invitrogen, Carlsbad, CA, USA). The tissue:Trizol ratio (mg:mL) was 100:1 for all samples (1-mL TRIzol per 50- to 100-mg tissue). Quantity and quality of isolated RNA were assessed UV absorbance (NanoDrop 2000; UV-Vis Spectrophotometer; Thermo scientific, Wilmington, DE, USA) at 260 nm and 260:280-nm ratio, respectively. Extracted total RNA was stored at –80 °C until further processing.

2.5. Primer design

All forward and reverse primers were designed from bovine mRNA sequences (National Center for Biotechnology Information) using the PrimerQuest PCR Design Tool (Integrated DNA Technologies, Coralville, IA, USA). The primer sequence, product length, and gene accession number are provided in Table 1.

2.6. Reverse transcription synthesis of cDNA

After extraction, reverse transcription reactions were performed by following the kit manufacturer's protocol. A total RNA sample of 2500 ng was treated with 1- μ L DNase (New England Biolabs, Ipswich, MA, USA) to digest any DNA left from the RNA extraction and were incubated for 10 minutes at 75 °C. Next, to prevent DNase I activity by chelating the divalent cations that it requires (Mg⁺⁺ and Ca⁺⁺), and also to prevent cation-related RNA cleavage, 0.25- μ L EDTA, ultrapure 0.5 M, PH 8.0 (Life Technologies, Burlington, ON, USA) was added to each sample and incubated for 10 minutes at 37 °C. When DNase treatment finished, a High Capacity cDNA Reverse Transcription Kit

Table 1

Primer sequences of analyzed genes from endometrium, CL, and conceptus tissues.

