The role of adenosine monophosphate activated protein kinase in luteal progesterone production

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The role of adenosine monophosphate activated protein kinase in luteal progesterone production

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Dissertation submitted
to Davis College of Agriculture, Natural Resources and Design
at West Virginia University

in partial fulfillment of the requirements for the degree of

Doctor of Philosophy
in
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Abstract

The role of adenosine monophosphate activated kinase in luteal progesterone production

Elizabeth Compton Bowdridge

Prostaglandin F<sub>2</sub> alpha (PGF<sub>2α</sub>) has in vivo luteolytic actions on the bovine corpus luteum (CL). Although PGF<sub>2α</sub> is utilized extensively in livestock to synchronize estrus, its actions in vitro are controversial and do not always result in a reduction in progesterone (P4) secretion. The mechanism of action of PGF<sub>2α</sub> is thought to involve activation of the phospholipase C-<sup>Ca<sup>2+</sup></sup> pathway. The luteolytic actions of PGF<sub>2α</sub> are mediated through an elevation of cytosolic Ca<sup>2+</sup> involving activation of protein kinase C (PKC). Expression of PKC isozymes and calcium/calmodulin-dependent protein kinase kinase 2 (CAMKK2) change as the CL develops to a mature stage. Either CAMKK2 directly affects steroidogenesis or works indirectly through an intermediary step, such as adenosine-monophosphate kinase (AMPK), which is activated through CAMKK2 targeted phosphorylation. Specifically, the hypothesis that activation of PGF<sub>2α</sub> receptor (FP) in the mature CL involves AMPK was examined. Expression of mRNA encoding a potential target AMPK for the action of CAMKK2, was increased in the mature versus developing CL. Furthermore, activation of FP induced rapid phosphorylation of AMPK. An AMPK-specific inhibitor, dorsomorphin dichloride (DM), eliminated effects of PGF<sub>2α</sub> on secretion of P4, supporting the hypothesis that activation of the FP receptor in mature CL involves participation of AMPK. Furthermore, in vitro two AMPK activators, 5-aminooimidazole-4-carboxamide ribonucleotide (AICAR) and metformin, decreased basal P4 secretion in mature but not in developing CL. Effect of FP activation on cholesterol transport in the mature CL was investigated. Both serum and luteal P4 declined as soon as 2 hours after PGF<sub>2α</sub> administration, which was paralleled by a decrease in protein concentration of low density lipoprotein receptor (LDLR). Human mural granulosal cells are collected routinely when a patient is undergoing in vitro fertilization (IVF) and are useful for studying granulosal lutein cells of the CL. Altering intracellular calcium in these cells through an ionophore, A23187, increased basal P4 production. However, at high concentrations a calcium chelator, BAPTA, decreased P4 from mural granulosal cells. Infertility is a problem that affects more than 6.1 million women and their partners across the United States. Causes of infertility range from female related tubal factors such as blockage, endometriosis, diminished ovarian reserve, to male related infertility due to abnormal sperm motility or morphology to unexplained infertility. Primary infertility types, age, BMI, and pregnancy outcome influenced ability of granulosal cells to increase P4 production when stimulated with human chorionic gonadotropin (hCG). AICAR decreased basal P4 production in these cells, which is similar to earlier observations in the mature bovine CL. Thus, AMPK appears to be a distal target in the pathway responsible for mediating the actions of PGF<sub>2α</sub>. Directly activating AMPK could lead to better estrous cycle manipulation in the cow, or overcoming luteal insufficiency in women undergoing IVF.
Dedication

“Here’s to strong women,
May we know them,
May we be them,
May we raise them.”
-Unknown

To my daughter, Charlotte, and to all the great women who have made me who I am.
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# Table of Contents

The role of adenosine monophosphate activated kinase in luteal progesterone production................................................................................................................................. i

Abstract............................................................................................................................................. ii

Dedication ........................................................................................................................................ iii

Acknowledgements ............................................................................................................................ iv

Table of Figures................................................................................................................................... viii

Chapter 1: Literature Review ............................................................................................................ 1

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

Development of the bovine corpus luteum ........................................................................................ 2

Neovascularization.............................................................................................................................. 2

Small and large luteal cells .................................................................................................................. 4

Progesterone production ................................................................................................................... 5

Lutienization of human mural granulosal cells............................................................................... 6

Luteotropic versus luteolytic signals ................................................................................................. 7

Luteotropic hormones ......................................................................................................................... 7

Luteolytic hormones ............................................................................................................................ 8

Signals for maintenance of the corpora lutea ................................................................................... 10

Estrous synchronization protocols ................................................................................................... 11

Responsiveness of corpora lutea to PGF\textsubscript{2α} ........................................................................ 11

Normal versus induced corpora lutea ............................................................................................... 12

Structural and functional regression of corpus luteum .................................................................. 12

Structural versus functional .............................................................................................................. 12

Factors involved in regression .......................................................................................................... 15

PGF\textsubscript{2α} signal transduction pathway .................................................................................... 21

Adenosine monophosphate activated kinase (AMPK) ................................................................. 21
### Pharmaceutical agents

Model of activation of AMPK in the corpus luteum

Statement of the problem:

| Chapter 2: Effects of prostaglandin F 2 alpha and adenosine monophosphate activated kinase on progesterone production in the bovine corpus luteum in vivo |
| Introduction |
| Materials and Methods |
| Results |
| Discussion |

| Chapter 3: Adenosine monophosphate activated protein kinase activation in vitro modulated progesterone secretion in mature but not developing bovine CL |
| Introduction |
| Materials and Methods |
| Results |
| Discussion |

| Chapter 4: Primary infertility affects progesterone production by luteinized human mural granulosal cells in vitro |
| Introduction |
| Materials and Methods |
| Results |
| Discussion |

| Chapter 5: Literature Cited |

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vii
Table of Figures

Figure 1.0: Model of actions of AMPK in luteal cells ................................................................. 27
Figure 1.1: Bivariate fit of serum (A) and luteal (B) P4 (ng/ml) by time from PGF₂α (h). ........ 39
Figure 1.2: Semiquantitative analysis of the densitometric data derived from Western blots using protein samples isolated from mature bovine CLs 0, 2, and 4 h after PGF₂α. .............. 40
Figure 1.3: Concentrations of serum progesterone after 15 or 30 (mg/kg of BW) IU of metformin ........................................................................................................................................ 41
Figure 1.4: Concentrations of serum progesterone pooled across experiments .......................... 42
Figure 1.5: Serum insulin concentrations one day prior to and during metformin treatment ..... 43
Figure 2.1: Stimulatory effect of PGF₂α as blocked by DM in mature (d10) but not developing CL (d4) .......................................................................................................................................... 50
Figure 2.2: Neither PGF₂α nor its activators, metformin and AICAR, had any significant effect on basal progesterone production in developing (D4) bovine CL .................................................................................. 51
Figure 2.3: At the greatest concentrations tested, both metformin and AICAR significantly decreased basal progesterone production in mature (D10) bovine CL. ...................................................... 52
Figure 3.1: Production of progesterone by granulosa cells from patients with male factor (A), endometriosis (B), diminished ovarian reserve (C), and tubal (D) primary infertility .... 65
Figure 3.2: Production of progesterone in mural granulosa cells affected by age. ...................... 66
Figure 3.3: Production of progesterone in mural granulosa cells affected by BMI. .................... 67
Figure 3.4: Production of progesterone in mural granulosa cells affected by pregnancy status post IVF. .................................................................................................................................................. 68
Figure 3.5: The AMPK activator, AICAR, inhibited P4 production by human luteinized mural granulosa cells ........................................................................................................................................ 69
Figure 3.6: Production of P4 as stimulated by PGF₂α in mural granulosa cells but not cumulus granulosa cells ........................................................................................................................................ 70
Figure 3.7: Biphasic effect of BAPTA on production of P4 in luteinized mural granulosa cells 71
Figure 3.8: Increases in intracellular Ca²⁺ increased production of P4 in mural granulosa cells. .................................................................................................................................................. 72
Chapter 1: Literature Review

