Expression of Glucocorticoid Receptor α and Its Regulation in the Bovine Endometrium: Possible Role in Cyclic Prostaglandin F2α Production

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Abstract. Cortisol (Cr), the most important glucocorticoid (GC), is well known to suppress uterine prostaglandin F2α (PGF) production. However, the details of the regulatory mechanisms controlling the cyclic changes in endometrial PGF production remain unclear. Here we investigated the expression of the GC receptor (GC-Rα), the actions of cortisol throughout the estrous cycle and the regulatory mechanism of GC-Rα in the bovine endometrium. The levels of GC-Rα protein were greater at the mid-luteal stage (Days 8–12) than at the other stages. Cr more strongly suppressed PGF production at the mid-luteal stage than at the follicular stage. GC-Rα expression was increased by progesterone (P4) but decreased by estradiol-17β (E2) in cultured endometrial stromal cells. The overall results suggest that ovarian steroid hormones control the cyclic changes in endometrial PGF production by regulating GC-Rα expression in bovine endometrial stromal cells.

Key words: Cattle, Endometrium, GC-Rα, Prostaglandins, Steroids

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The bovine endometrium synthesizes and secretes prostaglandin F2α (PGF), which is the main luteolysin responsible for regression of the corpus luteum (CL). In nonpregnant cows, the uterus increases PGF production between days 17 and 19 postovulation [1]. The increase in uterine PGF production and its pulsatile release seem to regulate the estrous cycle by inducing luteal regression in ruminants [2–4]. We reported that the amount of PGF production by the bovine endometrium was low at the mid-luteal stage, increased at the late-luteal stage and reached the highest levels at estrus [5]. Thus, PGF production and secretion by the endometrium seem to be precisely regulated throughout the estrous cycle.

Glucocorticoids (GCs) derived from the adrenal cortex have been shown to regulate female reproductive functions [6, 7]. Recently, we suggested that an active GC, cortisol (Cr), affects pregnancy rate and CL function in cattle [8]. In addition, Cr suppresses basal and tumor necrosis factor-α-stimulated PGF production in bovine endometrial stromal cells but not in the epithelial cells [9]. Moreover, 11β-hydroxysteroid dehydrogenases (HSD11B) play an important role in regulating GC availability in target tissues. HSD11B type 1 mainly converts cortisone to Cr in the target organs [10, 11]. A previous study demonstrated that the activity of HSD11B in the bovine endometrium was greater at the follicular stage and estrus than at the other stages of the estrous cycle [9]. In addition, PGF increased the protein expression of HSD11B in bovine endometrial stromal cells [9]. The above findings suggest that Cr plays some important roles in regulating PGF production throughout the estrous cycle in the bovine endometrium.

The above findings raise the possibility that the intracellular availability of Cr increases and that PGF production is thereby suppressed when HSD11B expression and activity are high at the follicular stage. However, PGF production by the bovine endometrium is highest at the follicular stage [5, 9]. These results suggest that the inhibitory effect of Cr on PGF production depends on other factors. The biological action of Cr is mediated through the activation of specific intracellular receptors, GC receptors (GC-R) [12–16]. Because Cr actions depend on the expression of GC-Rα in the target organs, endometrial PGF production appears to be regulated by not only changes in the levels of active GC (Cr) but also changes in GC-Rα expression throughout the estrous cycle. However, the regulatory mechanisms controlling the cyclic changes in endometrial PGF production remain unclear.

Progesterone (P4) and estradiol-17β (E2) secreted by the ovary play crucial roles in regulating functional and structural changes in the endometrium throughout the estrous cycle [17]. Furthermore, in ruminants, the uterine epithelium, stroma and myometrium all contain receptors for P4 and E2 [18]. Thus, we hypothesize that P4 and E2 regulate GC-Rα expression, which plays an important role in controlling PGF production in the bovine endometrium.