Gene symbol	Accession no.	Primer	Primer sequence	Product length (bp)
Sequences of primers used for qPCR analysis of endometrium tissue				
<i>GAPDH</i>	NM_001034034.2	F R	GAG ATC CTG CCA ACA TCA A CTT CTC CAT GGT AGT GAA GAC	83
<i>LGALSBP3</i>	NM_001046316.2	F R	CTC TGT CTC CTG GTC TTT GGG ATT GGA CTT GGA GTA	127
<i>SERPINA14</i>	NM_174821.2	F R	GAC AGA GTC ACC TCA GAT A CAT CGA GAA TAC CTC CTT TC	91
<i>CLD4</i>	NM_001014391.2	F R	CCC TCA TCG TCA TCT GTA T CCT TGG AGC TCT CAT CAT	99
<i>IDO</i>	NM_001101866.2	F R	AGC TAT GGT CTC CTT GAG GCC TCC AGT TCC TCT ATT	121
<i>MSX1</i>	NM_174798.2	F R	AAG CAG TAC CTG TCC ATC GGT TCT GAA ACC AGA TCT TC	88
<i>SPP1</i>	NM_174178.2	F R	GGA CTT CAC ATC ACA CAT AG CTC GCT ACT GTT GGT TTC	97
<i>IL-10</i>	NM_174088.1	F R	GCT CAG CAC TAC TCT GTT GTT GGC AAG TGG ATA CAG	97
<i>AXIN1</i>	NM_001191398.1	F R	GCC ATC TAC CGC AAA TAC CGA GAT GCA GTC CTT TAT G	93
<i>IGLL1</i>	NM_001083800.1	F R	GGA AGC AGC ACG AAT ATC GGG TCG ATA CTT ATC TTC ATA G	99
<i>TIMP2</i>	NM_174472.4	F R	GGT CAC GGA GAA GAA CAT TCC TCG ATG TCC AGA AAC	126
<i>MX2</i>	NM_173941.2	F R	CCA ATC AGA TCC CGT TCA TGA AGC AGC CAG GAA TAG	115
<i>TRD</i>	XM_603355.3	F R	GTC GCT TGT TTG GTG AAG CCA GGT GAG ATG GCA ATA	104
<i>CDH1</i>	NM_001002763.1	F R	CTG AGA ACG AGG CTA ATG T GGT CTG TGA CGA CGA TAA A	132
<i>RELN</i>	NM_001206458.1	F R	GGG TGT GCC AAT CAA TTC CTG GGT AAC AGC CTT CTT	100
<i>EMMPRIN</i>	NM_001075371.2	F R	GGT CAC CAT CAT CTT CAT CTA AGA GCC TAT GTC TTC ATC ATC	73
<i>LIFR</i>	NM_001192263.1	F R	GCT CTT GGA ATG GGA AAT AG CCA GAC TGA GAT GAG TTA CA	98
<i>SLPI</i>	NM_001098865.2	F R	GCC TTG GAG ATG AGA AAC GGT CCA GAC ATT CAG TTC	96
<i>MYL12A</i>	NM_001015640.2	F R	CAC CAT TCA GGA GGA TTA C GTC AAT AGG TGC TTC TCT G	100
<i>MYH10</i>	NM_174834.1	F R	GAC TAC CAG CGT GAA TTA G CCT GCA ACT GAA GGA TTT	115
<i>MYH9</i>	NM_001192762.1	F R	GAC AAG AGT GGC TTT GAG GTT CAC CTT CAC CTT CTT C	96
<i>IGHG1</i>	DQ452014.1	F R	GAC CCT CTG TCT TCA TCT GTT TAC CTC CAC GTT GTC	146
<i>FDZ4</i>	NM_001206269.1	F R	GTT CCA TCT GGT GGG TTA TTC GCT GCG ATG TGG AAA TAA GA	106
<i>FZD8</i>	XM_005214320.1	F R	CCT ATA TGC CCA ACC AGT TC CAT GCT GCA CAG GAA GAA	122
<i>WNT3</i>	NM_001206024.1	F R	AGA AGC GGA AGG AGA AGT CAC GTC ATA GAT GCG GAT AC	83
<i>AXIN2</i>	NM_001192299.1	F R	GGA GAA ATG CGT GGA TAC TT GTA GAT CGC TTT GGC TAC TC	103
<i>GSK3B</i>	NM_001101310.1	F R	GGG TCA TTT GGT GTC GTG TAT C GAT CTG GAG CTC TCG GTT CTT A	97
<i>GLYCAM1</i>	NM_174828.2	F R	CCT CTG CTC AGT TCA TCA GG TCT GAT CAC AAT TTG CTC TTT GG	97
<i>SELL</i>	NM_001076141.1	F R	GGT GGG AAC CAA CAA ATC CAC AGT CCT CCT TAC TCT TC	86
<i>WNT2</i>	NM_001013001.1	F R	TCC TGT GAC CCA AAG AAG GCA AAC TTG ATC CCA TAG TC	98
<i>CXCL10</i>	NM_001046551.2	F R	GTG TAC CTC TCT CTA GGA ATA C GGA TTG ACT TGC AGG AAT G	107
<i>PTX3</i>	NM_001076259.2	F R	CGC TGA TGC TGT GAT TTC CCA CCG AGT CAC CAT TTA	101
<i>DKK1</i>	NM_001205544.1	F R	CCA TGG GCT GGA GAT ATT GTG AAG CCT GGA AGA ATT AC	100
<i>MMP19</i>	NM_001075983.1	F R	ATC TTG AAC CTA CCG TCT AC GCC ACA TTG CTC CAA TAC	83

(continued on next page)

Table 1 (continued)