Introduction

Follicles, composed of the oocyte, granulosal and thecal cells, are the functional unit of the ovary. The somatic cells in the outermost vascular portion of the follicle include thecal and myoepithelial cells. Myoepithelial cells aid in rupture of the follicle during ovulation by increasing intrafollicular pressure through contraction. At least two populations of thecal cells are recognized: theca interna and theca externa. The theca interna expresses luteinizing hormone (LH) receptor and produces androstenedione, but does not contain aromatase, the enzyme needed for conversion of androstenedione to estrogen, which is expressed in granulosal cells (Hseuh et al., 1984). The avascular component of the follicle is delimited by a basement membrane upon which the other somatic cells of the follicle, the granulosal epithelial layer and the germ cell reside. Two populations of granulosal cells have been identified: mural and cumulus granulosal cells. Both populations of cells are organized as a stratified epithelium with the basal layer of mural cells contacting the basement membrane adjacent to the thecal cells and the cumulus cells surrounding the oocyte (Buccione et al., 1990). The 2-cell, 2-gonadotropin theory was first suggested 50 years ago (Short, 1961). The theory is essential in understanding the role that these two types of granulosal cells play in estrogen secretion from the growing follicle (Liu and Hsueh, 1986). In most mammalian species, formation of a follicular antrum, a fluid filled form of extracellular matrix (ECM; Rogers and Rogers, 2010), divides these two populations of granulosal cells. Presumably, the antrum is formed by an osmotic gradient that draws fluid from the vasculature that is adjacent to the thecal cell population (Rogers and Rogers, 2010). Granulosal cells produce both hyaluronan and versican, and these two molecules contribute to cumulus cell expansion (Rogers and Rogers, 2010). After antrum formation, mural granulosal
cells express both FSH and LH receptor as well as 3β-HSD and P450 SCC (Zoller and Weisz, 1979) the expression of these two enzymes is more a luteal feature in these cells. Cumulus granulosal cells remain with the oocyte after ovulation and respond to the ovulatory LH surge by assembling another specialized ECM by a process known as cumulus expansion (Russell and Salustri, 2006). This unique set of somatic cells possess an epithelial phenotype and are tightly connected to one another via intercellular processes, known as gap junctions, which allow for passage of small molecules that regulate signaling and metabolism by the oocyte (Russell and Salustri, 2006). This subset of granulosal cells has a gene expression profile that is specific for cumulus cell expansion, which distinguishes them from mural granulosal cells, which do not undergo expansion. Once ovulation has occurred, mural granulosal cells lose their ability to proliferate, undergo epithelial-mesenchyme transition and differentiate into the large luteal cells (LLC) of the corpus luteum (CL), which secrete high concentrations of basal progesterone (Chaffkin et al., 1992). Theca cells become small luteal cells (SLC) once luteinization has occurred. In the primate, the granulosal and thecal cells remain as two distinct populations and are separated by the remnants of the follicular basement membrane (Guraya, 1971). In contrast, the basement membrane separating these two cell populations is degraded, and thecal cells invade the granulosa to form a heterogeneous mixture of cells in the CL of nonprimate species (Niswender, 2000).

**Development of the bovine corpus luteum**

**Neovascularization**

The CL is a transient endocrine organ that begins to form immediately after ovulation and requires extensive neovascularization to ensure both structural and functional development. During the first eight to ten days of growth, the CL doubles in size and cell number every 60-70
hours (Reynolds et al., 1994). This rate of growth is comparable to the fastest growing tumors, yet, unlike tumors, cell proliferation in the CL is self-limiting and tightly regulated (Reynolds et al., 1994). Ovulated follicular tissues of approximately 40 mg grow to 600-700 mg of CL tissue within a few days (Reynolds, 2000). This growth can be attributed to hyperplasia of SLC, fibroblasts, and endothelial cells, as well as hypertrophy LLCs without hyperplasia in the ovine CL (Farin et al., 1986). However, *in vitro* both hypertrophy and hyperplasia occur in bovine luteal steroidogenic cells as measured by cell cycle (cyclin D2 and E1) and cell proliferation (Ki-67) genes (Yoshioka et al., 2013). To support such extensive cell growth, angiogenesis occurs rapidly and reaches a maximum 2 to 3 days after ovulation (Reynolds, 2000). This change is reflected by the fact that approximately 50-85% of the luteal cell proliferation occurs in the microvasculature unit, and endothelial cells and pericytes make up over half of the cell population in the mature CL (Reynolds et al., 1994). Increased vascularization enables the CL to receive more blood flow per unit of tissue than any other mature organ in the body (Reynolds, 2000). Most membranes of luteal cells are either adjacent to capillaries or adjacent to the interstitial space that is in close proximity to capillaries (Dharmarajan et al., 1985). Inadequate or subfunctional luteal development is known to occur in cattle (Redmer & Reynolds, 1996) and in humans (Reynolds et al., 2000) and is associated with reduced vascularization in both cases.

Several angiogenic factors contribute to the vascularization of the bovine CL. Two key mediators in neovascularization are fibroblast growth factor 2 (FGF2) and vascular endothelial growth factor (VEGF) A (Robinson et al., 2009). Both VEGF and its receptors VEGFR1 and VEGFR2 mRNA are expressed in the bovine CL and endothelial cells (Berisha et al., 2000). Additionally, VEGF is a mitogen for endothelial cells, and its expression is induced by the preovulatory LH surge (Garrido et. al, 1993). In the cow, FGF2 protein synthesis occurs in
endothelial cells and pericytes of the thecal layer occur prior to the LH surge and in the luteinizing granulosa mainly after ovulation (Berisha et al., 2006). After a few days FGF2 was absent from all steroidogenic cells but reappeared in endothelial cells (Berisha et al., 2006). Both endothelial cells (Robinson et al., 2008) and pericytes (Robinson et al., 2007) are targets for multiple angiogenic factors. Pericytes appear to play a dual role in neovascularization. First they act as a guide for the outgrowth of endothelial cells, and secondly they help stabilize newly formed vessels (Robinson et al., 2009). Changes in mRNA and protein expression in FGF2 that occur after the LH surge imply that FGF2 is crucial for the follicle-luteal transition in cattle (Robinson et al., 2007). The LH surge had no effect on VEGF concentrations, so VEGF is thought to play a constitutive, but essential, role in the development of luteal vasculature (Robinson et al., 2007). Both factors and their receptors are expressed at higher concentrations in the developing bovine CL and decrease significantly in the maintenance to regressing CL (Schams et al., 1994; Berisha et al., 2000).