In the present study, to determine the regulatory mechanisms of PGF production in the bovine endometrium, we investigated 1) GC-Rα protein expression and the biological actions of Cr in the bovine endometrium throughout the estrous cycle and 2) the roles of steroid hormones in the regulation of GC-Rα expression in bovine endometrial stromal cells.
Materials and Methods

Collection of endometrial tissues

Apparently healthy uteri of Holstein cows without a visible conceptus were obtained at a local slaughterhouse within 10–20 min of exsanguinations in accordance with protocols approved by the local institutional animal care and use committee. The stages of the estrous cycle were confirmed by macroscopic observation of the ovary and uterus as described previously [5, 19]. For protein determination, endometrial tissues (n=3/stage) were collected from cows at six different stages of the estrous cycle (estrus, day 0; early-luteal, days 2–3; developing, days 5–6; mid-luteal, days 8–12; late-luteal, days 15–17 and follicular stage, days 19–21). Intercaruncular endometrial tissues from the uterine horn, ipsilateral to the CL, were used for experiments. The endometrial tissues were immediately frozen in liquid nitrogen, and stored at −80°C until processed for protein isolation. For experiments involving tissue culture and cell culture, the uteri were submerged in ice-cold physiological saline and transported to the laboratory within 1–1.5 h on ice.

Culture of endometrial tissues

For tissue culture, endometrial tissues were obtained at the mid-luteal stage and at the follicular stage. Endometrial tissue culture was conducted as described previously [9]. Briefly, endometrial strips were washed three times in sterile saline solution containing penicillin (100 IU/ml) and streptomycin (100 μg/ml). The tissues were finally cut into small pieces (40 to 50 mg) with a scalpel and subsequently washed another three times in Hank’s balanced salt solution supplemented with penicillin (100 IU/ml), streptomycin (100 μg/ml) and 0.1% BSA. After hanging the tissues with steel needles (TOP, Tokyo, Japan; 8N01B), the individual endometrial tissues were placed into culture glass tubes (12 mm × 75 mm; Kimble Chase Life Science and Research Products, Vineland, NJ, USA; 73500-13100) containing 2 ml culture medium (DMEM/Ham’s F-12; 1:1 (v/v) [Invitrogen, Carlsbad, CA, USA; 12400-024]) and supplemented with penicillin (100 IU/ml), streptomycin (100 μg/ml) and 0.1% BSA under 5% CO2 in air. Endometrial tissues were exposed to Cr (10–100 nM) at 37.5°C for 4 h. At the end of incubation, 1 ml of the conditioned media were collected into 1.5 ml tubes containing 10 μl of a stabilizer solution (0.3 M EDTA, 1% (W/V) acetylsalicylic acid, pH 7.3) and frozen at −30°C. The tissues were blotted on filter paper and weighed to normalize the PGF concentration. The concentrations of PGF in the culture media after incubation were determined by enzyme immunoassay (EIA). The PGF standard curve ranged from 0.016–4 ng/ml, and the ED 50 of the assay was 0.25 ng/ml. The DNA content was used to standardize the PGF concentration in the culture media.

Isolation of endometrial cells

Early-luteal stage uteri (days 2–5) were used for isolation of endometrial cells. The epithelial and stromal cells from the bovine endometrium were enzymatically separated (0.05% collagenase; Worthington Biochemical, Lakewood, NJ, USA; #CLS1) using procedures described previously [20]. A polycarbonate catheter was inserted into the side of the oviduct, and the ends of the horn were tied to retain trypsin solution used to detach the epithelial cells as described herein. The uterine lumen was washed three times with 30–50 ml of sterile Ca²⁺- and Mg²⁺-free Hank’s balanced salt solution (HBSS) supplemented with 100 IU/ml of penicillin (Meiji Seika Pharma, Tokyo, Japan; 611400D3051), 100 μg/ml of streptomycin (Meiji Seika Pharma; 6161400D1034) and 0.1% (w/v) bovine serum albumin (BSA) (Roche Diagnostics, Manheim, Germany; 10735086001). Thirty to fifty milliliters of sterile HBSS containing 0.3% (w/v) trypsin (Sigma-Aldrich, St. Louis, MO, USA; T9201) was then infused into the uterine lumen through the catheter. After collection of the epithelial cells, the uterine lumen was washed with sterile HBSS supplemented with antibiotics and 0.1% (w/v) BSA. The horn was then cut transversely with scissors into several segments, which were slit to expose the endometrial surface. Intercaruncular endometrial strips were dissected from the myometrial layer with a scalpel and washed once in 50 ml of sterile HBSS containing antibiotics. The endometrial strips were then minced into small pieces (1 mm³). The minced tissues (~5 g) were digested by stirring for 60 min in 50 ml of sterile HBSS containing 0.05% (w/v) collagenase, 0.005% (w/v) DNase I (BBI Enzymes, Cardiff, UK; DNP2), and 0.1% (w/v) BSA. The dissociated cells were filtered through metal meshes (100 μm and 80 μm) to remove undissociated tissue fragments. The filtrate was washed three times by centrifugation (10 min at 100 g) with Dulbecco’s Modified Eagle’s Medium (DMEM) (Sigma-Aldrich; D1152) supplemented with antibiotics and 0.1% (w/v) BSA. After the washes, the cells were counted using a hemocytometer. The cell viability was greater than 85% as assessed by 0.5% (w/v) trypan blue dye exclusion.