Gene symbol	Accession no.	Primer	Primer sequence	Product length (bp)
<i>APC</i>	NM_001075986.2	F R	GAG CCC TTC ACA GAA TGA CTC AGG ATA CAC GGG ATA AG	118
<i>FZD7</i>	NM_001144091.1	F R	GGG TGT GCC AAT CAA TTC CTG GGT AAC AGC CTT CTT	138
<i>CTNNB1</i>	NM_001076141.1	F R	CCC TTT GTC CAG CAA ATC CTG TGT TCC ACC CAT AGA	119
<i>MX1</i>	NM_1733940.2	F R	AGT CCA TCC GAC TAC ATT TC CIT CTT CTG CCT CCT TCT C	102
<i>COX-2</i>	NM_174445	F R	AGGTGATGTATGAGTGTAGGA GTGCTGGGCAAAGATGCAA	484
Sequences of primers used for qPCR analysis of CL tissue				
<i>BAX</i>	NM_173894.1	F R	TCT GAC GGC AAC TTC AAC TG CCA TGA TGG TCC TGA TCA ACT C	98
<i>CYP11A1</i>	NM_176644.2	F R	GAA TTA CCC AGG CAT CCT CTA C TCT CCG TAA TAT TGG CCT TGA C	97
<i>BCL-2</i>	NM_001166486.1	F R	ATC GTG GCC TTC TTT GAG TTC TCA GGT ACT CGG TCA TCC AC	104
<i>NOS2</i>	NM_001076799.1	F R	GAG CTT CTA CCT CAA GCT ATC G TCT ATC TCC TTT GIT ACT GCT TCC	94
<i>NOS3</i>	NM_181037.3	F R	GAT GGT CAA CTA CAT CCT GTC C GGT CTT CTT CCT GGT GAT GC	100
<i>FGF2</i>	NM_174056.3	F R	CAA CAG AAG ACC TAG GGA AGA C ACA GCC AAC TCC TAA CAT CC	124
<i>StAR</i>	NM_174189.2	F R	TAC ACC ATG TGG AAT GTC AGG CCT GTG TCA GTT GTA CAG TCT C	104
<i>3BHSD</i>	NM_174343.3	F R	GGT AAC GTG GCC TGG ATG CIT GTA GGG CGA GIT GTC ATA G	123
<i>FPr</i>	D17395	F R	TTAGAAGTCAGCAGCACAG ACTATCTGGGTGAGGGCTGATT	98
<i>OXT</i>	M25648.1	F R	GTCTGCACCATGGCAGGTT CAGGGGGCAGTTCGAATGT	125
Sequences of primers used for qPCR analysis of embryo tissue				
<i>PLAU</i>	NM_174147.2	F R	CTA GGG AGA AAG AAG AGT TCC TCG ATG CCT CCT GTA GAT	125
<i>HOXB7</i>	NM_174342.2	F R	ACC TAC ACC CGC TAT CA TGA TCT GTC TTT CTG TGA GG	118
<i>FTH1</i>	NM_174062.3	F R	AGG TGG AAG CCA TCA AAG GGG TGT GCT TGT CAA AGA	102
<i>EEF1A1</i>	NM_174535.1	F R	CTG GAA GAT GGC CCT AAA T GGG AGG ATA ATC AGA GAA GC	102
<i>GPX4</i>	NM_174770.3	F R	GCT GGC TAT AAC GTC AAA TTC GCT GGA CTT TCA TCC ATT TC	91
<i>ISG15</i>	NM_174366.1	F	GTA CAA GCA GAC CAG TTC	84
<i>IL-6</i>	NM_173923.2	F R	CTT CAA ACG AGT GGG TAA AG TAC TTC ATC CGA ATA GCT CTC	97
<i>BMP15</i>	NM_001031752.1	F R	CAT ACA GAC CCT GGA CTT TC GAG AGG TGG GAA TGA GTT AG	108
<i>IFN-tau</i>	AF238612	F R	GCCCTGGTGTGCTCAGCTA CIT CAT GAG GCC GTA TTC	102

Abbreviations: F, forward; qPCR, quantitative polymerase chain reaction; R, reverse.

(Applied Biosystems, Foster City, CA, USA) was used to synthesize complementary DNA (cDNA) from RNA. To proceed for reverse transcription polymerase chain reaction (RT-PCR) master mix, 5 μ L of DNase-treated RNA was mixed with a 5- μ L reaction mixture containing 1 μ L of 10X random primers, 0.4 μ L of 0.8-mM deoxyribonucleoside triphosphate mixture, 1 μ L of 10X buffer, 0.5 μ L of 50 U/ μ L of reverse transcriptase, 0.25 μ L of 40,000 U/mL of RNase inhibitor (New England Biolabs), and 1.85 μ L of nuclease-free water (provided in the kit). Then, the mixture was centrifuged at 2000 rpm for 2 minutes at 4 $^{\circ}$ C. The conditions used for RT-PCR was set as follows: 37 $^{\circ}$ C for 30 minutes, 75 $^{\circ}$ C for 15 minutes, and 4 $^{\circ}$ C for the final step. Finally, the products were stored at -20 $^{\circ}$ C until the quantification polymerase chain reaction (qPCR) was performed.