**Small and large luteal cells**

Two morphologically distinct steroidogenic cells types comprise CL of cattle, sheep and many other mammalian species (Niswender and Nett, 1994). Small luteal cells (18-20 μm) are derived from the thecal cells of the follicle, contain receptor for luteinizing hormone (LH), respond to LH by producing cAMP and increasing secretion of progesterone, and usually contain multiple lipid droplets (Niswender, 2002). The number of SLC increased four-fold from day 4 to day 8 in the estrous cycle of the ewe, but these cells maintained a constant volume (Farin et al., 1986). Large luteal cells (25-40 μm) originate from granulosal cells, secrete high basal concentrations of progesterone, and have receptors for both LH and PGF$_2$α (Niswender, 2002). Numbers of LLC did not change throughout the estrous cycle of the
ewe but volume and diameter increased until day 16 of the cycle (Farin et al., 1986). Basal secretion of progesterone differs between these two luteal cell types with LLC producing 2-40 times more progesterone than SLC (Niswender et al., 2000). Both LLC and SLC differ in their response when stimulated with hormones and/or intracellular mediators. Physiological concentrations of LH increased secretion of progesterone from bovine SLC but not LLC in vitro (Alila et al., 1998). Steroidogenic cells in the CL synthesize large amounts of progesterone. Increased synthesis of progesterone occurs as the CL develops and is maintained if a pregnancy is established, or is abolished due to CL regression.

**Progesterone production**

All steroids, including progesterone, are derived from cholesterol, which can come from the plasma membrane, cellular stores (lipid droplets), lipoproteins (low density lipoproteins [LDL] and high density lipoproteins [HDL]), and de novo synthesis (Hu et al., 2010). Each LDL particle contains approximately 2500 cholesterol molecules, which makes this pathway very efficient for progesterone production. Uptake of LDL by luteal cells occurs by receptor-mediated endocytosis (Hu et al., 2010). During this process, LDL binds to its plasma membrane (PM) bound receptor, which then translocates to specialized regions of the PM called coated pits to invaginate and pinch off to form clathrin-coated vesicles (Hu et al., 2010). These vesicles rapidly fuse with lysosomes where the protein of the LDL is hydrolyzed to amino acids and cholesteryl esters are hydrolyzed by a lysosomal acid lipase (Brown and Goldstein, 1979). The LDL-receptor complex rapidly dissociates with the full intact LDL-receptor being sorted and returned to the PM. The resulting cholesterol from the LDL particle can be released and transported to the mitochondria for steroid hormone synthesis (Hu et al., 2010) or undergo esterification by A:cholesterol acyltransferase 1 (ACAT1). Additionally, ACAT1 is responsible
for esterification of de novo synthesized cholesterol. Cholesterol esters are stored in lipid droplets, which have been shown to be hormonally regulated in steroidogenic cells of the adrenal, ovary and testis (Behrman and Greep, 1972; Hu et al., 2010).

The HDL is taken up by the cell by binding to the plasma membrane-bound HDL binding protein (Lestavel and Fruchart, 1994), and cholesterol is mobilized into the cell by an undefined mechanism. Free cholesterol is transported to the mitochondria along the cytoskeletal network and by sterol carrier proteins where it is then transported from the outer to inner mitochondrial membrane by steroidogenic acute regulatory protein (StAR) (Niswender, 2002). Cholesterol is converted to pregnenolone by an inner mitochondrial enzyme, cytochrome P450 side chain cleavage (P450scc). Pregnenolone travels out of the mitochondria and is converted to progesterone by the smooth endoplasmic reticulum enzyme 3β-hydroxysteroid dehydrogenase (3β-HSD) (Niswender, 2002). Additionally, peripheral-type benzodiazepine receptors (PBR) may play a role in cholesterol transport as it is moved from the outer to inner mitochondrial membranes (Niswender, 2002). Targeted deletion of the PBR from Leydig cells drastically reduced steroid secretion, and reintroduction of PBR allowed steroid secretion to be restored (Papadopoulos et al., 1997). Additionally, phosphorylation of PBR by protein kinase A (PKA) enhanced cholesterol transport (Papadopoulos et al., 1997). It is possible that PBR works as a cholesterol channel.

**Luteinization of human mural granulosal cells**

During follicular development granulosal cells primarily secrete estrogen (E₂), but as ovulation approaches they begin to produce progesterone (P₄). During the preovulatory period E₂ secretion is drastically reduced, and granulosal cells begin to synthesize P₄ (McNatty et al., 1979). Additionally, these cells undergo loss of aromatase and mitotic activity (Robertson and
Baker, 1969; Friedrich et al., 1975). Interestingly, P₄ receptors (PGR) cannot be detected with immunohistochemical analysis in human granulosal cells at any follicular stage but are detected readily in human luteal tissue (Suzuki et al., 1994). This finding supports the hypothesis that LH induces PGR expression (Hild-Petito et al., 1988).

Human granulosal cells recovered during oocyte retrieval in patients undergoing in vitro fertilization have been shown to luteinize in culture and, therefore, have been used as a model to study granulosal lutein cells of the CL (Stewart and Vandevoort, 1997). This model has several advantages: no collagenases or other proteolytic enzymes that may influence hormone secretions are needed for cell isolation; unlike ovarian cell line cultures, primary granulosal cell cultures retain their ability to produce and respond to steroid hormones; and serum-free media can be used to examine circulating hormones or serum-derived factors (Hsueh et al., 1984). This model has limitations: absence of vascularization and the influence of angiogenic factors; inability of other follicular cell types such as theca interna or oocyte-cumulus cells to influence granulosal cell differentiation (Hsueh et al., 1984); and representative of only one stage of development of the CL (which has not been clearly defined) and do not take into account the heterogeneity of a granulosal cell population or the fertility status of the patient from which they were isolated.

Luteotropic versus luteolytic signals

Luteotropic hormones

Luteotropic hormones support the growth and/or function of the CL. Luteinizing hormone (LH), growth hormone (GH), prolactin, insulin-like growth factor 1 (IGF-1), prostaglandin (PG) E₂ (PGE₂) and prostacyclin (PGI₂) are all luteotropic hormones. In order to increase P₄ LH acts on the SLC of most species and the LLC in pregnant and pseudopregnant rats as well as nonhuman primates (Hild-Petito et al., 1989; Niswender et al., 2000). Most
luteotropic hormones work by initiating the same intracellular mediator cascade. For example in bovine luteal tissue or cells, LH binds to its G-protein-coupled receptor in SLC and activates adenylate cyclase, which increases concentrations of cyclic adenosine monophosphate (cAMP) and activates PKA (Hoyer et al., 1984). Increase in P4 is achieved through increased transport of cholesterol to the P-450 SCC enzyme complex to the mitochondria (Wiltbank et al., 1993). The LLC also express LH receptor but secrete more basal P4 than SLC, so an increase in response to LH may be due to constitutively expressed PKA (Hoyer et al., 1984).

**Luteolytic hormones**

Luteolytic hormones promote luteal regression. In many species, including cattle, PGF$_2$α is the primary uterine factor responsible for functional and structural luteolysis (McCracken, 1971). Additionally, oxytocin and luteal PGF$_2$α play roles in luteolysis. Increases in protein kinase C (PKC) enzyme activity and intracellular calcium mediate the anti-steroidogenic effects of PGF$_2$α (Niswender et al., 2000). In fact, PGF$_2$α-induced accumulation of extracellular calcium in the cytosolic compartment enhances PKC enzyme activity (Niswender, 2000).