Culture of endometrial stromal cells

The final pellet of the isolated stromal cells was resuspended in culture medium (DMEM/Ham’s F-12; 1:1 (v/v); Invitrogen; 12400-024) supplemented with 10% (v/v) bovine serum (Invitrogen; 16170078), 20 μg/ml gentamicin (Sigma-Aldrich; G1397) and 2 μg/ml of amphotericin B (Sigma-Aldrich; A9528) [21]. The stromal cells were seeded at a density of 1×10⁵ viable cells/ml in 24-well cluster dishes (Greiner Bio-One, Frickenhausen, Germany; #662160) or 1×10⁶ viable cells/ml in 75 cm² culture flasks (Greiner Bio-One; #658175) for GC-Rα protein determination (Figs. 3–5), and were cultured at 37.5°C in a humidified atmosphere of 5% CO2 in air. For thestromal preparation, the medium was changed 2 h after plating, by which time selective attachment of stromal cells had occurred [21, 22]. The homogeneity of stromal cells was evaluated using immunofluorescent staining for specific markers of epithelial (cytokeratin) and stromal (vimentin) cells as described previously [23]. Epithelial cell contamination of stromal cells was about 1%. When cells were confluent (6–7 days after the start of the culture), the medium was replaced with fresh DMEM/Ham’s F-12 supplemented with 0.1% (w/v) BSA, 5 ng/ml sodium selenite (Sigma-Aldrich; #S5261), 0.5 mM ascorbic acid (Wako Pure Chemical Industries, Osaka, Japan; 031-12061), 5 mg/ml transferrin (Sigma-Aldrich; T4132), 2 mg/ml insulin (Sigma-Aldrich; I-4011) and 20 mg/ml gentamicin.

Stromal cells were exposed to Cr (1–100 nM), and P4 (0.1–10 nM) and E2 (0.1–10 nM) for 24 h (n=3). After incubation, total RNA was extracted for determination of GC-Rα mRNA expression. For protein analysis, the cultured cells were scraped and placed in ice-cold
Western blotting analysis

GC-Rα protein levels in endometrial tissues and stromal cells were assessed by Western blotting analysis. Endometrial tissues were homogenized on ice in the homogenization buffer by a tissue homogenizer (Physcion; Microtec Do., Chiba, Japan; NS-50), followed by filtration with a metal wire mesh (150 μm). For GC-Rα protein analysis, nuclei were isolated from the tissue homogenates by centrifugation at 600 x g for 30 min. The cultured stromal cells were lysed in 200 μl of lysis buffer (20 mM Tris-HCl, 150 mM NaCl, 1% Triton X-100, 10% glycerol [Sigma-Aldrich; G7757], Complete, pH 7.4). Protein concentration was determined by the method of Osnes et al. [24] using BSA as a standard. The proteins were then solubilized in SDS gel-loading buffer (10% glycerol, 1% β-mercaptoethanol [Wako Pure Chemical Industries; 137-06862], pH 6.8), and heated at 95°C for 10 min. Samples (30 μg protein) were subjected to SDS-PAGE (12%) for 1.5 h at 200 V. The separated proteins were electrophoretically transblotted to a nitrocellulose membrane (Amersham Biosciences, Piscataway, NJ, USA; RPN787D) for 3 h at 250 mA in transfer buffer (25 mM Tris-HCl, 192 mM glycine, 20% methanol, pH 8.3). The membrane was washed in TBS-T (0.1% Tween 20 in TBS [25 mM Tris-HCl, pH 7.5, 137 mM NaCl]), incubated in blocking buffer (4% nonfat dry milk in TBS-T) overnight at 4°C, incubated at room temperature with a primary antibody specific to each protein (GC-Rα antibody [95 kDa; Santa Cruz Biotechnology, Santa Cruz, CA, USA; sc-1002; 1:200, 1 h] and β-actin (ACTB) antibody [42 kDa; Sigma-Aldrich; A2228; 1:4,000, 1 h]), incubated in blocking buffer for 10 min at room temperature, washed two times for 10 min in TBS-T at room temperature, incubated with secondary antibody [GC-Rα (1:10,000), anti-rabbit Ig, HRP-linked whole antibody produced in donkey, Amersham Biosciences, NA934; ACTB (1:40,000), anti-mouse Ig, HRP-linked whole antibody produced in sheep, Amersham Biosciences, NA931] for 1 h, washed two times in TBS-T for 10 min and then washed in TBS for 10 min at room temperature. The signal was detected using an ECL Western Blotting Detection System (Amersham Biosciences; RPN2109).