2.7. Quantitative real-time PCR

To perform transcription analysis and gene expression of reproductive tissues, 58 genes in total were selected on the basis of evidence in the literature showing their impact on endometrium remodeling, CL function, and embryo survival: 39 genes for endometrium, 10 genes for CL, and 9 genes for the conceptus (Table 1). These genes have been grouped on the basis of their roles during endometrium preparation, embryo and CL development (Table 2).

Transcript abundance was compared for a set of genes in the endometrium, CL, and embryonic tissue with three replicates per sample using quantitative real-time PCR (qPCR). The qPCR analysis was performed using the Rotor-Gene Q real-time cyler (Qiagen, Hilden, Germany),

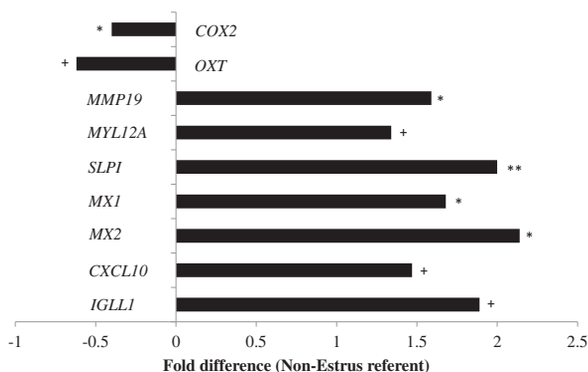


Fig. 2. Effect of estrous expression on endometrium gene expression. Significant fold difference based on nonestrous expression as a referent has been shown for genes with significant pattern of expression in endometrium tissue. For this graph, the asterisks (*, **) and (+) refer to $P \leq 0.05$, $P \leq 0.01$ and $P \leq 0.10$, respectively.

3.2. Corpus luteum gene expression

Among the analyzed genes from the CL tissue, *FPr* ($P = 0.05$), *CYP11A* ($P = 0.01$), and *BAX* ($P = 0.05$) were significant downregulated in estrus cows, with a 0.7-, 0.7-, and 0.8-fold difference in mRNA expression, respectively. The remaining genes analyzed (*NOS2*, *NOS3*, *FGF2*, *OXT*, *3βHSD*, *StAR*, and *BCL2*) were statistically unaltered by estrous expression ($P > 0.20$). All values of fold increase and significance are depicted in Figure 3.

3.3. Gene expression in the embryo associated with estrus

Downregulation in two different groups was observed in the embryos collected from cows in the estrus group compared with the nonestrous group. The *ISG15* gene, from the maternal recognition of the pregnancy group, was observed a 0.56-fold decrease ($P = 0.05$) in embryos collected from estrus cows. The *EEF1A1* ($P = 0.09$) and *PLAU* ($P = 0.01$), both transcripts that belong to the morphogenesis group, were also different between groups with a

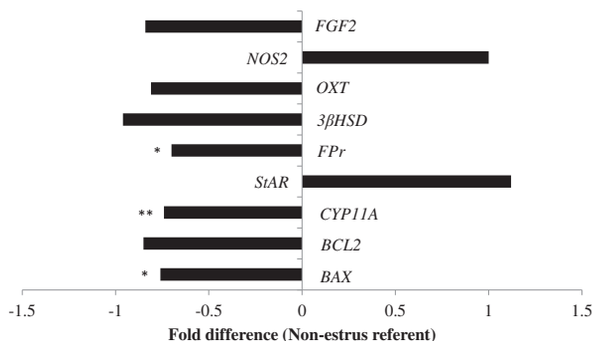


Fig. 3. Effect of estrous expression on CL genes involved in steroidogenesis, angiogenesis, and apoptosis. Significant fold difference based on nonestrous expression as a referent has been shown for genes with significant pattern of expression in CL tissue. For this graph, the asterisks (*) and (**) refer to $P \leq 0.05$ and $P \leq 0.01$, respectively.