Metabolism and actions of intraluteal PGs, including but not limited to PGF$_2$α, affect the lifespan of the CL. Actions of luteal-derived PGF$_2$α are likely mediated by the G-protein coupled plasma membrane receptor, FP, that uterine PGF$_2$α acts on (Wiltbank and Ottobre, 2003). Expression of FP receptor mRNA is present at 100-fold greater concentrations in the CL than in any other tissue, EP3 receptor is also expressed (Anderson et al., 2001). Koybayashi et al. (2001) found that developing bovine CL (day 3) produce more PGF$_2$α than mid phase CL (days 4-6). In agreement with those results, luteal concentrations of cyclooxygenase-2 (COX-2) mRNA were greater in the developing than mature to regressing CL. Additionally, PGF$_2$α increased the availability of arachidonic acid (AA) and COX-2 activity, both essential for PG
production in the CL (Niswender et al., 2000). This evidence supports the ability of PGF$_2\alpha$ to regulate its own production in the CL. However, a temporal relationship exists in the ratio of luteotropic (PGI$_2$) and luteolytic (PGF$_{2\alpha}$) PGs throughout the estrous cycle (Milvae and Hansel, 1983). Luteal PGs are stimulated by cytokines such as INF$\gamma$ (Fairchild and Pate, 1991) and TNF$\alpha$ (Benyo and Pate, 1992) but suppressed by high intraluteal concentrations of progesterone (Pate, 1988). Although uterine-derived PGF$_2\alpha$ is the main luteolytic factor in ruminants, other PGs appear to play a role in determining the lifespan of the CL.

Luteal regression is not mediated by uterine PGF$_2\alpha$ in primates (Beling, 1970). Infusion of PGF$_2\alpha$ directly into the CL reduced P4 and caused premature menses in monkeys (Auletta et al., 1984). It has been suggested that luteal-derived PGF$_2\alpha$ acts in a paracrine and/or autocrine manner to induce luteal regression (Auletta and Flint, 1988). Despite this evidence, the source and actions of PGF$_2\alpha$ in the primate still remain controversial.

Oxytocin is secreted mainly by the posterior pituitary during and after parturition and lactation in order for milk letdown to occur, but also it is an important factor during luteal regression. Flint and Sheldrick (1982) showed that a rapid increase of ovarian oxytocin was elicited within five minutes of administering a PGF$_{2\alpha}$ analogue (Estrumate®, 125 $\mu$g i.m.). In part due to this finding, McCracken et al. in 1996 suggested that the initiating factor for luteolysis is stimulation of posterior pituitary oxytocin by E$_2$ or the removal of P4 (McCracken et al., 1996), which in turn stimulate secretion of uterine PGF$_{2\alpha}$ (Fairclough et al., 1980). This cascade leads to a positive feedback loop that causes the release of luteal oxytocin as well as luteal and more uterine PGF$_{2\alpha}$ (Silvia et al., 1991) leading to luteal regression. One explanation for the time between pulses of naturally occurring uterine PGF$_{2\alpha}$ is refractoriness of the uterus to oxytocin. However, depletion of oxytocin or its receptors in the bovine CL did not delay
luteolysis (Kotwica and Skarzynski, 1999), implying that it is not critical for luteal regression. The role of oxytocin in luteal regression remain controversial.

**Signals for maintenance of corpora lutea**

During gestation, maternal recognition of pregnancy must occur so that P4 synthesis and secretion continue uninterrupted. Considerable variation exists among species, but, in general, the conceptus secretes factors that are anti-luteolytic (anti-PGF$_{2\alpha}$) or luteotropic (Roberts et al., 1996) to maintain the CL and P4 secretion. In the case of the human, [human] chorionic gonadotropin (hCG) is secreted by the trophoblastic cells of the implanting embryo (Hearn et al., 1991) and binds to the LH receptors in the CL to increase P4 secretion as well to protect the CL from PGF$_{2\alpha}$ (Patton and Stouffer, 1991). In contrast, in the cow, secretion of the protein interferon tau (IFτ) by the trophectoderm of the elongating blastocyst has been postulated to suppress the pulsatile release of PGF$_{2\alpha}$ from the uterus (Roberts et al., 1996).

During a non-fertile luteal phase, luteotropic signals allow the CL to increase in size and ability to secrete P4 (Niswender et al., 2000). However, if a pregnancy is not established, the CL must regress in order for another follicle to grow, ovulate, and a new cycle begin. As mentioned earlier, uterine PGF$_{2\alpha}$ is the luteolytic signal in ruminants. In the cow, both large and small luteal cells are capable of responding to PGF$_2\alpha$ with an increase in intracellular Ca$^{+2}$ (Choudhary et al., 2004). The binding of PGF$_{2\alpha}$ to its receptor in these luteal cells activates two pathways that affect steroidogenesis in the CL. The first is activation of PKC, which directly inhibits progesterone production (Niswender, 2002). Additionally, increases in free intracellular Ca$^{+2}$ lead to apoptosis and cell death (Wiltbank et al., 1989). In primates, the role of PGF$_{2\alpha}$ is controversial but may play a role when it is of ovarian origin. Due to these actions PGF$_{2\alpha}$ is utilized extensively in livestock species to synchronize estrus. However, the bovine CL is
resistant to luteal regression by exogenous PGF$_{2\alpha}$ before day 5 (D 5) of the estrous cycle.

**Estrous synchronization protocols**

The discovery of PGF$_{2\alpha}$ as the luteolysin in ruminant species led to an explosion of synchronization protocols specifically in dairy cattle (Pursley et al. 1995). Synchronizing estrus with exogenous PGF$_{2\alpha}$ allowed for more easily detecting estrus in large herds as well as more efficient artificial insemination (AI) programs (Stevenson and Pursley, 1994). Originally, PGF$_{2\alpha}$ was given twice, 11 days apart (Lucy et al., 1986). However, it was found that while 60% of cows responded to the first PGF$_{2\alpha}$ injection by CL regression only 72% responded to the second injection when all animals should have had CL older than 5 days (Lucy et al., 1986). Due to variability in time to estrus and low conception rates for fixed time AI, GnRH was incorporated into estrous synchronization protocols in order to initiate a new follicular wave with a dominant follicle prior to PGF$_{2\alpha}$ and to more tightly synchronize ovulation 24 hours before breeding (Pursley et al., 1995).

**Responsiveness of corpora lutea to PGF$_{2\alpha}$**

The ability of the CL to respond to the luteolytic actions of PGF$_{2\alpha}$ increases with maturation in cattle. The developing CL (days 1-5) will not regress if a single administration of PGF$_{2\alpha}$ is given, but in the mature CL (days 6-15), the same dose of exogenous PGF$_{2\alpha}$, will induce regression, and the cow will come into estrus within 48-72 hours (Rowson, Tervit and Brand, 1972; Inskeep, 1973). However, differences in response to PGF$_{2\alpha}$ have been noted across timespans during the mature phase of the CL. In 1984, Tanabe and Hann administered a single injection of PGF$_{2\alpha}$ on days 7, 11, or 15 of the estrous cycle in heifers and found that the percentage of heifers exhibiting synchronized estrus at 80 hours after PGF$_{2\alpha}$ tended to increase with stage of estrous cycle (day 7-86%, day 11-90%, and day 15-98%). This area of research has
been under intensive investigation for many years, but the mechanism for the developmental difference is still not understood. However, lack of receptors (Wiltbank, 1995; Wright et al., 2014) or the ability of PGF$_{2\alpha}$ to produce physiological responses (Tsai and Wiltbank, 1998; Sen et al., 2005) do not explain this difference in the developing CL.