The intensity of the immunological reaction (GC-Rα and ACTB)
in the tissues was estimated by measuring the optical density in the defined area by computerized densitometry using NIH Image (National Institutes of Health, USA).

**Reverse transcription and real-time PCR**

Total RNA was extracted from endometrial tissue using TRIzol (Bioline, London, UK; #BIO-38033) according to the manufacturer’s directions. Extracted DNA from each sample was quantified using a NanoDrop Lite Spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA). DNA in the solution was quantified by the absorbance of light (260 nm) in the spectrophotometer. One microgram of each total RNA was reverse transcribed using a ThermoScript RT-PCR System (Invitrogen), and 10% of the reaction mixture was used in each PCR using specific primers for GC-Rα and GAPDH from the bovine sequence. The primers were chosen using Primer3, an online software package (http://primer3.sourceforge.net/).

Gene expression was measured by real-time PCR using an Mx3000P QPCR System (Stratagene, La Jolla, CA) and a QuantiTect SYBR Green PCR system (Qiagen, Hilden, Germany) starting with 2 ng of reverse-transcribed total RNA as described previously [25]. Briefly, GAPDH expression was used as an internal control. The sequences of the GC-Rα primers, 5’-CCATTTCGTTACCGGTGTG-3’ (5’ primer, 20 mer) and 5’-CTGAACCGACAGGGAATTG-3’ (3’ primer, 20 mer), were synthesized according to bovine GC-Rα cDNA (GenBank accession number AY238475), and these primers generated a specific 132-base pair (bp) product from all cell types. The primers for GAPDH were 5’-CACCTCAAGATTGTCAGCA-3’ (5’ primer, 20 mer) and 5’-GGTCATAAGTCCCTCCACGA-3’ (3’ primer, 20 mer). These primers generated a specific 103-bp product from all cell types. For quantification of the mRNA expression levels, the primer length (20 bp) and guanine and cytosine contents of each primer (50–60%) were selected, and PCR was performed under the following conditions: 95°C for 15 min, followed by 55 cycles of 94°C for 15 sec, 60°C for 20 sec, and 72°C for 15 sec. Use of the QuantiTect SYBR Green PCR system at elevated temperatures resulted in reliable and sensitive quantification of the RT-PCR products with high linearity (Pearson product-moment correlation coefficient, r=0.99).

**Statistical analysis**

All experimental data are shown as the mean ± SEM of values obtained in 3–4 separate experiments, in which each experiment was performed in triplicate using stromal cells from a single bovine endometrium. The statistical significance of differences in the effects of Cr, P4 and E2 on GC-Rα mRNA and protein expressions and the effects of Cr on PGF production was assessed by analysis of variance (ANOVA) followed by a Fisher’s protected least-significant difference procedure (PLSD) as a multiple comparison test. Statistical significance of differences in GC-Rα protein expression was assessed by Student’s t-test.

**Results**

**GC-Rα protein expression throughout the estrous cycle**

GC-Rα protein was detected in the bovine endometrium throughout the estrous cycle (n=3 cows/stage). GC-Rα protein expression was higher at the mid- and late-luteal stages than at other stages (Fig. 1, P<0.05).