0.81- and 0.74-fold difference, respectively, when comparing embryos from estrus versus nonestrous cows. A fold difference of 0.19 was observed for *BMP15*, which was downregulated in estrus cows compared with nonestrous cows ($P = 0.10$; Fig. 4). The remaining genes (*HOXB7*, *FTH1*, *IL6*, *IFNT*) were not significantly different (Fig. 4).

3.4. Ovarian and embryo parameters

Estrous expression positively affected the dimensional development of the embryos ($P = 0.02$) as they were around 10 cm longer when collected from cows in the estrus group (Table 3). The IFNT concentration within the uterine flushing media was not different between the estrus and nonestrous groups ($P = 0.47$). Follicle size was not affected by estrous expression as well (Table 3; $P = 0.89$). The CL tended to be smaller ($P = 0.10$) although concentrations of P4 were not statistically significant when comparing estrus and nonestrous cows on Day 7 ($P = 0.34$; Table 3). By Day 18, the volume of the CL was not different between groups ($P = 0.45$). There was a tendency for a greater BCS in nonestrous cows compared with estrus cows ($P = 0.10$; Table 3).

3.5. Effect of concentration of P4 on Day 7

Effect of concentration of P4 (high and low; based on median value) on Day 7 as a main factor affecting gene expression was analyzed. Gene expression in the endometrium was affected by P4 concentration. Immune-related genes within the endometrium such as *TRD*, *IGLL1*, *MX2*, and *SLPI* showed a significant upregulation when comparing the high versus low P4 concentrations ($P < 0.05$; Fig. 5). Other groups of genes which showed upregulation in the high-P4 group compared with the low-P4 group belong to adhesion molecules (*GLYCAM1* [$P = 0.003$] and *MYL12A* [$P = 0.02$]), the wnt signaling pathway (*APC* [$P = 0.001$] and *WNT2* [$P = 0.01$]). The *IL10* ($P = 0.09$), *CXCL10* ($P = 0.07$), *MX1* ($P = 0.07$), and *CDH1* ($P = 0.09$) also showed a tendency for upregulation in the high-P4 group compared with the low-P4 group. Embryo gene expression was also not affected by concentration of P4 on Day 7. The interaction between estrus effect and P4

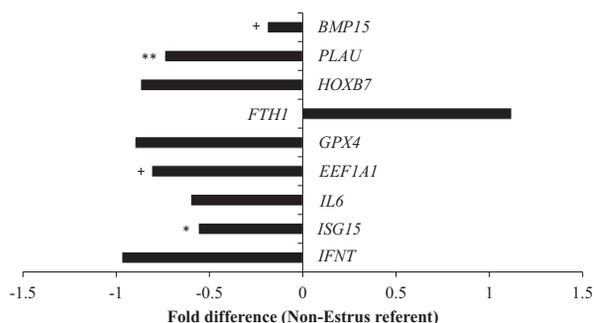


Fig. 4. Effect of estrous expression on embryo genes involved in morphogenesis, immune system, and protein synthesis. Significant fold difference based on nonestrous expression as a referent has been shown for genes with significant pattern of expression in endometrium tissue. For this graph, the asterisk (*, **) and (+) refer to $P \leq 0.05$, $P \leq 0.01$ and $P \leq 0.10$, respectively.

Table 3

Reproductive parameters collected on Days 7 and 19 of pregnancy from cows in the estrus and nonestrus groups.