**Normal versus induced corpora lutea**

There is evidence that a CL formed after a spontaneous ovulation is characteristically different than one formed when the previous CL is regressed by pharmaceutical methods. In a study by Keisler and Keisler in 1989, CL that were induced in the absence of an existing CL were short-lived whereas CL that were induced during the luteal phase of the estrous cycle did not affect lifespan of spontaneous CL. However, in Brahman cattle, no difference in CL weight, progesterone content, or number of LH receptors were found between spontaneous and induced CL (Hansen et al., 1987).

In multiple species, PGF$_{2\alpha}$ elicits both inhibitory and stimulatory responses to P4 in luteal cells *in vitro*. Most investigators used a non-physiological dose of PGF$_{2\alpha}$ when treating luteal cells *in vitro*. The fact that PGF$_{2\alpha}$ at high concentration is able to activate the PGE$_2$ receptor (Rao, 1974) could, in part, explain increases in progesterone production. Additionally, the entire population of cells that would normally be present in addition to luteinized granulosal cells are not present in all *in vitro* experiments. The inhibitory action of PGF$_{2\alpha}$ on human luteinized granulosal cells may be enhanced by the presence of endothelial and/or immune cell types (Liptak et al., 2005).

**Structural and functional regression of corpora lutea**

**Structural versus functional**

Through decades of luteal physiology research, luteal regression has come to be defined
as either structural or functional (Bowen-Shauver and Telleria, 2003). Functional regression is defined as the cessation of P4 production, whereas structural regression refers to a decrease in weight and size of the CL (Goyeneche et al., 2002) and then disappearance of the structure on the ovary (Bowen-Shauver and Telleria, 2003). Progress has been made in separating these two events in vivo, as well as distinguishing between the two in the literature. However, it is important that these terms are defined in the context of both the species and the reproductive stage being studied (Davis and Rueda, 2002).

Signs of functional regression occur prior to structural regression; presumably the loss of P4 production by the CL ultimately leads to its morphological changes and ultimately structural regression. Initial changes seen in serum P4 are not due to loss of steroidogenic luteal cells, because numbers of large luteal cells do not decrease until after a dramatic decrease in P4 (Braden et al., 1988). Therefore, a decrease in blood flow to the CL and decreased steroidogenic capacity of luteal cells is one possible explanation for this decrease in P4. However, a decrease in P4 prior to a decrease in luteal blood flow has been seen in rodents (Behrman et al., 1979) rabbits (Keyes et al., 1983) and domestic species (Ginther et al., 2007). Studying spontaneous luteolysis has proven to be difficult because timing, duration and number of pulses of uterine PGF\(_{2\alpha}\) differ among individuals. Decreased mRNA and protein of enzymes (3β-HSD, StAR, etc.) that contribute to luteal regression also are expressed at inconsistent times, which makes finding significant changes within a group of animals difficult due to the variation. Due to this fact, researchers have focused on subsequent changes in these key factors after administration of PGF\(_{2\alpha}\).

Structural regression occurs due to a series of physiological changes that result in distinct morphological changes in the CL. One such event that contributes greatly to the demise of the
CL is apoptosis, which is promoted by PGF$_{2\alpha}$ (Sawyer, et al., 1990). Apoptosis, also known as programmed cell death, is defined as an active, energy dependent process by which nonessential populations of cells are removed by other cells from tissue (Kerr et al., 1972), a mechanism that allows for control of cell numbers and tissue size. Apoptosis in endocrine tissues commonly occurs after the tropic hormone is removed or a negative stimulus is activated. Cells shrink, and fragments of both nuclear and cytoplasmic origin appear (Kerr et al., 1972). Apoptosis is categorized into the intrinsic and extrinsic signaling cascade. The intrinsic pathway is activated by apoptotic signals that originate within the cell in response to insults such as drugs, radiation, or growth factor withdrawal. As a result, a change in mitochondrial permeability occurs through modifications in the ratio of pro- to anti-apoptotic Bcl-2 family members (Adams and Cory, 1998). Alternatively, the extrinsic pathway is activated by extracellular signals, such as Fas, or tumor necrosis factor alpha (TNF-\(\alpha\)), that bind with cell surface receptors to initiate cell death (Nagata, 1997). Caspases, a family of intracellular proteases, are activated by both pathways and cleave multiple cellular components that contribute to the typical morphological appearance in apoptotic cells (Martin and Green, 1995). These fragments, known as apoptotic bodies, are targets for macrophages and other immune cells.

Specifically, the proportion of steroidogenic luteal cells in the CL of ewes decreased within 24 hours of PGF$_{2\alpha}$ treatment during the maintenance phase (Braden et al., 1998). Both the number and size of LLCs decreased, and this was followed by a decrease in the number of SLC (Braden et al., 1998). Morphological changes in the steroidogenic cell population of the CL do not become visible until 24-36 hours post PGF$_{2\alpha}$ (Sawyer et al., 1990), yet their ability to produce P4 has already been compromised. This indicates that the structural changes in the CL are a consequence of reduced P4 production and not vice versa. However, it could indicate that
functional regression is regulated by a different mechanism than apoptosis, such as autophagy.

Indeed, autophagy has been shown to play a role in structural regression in human and rats. Autophagy is an intracellular bulk degradation system in which portions of cytoplasm are enveloped into autophagosomes that then undergo maturation and fusion with lysosomes for degradation (Klionsky and Emr, 2000). The appearance of cytoplasmic vacuoles, prominent lysosomes, and the accumulation of lipofuscin pigment without any changes in nuclear morphology in the regressing primate CL strongly indicate that autophagy induces or at least plays a part in the death of steroidogenic luteal cells (Morales et al., 2000). Additionally, autophagy has been shown to play a role in luteal regression in the rat by accumulation of autophagosomes, which then induce apoptotic cell death in luteal cells (Choi et al., 2011). Autophagy is promoted by adenosine monophosphate activated protein kinase (AMPK; Kim et al., 2011). The role autophagy plays in bovine luteal regression has yet to be examined. However, AMPK reduced P4 production in murine, galline and bovine granulosal cells (Tosca et al., 2005; Tosca et al., 2006; Tosca et al., 2007), which will be discussed at greater length later in this review. The link between AMPK and autophagy induction in bovine luteal tissue has not been determined.