**Effect of Cr on endometrial PGF production at the mid-luteal stage and at the follicular stage**

Cr (10 nM) decreased PGF production to 44% of the control at the mid-luteal stage (P<0.05), whereas Cr tended to decrease PGF production to 68% of the control at the follicular stage (P>0.05) (Fig. 2).
Effect of Cr, P4 and E2 on GC-Rα mRNA and protein expression in cultured bovine endometrial stromal cells

The suppressive effect of Cr on GC-Ru protein was significant at 100 nM. However, the suppressive effect of Cr on GC-Rα mRNA expression was apparent at 1 nM in endometrial stromal cells (Fig. 3). P4 increased GC-Ru mRNA and protein expression. The effect was significant at 10 nM in cultured bovine endometrial stromal cells (Fig. 4, P<0.05). Exposing cultured bovine endometrial stromal cells to E2 did not affect GC-Rα mRNA expression (Fig. 5A) but significantly decreased GC-Rα protein expression (Fig. 5B, P<0.05).

Discussion

The biological action of Cr is mediated through intracellular GC-R. Two isoforms of GC-R (GC-Ru and GC-Rβ) originate from the same gene by alternative splicing of the GC-R primary transcript [26–28]. Since the ligand-dependent GC-Ru stimulates gene transcription in Cr target tissues, GC-Ru is thought to be the active receptor isoform [26]. GC-Ru protein was detected in the nuclei of most cell types in the endometrium of cyclic and pregnant cows [29]. In the present study, we examined whether changes in GC-Ru expression during the estrous cycle are associated with PGF production in the bovine...
endometrium. The protein expression of GC-Rα was significantly higher at the mid-luteal stage than at the other stages. Since PGF production is significantly lower at the mid-luteal stage than at the follicular stage [5], the cyclic changes in GC-Rα seem to be associated with the regulation of PGF production in the bovine endometrium. In fact, the suppressive effect of Cr on PGF production by the bovine endometrial tissue was greater at the mid-luteal stage than at the follicular stage in the present study. These findings suggest that changes in GC-Rα expression are relevant for the suppressive effect of Cr on PGF production in the bovine endometrium.

Glucocorticoids and other steroid hormones downregulate the levels of their cognate receptors in a number of target tissues and in many different cell lines [30, 31]. GC-Rα was downregulated by its own ligand in different cell types [32, 33]. This effect is thought to be a feedback promoter mechanism that would avoid deleterious effects of prolonged exposure to a hormone [34, 35]. In the present study, Cr also inhibited GC-Rα mRNA and protein expression in bovine endometrial stromal cells. The above findings suggest that Cr has a role in regulating GC-Rα expression in bovine endometrial stromal cells. Although the plasma concentration of Cr does not change throughout the estrous cycle [36], the local concentration of Cr in the bovine endometrium has been suggested to be regulated by the levels of HSD11B [9]. HSD11B protein expression and activity were greater at the follicular stage than at the mid-luteal stage [9]. Thus, the reason why GC-Rα expression was low at estrus and at the follicular stage in the present study may be downregulation by a high level of local Cr.

Both P4 and E2 are sex steroid hormones that regulate several female reproductive functions [37]. The GC-Rα mRNA [9] and protein expressions observed in our study were highest at the mid-luteal stage in the bovine endometrium. The changes in GC-Rα expression throughout the estrous cycle were similar to the cyclic changes in the plasma P4 concentration [2], suggesting that P4 is one of the regulators of GC-Rα expression throughout the estrous cycle. In fact, P4 stimulated GC-Rα expression in cultured stromal cells in the present study. In contrast to P4, the plasma E2 concentration increases during the follicular stage and reaches a peak at estrus, when endometrial PGF production is higher than at the other stages of the estrous cycle [21]. There is a great difference in the concentrations of P4 and E2 in bovine peripheral blood. The concentrations of P4 and E2 (0.1–10 nM) used in the present study were within the range of the concentrations used in previous studies [38, 39]. E2 has been shown to downregulate GC-Rα expression and influence the sensitivity to Cr in human breast cancer cells [40]. In agreement with these reports, E2 inhibited GC-Rα protein, but not GC-Rα mRNA expression in cultured bovine endometrial stromal cells in the present study, suggesting that E2 may influence the total amount of GC-Rα protein by increasing protein catabolism or by reducing translational efficiency in bovine endometrial stromal cells. Thus, P4 and E2 may regulate PGF production by promoting or suppressing the expression of GC-Rα, at least in stromal cells.

In conclusion, the findings of this study suggest that the expression of GC-Rα is important in regulation of PGF production in the bovine endometrium and that sex steroid hormones and Cr control the cyclic changes in endometrial PGF production in the bovine endometrium at least in part by regulating GC-Rα expression.

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