Parameters	Estrus cows	Nonestrus cows	P value
BCS (1–5 scale)	3.30 ± 0.10	3.45 ± 0.10	0.10
Follicle diameter (mm)	14.0 ± 1.0	14.2 ± 1.0	0.89
P4 on Day 7 (ng/mL)	3.8 ± 0.9	5.2 ± 1.0	0.34
P4 on Day 18 (ng/mL)	3.9 ± 0.7	4.4 ± 0.8	0.62
CL diameter on Day 7 (cm)	6.9 ± 0.8	8.8 ± 0.8	0.10
CL diameter on Day 18 (cm)	10.5 ± 1.0	9.4 ± 1.0	0.45
Embryo length (cm)	38.3 ± 2.8	28.2 ± 2.9	0.02
IFNT concentration (pg/mL)	8.3 ± 1.7	10.2 ± 1.9	0.47

Abbreviations: BCS, body condition score; IFNT, interferon-tau; P4, progesterone.

concentration on Day 7 and their synergistic effect on endometrium gene expression were significant for immune-related genes such as *MX1* ($P = 0.003$), *MX2* ($P = 0.04$), *TRD* ($P = 0.05$), and *SLPI* ($P = 0.003$). *GLYCAM1* ($P = 0.04$), *APC* ($P = 0.01$), and *IgLL1* ($P = 0.08$; Fig. 6) also showed a differential gene expression on the basis of the interaction between expression of estrus and concentration of P4 on Day 7.

Other nongenomic results showed that IFNT concentration, CL volume on Days 7 and 18, follicle diameter, and BCS were not affected by categorization based on concentration of P4 on Day 7 ($P > 0.15$).

3.6. Effect of conceptus size

Animal variables and embryo gene expression were analyzed against embryo size (large and small; based on medium length [34 cm]). Embryo size did not affect IFNT concentration, CL volume on Days 7 and 18, concentration of P4 on Days 7 and 18, follicle diameter, and BCS. There were only two conceptus transcripts downregulated in the large-conceptus group (*BMP15* [fold difference = 0.05; $P = 0.005$] and *GPX4* [fold difference = 0.81; $P = 0.05$]).

4. Discussion

The aim of this study was to investigate the association of estrous expression at the time of AI with expression of

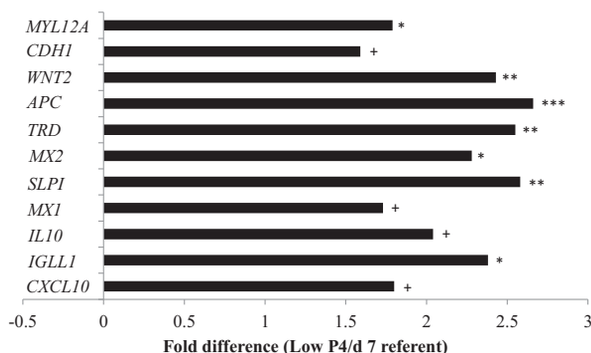


Fig. 5. Effect of progesterone (P4) concentration at Day 7 on endometrium gene expression. Significant fold difference based on the low-P4 group as a referent has been shown for genes with significant pattern of expression in endometrium tissue. For this graph, the asterisks (*, **, ***) and (+) refer to $P \leq 0.05$, $P \leq 0.01$, $P \leq 0.001$, and $P \leq 0.10$, respectively.

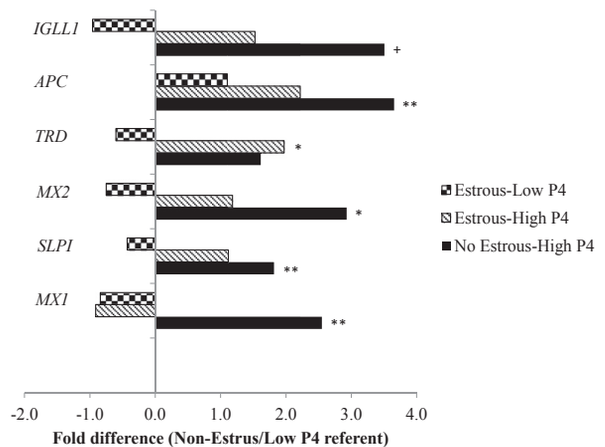


Fig. 6. Interaction between estrous expression and concentration of progesterone (P4) on Day 7 on endometrium gene expression. For this graph, the asterisks (*, **) and (+) refer to $P \leq 0.05$, $P \leq 0.01$, and $P \leq 0.10$, respectively.