**Factors involved in regression**

**Luteal blood flow**

Changes in luteal blood flow have been proposed to be one of the main mechanisms by which PGF$_{2\alpha}$ exerts its luteolytic actions (Nett et al., 1976; Knickerbocker, 1988). In ewes, PGF$_{2\alpha}$ reduces blood flow to the CL concomitantly with decreased secretion of P4 (Nett et al., 1976). Color Doppler ultrasound is a non-invasive and useful technique to examine blood flow within the CL during its development (Acosta et al., 2003) and demise (Acosta et al., 2002). On
days 16-18 of the estrous cycle in cattle, during spontaneous luteolysis, an increase in blood flow surrounding the CL was observed followed by decreased plasma P4 the next day (Shirasuna et al., 2004; Miyamoto et al., 2005). An acute increase in blood flow (0.5 to 2 hours) also was seen during PGF$_{2\alpha}$-induced luteolysis in mature (day 10) but not developing (day 4) bovine CL (Acosta et al., 2002). This increase was followed by a decrease in plasma P4 at 0.5 hours and blood flow at 8 hours (Acosta et al., 2002). Expression of vasoactive factors must increase in order to mediate this increase in blood flow. One such factor, nitric oxide (NO), a vasodilator, was proposed to be stimulated by PGF$_{2\alpha}$ (Yamamoto et al., 2004). Indeed, in a different tissue, it has been postulated that uterine PGF$_{2\alpha}$ induces NO production from large arterioles to acutely increase luteal blood flow and initiate luteolysis (Miyamoto et al., 2005). However, other investigators were unable to find an increase in luteal blood flow at the initiation of luteolysis (Ginther et al., 2007). In fact, a decrease in blood flow was not observed until the day after P4 decreased (Ginther et al., 2007).

**Protective effect of progesterone**

Luteal P4 has been shown to exert protective effects on CL as a luteotropin by increasing the number of LH receptors in the early CL (Rothchild, 1981) and through anti-apoptotic activity (Rueda et al., 1997). Direct supplementation of P4 to dispersed mature bovine luteal cells initiated a decrease in prostacyclin and PGF$_{2\alpha}$ synthesis (Pate, 1988), which indicates that P4 action could also be anti-luteolytic. This evidence shows that P4 can protect the lifespan of the CL by blocking a decrease P4 production, reducing cell death, and preventing luteolytic factors such as PGF$_{2\alpha}$ from regressing the CL. The P4 receptor, PR, is present in LLC and some SLC, but not endothelial cells (Rueda et al., 2000), and it was further delineated that two isoforms of PR, PR-A and PR-B, are expressed in the bovine CL (Kotwica et al., 2004). Protein synthesis of
these classical genomic receptors changes throughout the estrous cycle and pregnancy with the highest concentrations found at days 5-10 of the cycle and no expression in the regressing CL (Kotwica et al., 2004). Actions of P4 could be exerted through the genomic receptor; however, these effects could be mediated by the glucocorticoid receptor (Sugino et al., 1997) or through the non-genomic plasma membrane bound PR (Peluso, 1997).

**Pulsatile uterine PGF$_{2\alpha}$**

Prostaglandin F2α is released in a series of pulses from the non-pregnant uterus in order to regress the bovine CL (Niswender, 2002). Uterine release of PGF$_{2\alpha}$ is transferred to the ovary through the utero-ovarian vein via a counter current exchange mechanism (McCracken et al., 1972). This helps ensure that PGF$_{2\alpha}$ is not enzymatically inactivated in the lungs by PG dehydrogenase (PGHD) (Piper et al., 1970). Transport of PGs through plasma membranes is poorly understand, but prostaglandin transporter (PGT) mediates the efflux and influx of PGF$_{2\alpha}$ (Schuster, 1998). Recently Lee et al. (2013) showed PGT protein was required for pulsatile release of PGF$_{2\alpha}$ from the endometrium and to maintain a functional CL in sheep. Number and frequency of pulses vary considerably during natural ruminant luteal regression but approximately 5 distinct pulses, each around 4-hour duration over a period of approximately 30 hours, are required for luteal regression in heifers (Mann and Lamming, 2006). In cattle, P4 decreases during the ascending portion of a naturally occurring uterine PGF pulse but increases during the descending portion of the PGF pulse. This is due to a pulse of LH stimulated by decreased P4, as well as loss of the negative effect of PGF. A complete rebound in P4 occurs after each of a series of pulses of PGF$_{2\alpha}$ before the first luteolytic pulse of PGF$_{2\alpha}$ (Ginther and Beg, 2013) that causes a terminal decrease in P4.
Luteal endothelial cells

Luteal endothelial cells contribute to luteal demise. Endothelial cells express PGF$_{2\alpha}$ receptor (Mamluk et al., 1998), and administration of PGF$_{2\alpha}$ caused degeneration of these cells (O’Shea et al., 1979) leading to reduced capillary density (Braden et al., 1988) and ultimately to decreased blood flow to the luteal parenchyma. Endothelial cells appear to be one of the first cell types to be affected by PGF$_{2\alpha}$ treatment by showing signs of apoptosis (Sawyer et al., 1990). Endothelial cells and more specifically the luteal endothelial cell product endothelial-I (ET-1), could mediate the luteolytic actions of PGF$_{2\alpha}$ (Girsh et al., 1995). Choudhary et al., 2005 demonstrated that ET-1 increased concentration of intracellular Ca$^{2+}$ in bovine luteal steroidogenic cells in vitro. Additionally, SLC and LLC from mature (day-10) CL responded with a greater increase in Ca$^{2+}$ than SLC and LLC of developing (day-4) CL (Choudhary et al., 2002). However, when different doses of ET-1 or PGF$_{2\alpha}$ were administered to bovine luteal endothelial cells in vitro, no differences between the developing (day-4) and mature (day-10) CL were observed (Choudhary et al., 2005). Basal and LH-stimulated P4 were reduced by ET-1 in the developing and mature bovine CL with similar responses (Choudhary et al., 2005). The authors concluded that ET-1 is a tonic inhibitor of P4 secretion rather than a mediator of the action of PGF$_{2\alpha}$, and does not contribute to the insensitivity of the developing CL to luteolytic actions.

Luteal immune response

Immune cells such as macrophages, neutrophils, T cells, and eosinophils are present in both the developing and regressing CL. In the bovine CL, markers of CD5+ or CD8+ T cells increased during stage IV (days 19-21) of the estrous cycle and from day 16 onward in natural luteal regression (Penny et al, 1999). Similarly, markers of leukocytes, specifically macrophages
and T lymphocytes, significantly increased during luteal regression in cattle (Penny et al., 1999; Bauer et al., 2001). During induced luteolysis of mature bovine CL, mRNA of TNF-α, interferon gamma (IFN-γ), and interleukin 1-beta (IL-1β) were all increased as soon as two hours after injection of PGF$_{2α}$ and continued to be elevated throughout 12 hours post PGF$_{2α}$ (Neuvians et al., 2004). The wide range in time spanned by this influx of cytokines may indicate that they play roles in both functional and structural regression. Up to 70% of proliferating cells in the regressing bovine CL are CD14+ macrophages (Bauer et al., 2001). Recruitment of macrophages to the CL by the cytokine monocyte chemoattractant protein 1 (MCP-1) is enhanced by the ability of immune and endothelial cells to interact with one another (Shirasuna et al., 2012). Penny (2000) proposed that on day 18 of the estrous cycle, PGF$_{2α}$ acts on various cell types in the CL, such as T-lymphocytes, endothelial cells, fibroblasts, and possibly immune cells directly, to induce MCP-1 production. In the next 24 hours an influx of macrophages occurs in response to MCP-1, which is proposed to serve as an additional source of MCP-1 as well as playing an active role in structural regression (Penny, 2000). Additionally, activated immune cells seem to produce a factor that impairs luteal cell P4 production because PGF$_{2α}$ enhances the production of luteal chemoattractant, and macrophages and eosinophils decrease P4 secretion in the midphase CL in cattle (Pate, 1995).

**Signal Transduction Mechanism**

The mechanism governing the action of PGF$_{2α}$ is thought to involve activation of the phospholipase C-Ca$^{2+}$ pathway (Davis et al., 1987). The binding of PGF$_{2α}$ to its cognate G$_q$-protein-coupled receptor causes dissociation of the α subunit from the β/γ subunit. The α subunit exchanges GDP for GTP to activate phospholipase C (PLC; Davis et al., 1988). Then, PLC cleaves plasma membrane-bound phosphatidylinositol 4,5-bisphosphate (PIP$_2$) to generate free
IP3 (Berridge et al., 1983) and membrane-bound 1,2-diacylglycerol (DAG) (Berridge, 1987). Increase in both the concentration of cytoplasmic Ca²⁺ and activation of PKC are, in part, intracellular mediators of the actions of PGF₂α in luteal cells (Wiltbank et al., 1990). Even though it is generally referred to as PKC in signal transduction pathways, PKC is a family of protein kinases that can be divided into three subfamilies: classic (cPKC; α, βI, βII, and γ), novel (nPKC; δ, ε, η, and θ), and atypical (aPKC; ζ, τ, and λ) (Kang, 2014). Sen et al. (2003) hypothesized that differential expression of PKC isozymes during the estrous cycle might explain inability of the developing CL to mount the same luteolytic response to PGF₂α that is seen in the mature CL. Indeed, both PKCε and PKCβII were expressed at higher concentrations in the mature CL (day 10) compared to the developing CL (day 4). In both developing and mature CL, PGF₂α had the capacity to activate conventional PKC isozymes. Therefore the authors reasoned that these isozymes do not contribute to the insensitivity of developing CL to luteolytic actions of PGF₂α (Sen et al., 2003). Specific inhibitors of PKCε significantly decreased the PGF₂α-induced rise in intracellular Ca²⁺ in LLC and SLC that in turn had consequences (at least in part) in the ability of PGF₂α to inhibit LH-stimulated P4 secretion at this developmental stage (Sen et al., 2005). Additionally, downregulating expression of PKCε reduced the ability of PGF₂α to inhibit P4 synthesis/secretion (Goravanahally et al., 2007).

In other studies, the next intracellular mediator, cytosolic Cat⁺², was investigated. Developmental differences in the ability of PGF₂α to induce increases in intracellular Ca⁺² have been demonstrated in both LLC and SLC (Choudhary et al., 2005). In vitro, PGF₂α evoked a larger increase in intracellular Ca⁺² in both SLC and LLC from mature (day-10) than developing (day-4) CL, and percentage of day-10 SLC responding increased as a function of PGF₂α concentration (Choudhary et al., 2005).
Goravanahally et al., (2009) provided evidence that the luteolytic actions of increases in intracellular Ca\(^{2+}\) may be mediated through calmodulin dependent kinase kinase 2 beta (CAMKK2) because mRNA expression of CAMKK2 was upregulated in mature versus developing bovine CL. Additionally, CAMKK2 gene expression was upregulated in mature bovine CL when PGF\(_{2\alpha}\) was administered in vivo 24 hours before CL removal (Goravanahally et al., 2009). Whether CAMKK2 affects steroidogenesis directly or indirectly by acting upon another intermediary target is not known. One such target is adenosine monophosphate activated protein kinase (AMPK; Wong, 2009) and it is a possible mediator for the luteolytic actions of PGF\(_{2\alpha}\).

**PGF\(_{2\alpha}\) signal transduction pathway**

**Adenosine monophosphate activated protein kinase (AMPK)**

Adenosine monophosphate activated protein kinase is an important regulatory protein for cellular energy balance in multiple cell types. In all eukaryotic cells, AMPK is a heterotrimeric complex composed of one catalytic subunit \(\alpha\) (with two isoforms: PRKAA1 and PRKAA2), and two regulatory subunits \(\beta\) (with two isoforms: PRKAB1 and PRKAB2) and \(\gamma\) (with three isoforms: PRKAG1, PRKAG2, and PRKG3) (Wong et al., 2009). Different types of cells and tissues express distinctly different combinations of these subunits. Activation of AMPK by phosphorylation of threonine 172 of the \(\alpha\) subunit by upstream kinases, LKB1 or CAMKK2, is required for its activity (Wong et al., 2009). A serine/threonine kinase, LKB1, is the primary upstream kinase of AMPK and appears to be constitutively active (Hardie, 2008). When energy concentrations are depleted resulting in a decrease in ATP along with an increase in AMP, LKB1 phosphorylates Thr\(^{172}\).

Specific tissues, such as neurons and endothelial-derived cells, express an alternate
pathway for activation of AMPK (Hardie, 2008). Increases in cytoplasmic Ca\(^{+2}\) lead to activation of CAMKK2, which is capable of phosphorylating AMPK at Thr\(^{172}\). Free catalytic α subunits are usually inactive due to the presence of an autoinhibitory domain that is located in the center of the subunit. Autoinhibition is eliminated when the α subunit forms a functional complex with the β and γ subunits. The β subunit has key autoregulatory function in the AMPK complex, partially due to a specific sequence that binds to glycogen molecules (Guzman et al., 2009). A myristoylation site serves as a switch for reversible membrane binding of the complex, and a direct autoinhibitory domain is located within the β subunit (Guzman et al., 2009). The γ subunit is defined by the two Bateman domains, which selectively bind molecules that contain adenosine such as AMP or ATP (Bateman, 1997). The binding of AMP to the γ subunit activates AMPK by inducing conformational changes in the complex that promote phosphorylation of Thr\(^{172}\). Binding of ATP inhibits these changes. Phosphorylation by AMPK of proteins that inhibit anabolic processes, such as those involved in metabolic pathways that reduce protein synthesis and cell proliferation and increase catabolic pathways that create energy such as glucose uptake, mitochondrial biogenesis, glycolysis and lipid oxidation (Hardie, 2008). Recently, activated AMPK has been shown to decrease P4 secretion in rat, cow, and primary F1 hen granulosal cells (Tosca et al., 2005; Tosca et al., 2007; Tosca et al., 2006). All subunits and their isoforms of AMPK have been characterized in bovine whole ovary, CL, small and large follicles as well as granulosal cells (Tosca et al., 2007).

**Pharmaceutical agents**

Thiazolinediones (TZDs) have been used as a form of medication for type II diabetes mellitus since the 1990’s. This is the only approved use, TZD is being experimentally tested to treat polycystic ovarian syndrome (PCOS; Belfort et al., 2006), autism (Boris et al., 2007), and
ovarian hyperstimulation syndrome (Shah et al., 2010). Peroxisome proliferator-activated receptors (PPAR), which are endogenously activated by free fatty acids and prostanoids, are also activated by TZDs. Once activated, PPAR dimerizes with retinoid X receptor (RXR) to induce a specific cohort of gene expression. In general, TZDs decrease triglycerides and increase high-density lipoprotein cholesterol (HDL-C).