critical genes in the endometrium, CL, and embryo during the preimplantation period. In addition, the difference in estrous expression was evaluated for reproductive parameters such as CL volume, conceptus size, concentration of P4 in plasma, and follicle diameter. Evidence from this study supports our hypothesis that estrous expression positively influences the expression of target genes important for embryo survivability. Cows that expressed estrous behavior near AI had a significant improvement in the profile of endometrium gene expression critical for suppressing the local maternal immune system and adhesion between endometrium epithelial cells and the conceptus, as well as partly inhibiting the mRNA machinery for *PGF2 α* synthesis. Genes related to the immune system and adhesion group in the endometrium were also significantly affected by P4 concentration on Day 7. The results from the gene analysis of the CL also confirmed downregulation of cellular pathways associated with apoptosis and *PGF2 α* synthesis which favors CL maintenance and secretion of P4, both key to sustain pregnancy.

The early embryonic development until implantation is arguably the most important period that define a successful pregnancy. A significant proportion of all embryonic losses in lactating cows occurs between Days 8 and 21 of pregnancy [35]. Because of operational limitations, it was not possible to check for length of dominance or P4 levels during the growth of the preovulatory follicle. The specific causes that lead to the presence or absence of estrous expression are unknown on the basis of the data collected in this study and warrant further investigations. The expression of estrus can indicate the state of sensitivity of the hypothalamus to E2 and perhaps the best timing for the optimal function of all other reproductive tissues related with the survivability of the early embryo.

The upregulation of immune system-related genes involved in endometrium receptivity (*MX1*, *MX2*, *IGLL1*, *SLPI*, and *CXCL10*) is in agreement with previous studies [36–39]. The *CXCL10* acts to attract trophoblasts to the endometrium and promote adhesive activity in ruminant

species [40,41] and has been shown to have more than a 11-fold upregulation in pregnant cows [39]. In a study by Walker et al. [42], *CXCL10* was downregulated in subfertile dairy cows compared with fertile cows. Myxoviruses are integral components of the innate immune system and were identified in blood leukocytes as a potential marker for pregnancy diagnosis in dairy heifers [43]. Hicks et al. [44] indicated a 15-fold increase in *MX1* and *MX2* from Days 12 to 15 after AI caused by pregnancy. Others have shown a temporal difference in the expression of these genes as indicated by greater expression of *MX2* on Days 18 and 20 compared with Days 14 and 16 of pregnancy [45]. The *IGLL1* expression positively impacts B cells development which are critical members of the adaptive immunity [46] and can indirectly enhance *MX1* and *MX2* activity. *SLPI* has the ability to interrupt the activation of transcription factor NF- κ B and possibly cause a reduction in *COX-2* expression, favoring CL maintenance. Some studies showed that hypoxia-induced *COX-2* expression also happens through the NF- κ B pathway [47,48].

The extensive molecular and structural changes taking place during the preimplantation stage in the endometrium are necessary for the reorganization of the glandular endometrium [49]. *MMP19* has been shown to be important for the regulation of conceptus attachment in bovine endometrium [50], whereas *MYL12A* expression is important for the regulation of protrusion and adhesion-generated signaling [51,52] as well as for cadherin clustering [53,54] and the stability of the cell–cell junction.

Data from the present study showed a decrease in the expression of *OTR* in the endometrium in the estrus group. It was reported that the expression of *OTR* is impacted by P4 and E2 concentrations [55,56] and key for the synthesis of PGF 2α and consequent maintenance of the CL [50,55,57,58]. The downregulation of *COX2*, a major enzyme necessary for the synthesis of PGF 2α , is probably a product of the lower expression of *OTR*. The optimal reduction in the expression of *OTR* and *COX2* signals on Day 19 of the estrous cycle may only appear when the complete estrous cycle, including proper expression of estrus, is allowed.

Results regarding the role of the wnt signaling pathway showed no significant difference in gene expression between animals that did or did not express estrus at the time of AI. The influence of the wnt signaling pathway could be dependent on the stage of embryo development. The activation of wnt signaling in bovine embryos by inhibitors of GSK3 β either blocks or increases development to the blastocyst stage [29]. It is known that at the morula stage, the embryo undergoes major genome activation [59] and perhaps the wnt signaling may have been already deactivated on Day 19 of pregnancy.

Analysis of target genes in the CL showed a significant decrease in genes related to apoptosis, PGF 2α and P4 synthesis. Downregulation of *BAX* may be due to the anti-luteolytic effects of *IFNT* (increase) or *COX2* (decrease). Sugino et al. [60] reported high *BCL2* and low *BAX* expression in the CL during the midluteal phase and early pregnancy in humans, whereas low *BCL2* and high *BAX* expression were found in the regressing CL. The PGF 2α receptors (*FPr*) are required to interact with PGF 2α released from uterus at the time of luteal regression [61], but during

pregnancy, the number of PGF 2α receptors in CL is reduced to allow CL maintenance. The PGF 2α synthesis is indirectly regulated by endometrial *COX2*, and its expression is necessary before luteolysis [62–64], which is corroborated by the results of the present study.

The gene expression of the conceptus had a significant reduction in *ISG15* and *PLAU* expression in the estrus group. In addition, *eEF1A1* and *BMP15* showed a tendency for downregulation. *ISG15* synthesis is stimulated by IFNT secretion from the conceptus and early detected on Day 17 of pregnancy but with peak levels between Days 18 and 23 and back to baseline levels by Day 45 in cows [65]. No difference between estrus and nonestrus cows regarding IFNT concentration on Day 18 conceptus tissue was observed in the present study, in spite of the difference in conceptus length favoring the estrus group. The benefit of a larger conceptus is likely the physical occupation of the lumen and increased likelihood of promoting IFNT-driven changes in as much endometrium tissue as possible. Although in some studies, they have reported a correlation between IFNT secretion and embryo size [66], they have not observed a relationship between IFNT concentration or embryo size and *IFNT* mRNA expression. We also observed a reduction in *BMP15* expression of cows in the estrus group which possibly relates to the temporal genome activation of the embryo. In a study by Pennetier et al. [67], these authors found *BMP15* transcripts until the five- to eight-cell stage but only trace levels in the morulae stage. According to our results, cows with smaller embryo size had greater expression of *BMP15* and *GPX4* in estrus versus nonestrus cows. The target genes affected by estrous expression in the conceptus seem of significant importance, but their interpretation is rather unclear. Further studies are necessary to clarify their roles and relationship with the endometrium status.

Ultimately, the present study found a correlation between P4 concentration and endometrial gene expression, which was mainly pronounced in immune system-related genes (*IL-10*, *MX1*, *SLPI*, *MX2*, *TRD*, *CXCL10*, and *IGLL1*), adhesion molecules (*GLYCAM1*, *CDH1*, and *MYL12A*), and wnt signaling (*APC* and *WNT2*). Other variables such as conceptus gene expression or animal physiological factors were not affected by P4 concentration on Day 7 of gestation. There was an interaction between estrous expression and P4 concentration which significantly affected expression of genes in the endometrium, specifically when the combination of estrous expression and low P4 concentration was in place. The upregulation of critical groups of genes in the endometrium under these circumstances of estrous expression and low P4 could be of great importance, particularly in beef cows. It is likely that a combination of factors leading to the day of collection (e.g., expression of estrus, endocrine milieu during the preimplantation phase) leads to the optimal function of reproductive tissues and embryonic receptivity.

4.1. Conclusions

The expression of estrus promoted changes in the preimplantation endometrium, CL, and conceptus gene expression. Critical cellular pathways related to

suppression of the maternal immune system, attachment between the conceptus and the endometrium, and CL maintenance during pregnancy were favorably expressed in cows that expressed estrus near AI. Moreover, cows in the estrus group yielded longer conceptuses, which can be associated with better chances of survival. The effects of expression of estrus seem to interact with P4 concentration on Day 7 of the estrous cycle in a way that positively influences endometrium receptivity and embryo development.

Acknowledgments

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