Dorsomorphin dihydrochloride (DM), also known as Compound C, is a potent selective, cell-permeable and reversible AMPK inhibitor (Liu et al., 2014). Dorsomorphin inhibited AMPK induced either by 5-aminoimidazole-4-carboxamide-1-β-4-ribofuranoside (AICAR) (Tang et al., 2011) and metformin (Isakovic et al., 2007). However, DM inhibited other kinases in addition to AMPK (Bain et al. 2007).

One cell-permeable allosteric activator of AMPK is AICAR. It is metabolized intracellularly to ZMP (5-aminoimidazole-4-carboxamide-1-β-D-ribofuranotide) by adenosine kinase (Merrill et al., 1997). Phosphorylated AICAR, ZMP, is an AMP analog that interacts with the γ subunit of AMPK and induces kinase activation through allosteric changes in AMPK conformation (Merrill et al., 1997). Although AICAR is used extensively in the laboratory, it is not ideal for clinical scenarios because of its short half-life, requirement for intravenous infusion, and variable effectiveness among patients (Wong et al., 2009).

Metformin is another well-known activator of AMPK, although it activates the heterotrimeric complex indirectly. Metformin is a biguanide that is developed from galegine, which is a derivative of the guanidine found in *Galega officinalis* (French lilac). Metformin specifically works in the mitochondria by inhibiting the mitochondrial respiratory chain complex I. Although the exact mechanism of action has yet to be elucidated, metformin inhibits complex I without affecting any other steps in the respiratory chain (Viollet et al., 2012). Ultimately,
Metformin reduces the synthesis of ATP by decreasing not only NADH oxidation but also proton pumping across the inner mitochondrial membrane and rate of oxygen consumption (Viollet et al., 2012). This reduction in ATP results in a shift in the AMP: ATP ratio and causes the AMPK complex to become phosphorylated and therefore active (Viollet et al., 2012).

Metformin is the most widely used drug for the treatment of type II diabetes, being prescribed to nearly 120 million people worldwide (Viollet et al., 2012). It inhibits hepatic gluconeogenesis and augments the rate of glucose uptake in skeletal muscle (Bailey, 1996). Recently, metformin was used to treat women experiencing PCOS who were overweight or insulin resistant. Metformin’s ability to increase insulin sensitivity in individuals made it an ideal candidate for the treatment of PCOS. Metformin increased insulin sensitivity in patients with PCOS and it increased ovulation rate, improved menstrual cyclicity, and reduced serum androgen (Tang et al., 2010). However, in 2007 Legro et al. showed that clomiphene citrate was superior to metformin in achieving live birth rates in infertile women with PCOS. Metformin appears to work in PCOS patients by both direct ovarian effects and secondarily through decreasing insulin resistance. In bovine granulosal cells isolated from small antral follicles steroidogenesis was affected by metformin treatment (Tosca et al., 2007). Metformin (10 mM) reduced both P4 and E2 production as well as key enzymes (HSD3β, CYP11A1, and STAR), with or without FSH and/or IGF1 in the media (Tosca et al. 2007).

Interestingly, metformin is a hydrophilic base that is excreted unchanged in the urine. Metformin cannot easily diffuse through cell membranes and is not metabolized into other variants when being eliminated from the body (Gruzman et al., 2009). Metformin is a small molecule that has many transporters, known as organic cation transporters, in the kidney that lead to its high rate of renal clearance (Wong et al., 2009).
Model of activation of AMPK in the corpus luteum

Changes in P4 biosynthesis due to activation of AMPK in the bovine CL stem from increases in intracellular Ca$^{+2}$. In addition to activating PKC, rises in Ca$^{+2}$ activate calmodulin and therefore CAMKK. The latter is responsible for phosphorylating AMPK and converting it to its active form that traditionally monitors the energy balance of the cell, but AMPK may not have this role in steroidogenic cells. Typically, activation of AMPK results in a shift from anabolic processes, such as synthesis of proteins like enzymes involved in P4 production, to catabolic processes. Additionally, AMPK activation leads to phosphorylation of fatty acid synthase (FAS), and acetyl-CoA-carboxylase (ACC1), key enzymes in de novo lipid synthesis. It may also regulate cholesterol uptake through lipoprotein receptors. Progesterone is a steroid hormone and is derived from cholesterol that is made up of lipids. This provides another mechanism to reduce overall P4 production (Figure 1.0).

Statement of the problem

The literature established that developmental differences exist in multiple facets of physiology between developing and mature bovine CL. Changes in FP expression or PGF$_{2\alpha}$ ligand, luteal blood flow, endothelial system, immune system, and the signal transduction mechanism have all been investigated in an attempt to explain the insensitivity of the developing CL to a luteolytic dose of PGF$_{2\alpha}$. However, FP receptor expression is similar among developing and mature bovine CL, and, furthermore, PGF$_{2\alpha}$-mediated actions have been demonstrated at these two luteal developmental stages Therefore, a difference in the components of the signal transduction mechanism activated by PGF$_{2\alpha}$ upon binding to its cognate FP exits between developing and mature CL. In agreement with this idea, PGF$_{2\alpha}$ elicited a difference in Ca$^{+2}$
response between the two developmental stages of the bovine CL, and a down-stream target of 
Ca\(^{2+}\), CAMKK2, was differentially expressed in developing and mature CL. CAMKK2 is 
known to phosphorylate AMPK, but this component of the signal transduction pathway has not 
been investigated in relation to reproductive endocrinology. Therefore, the goal of this project is 
to investigate alternative downstream components activated by the rise in cytosolic [Ca\(^{2+}\)] 
initiated when the FP receptor is activated in both bovine luteal tissue and luteinized human 
granulosal cells. The objectives of the research were: to determine the changes in P4 secretion \textit{in 
vitro} and \textit{in vivo} in the presence of an AMPK inhibitor or activator in developing and mature 
bovine CL; define changes in AMPK that occur as a result of FP activation; define changes in 
cholesterol transport that occur as a result of PGF\(_{2\alpha}\) binding to FP, which may be monitored by 
AMPK; define culture conditions and responses to luteotropic and luteolytic signals by luteinized 
human granulosal cells derived from patients with various infertility causes. The proposed 
research has one central hypothesis: developmental differences in components of the signal 
transduction associated with FP expressed in developing and mature CL explain, at least in part, 
the developmental difference in the ability of PGF\(_{2\alpha}\) to induce functional luteal regression.
Figure 1.0: Model of actions of AMPK in luteal cells. Changes in progesterone biosynthesis due to activation of AMPK in the bovine CL stem from increases in intracellular Ca^{2+}, calmodulin, and CAMKK2. CAMKK2 phosphorylates AMPK and converting it to its active form results in a shift from anabolic processes to catabolic processes. Additionally, AMPK may regulate cholesterol uptake through lipoprotein receptors (LDLR and SRB-1). This provides another mechanism to reduce overall progesterone production in addition to the established role of PKC.
Chapter 2: Effects of prostaglandin F 2 alpha and adenosine monophosphate activated kinase on progesterone production in the bovine corpus luteum in vivo

Introduction

The ability of bovine corpus luteum (CL) to respond to luteolytic actions of prostaglandin F 2 alpha (PGF$_{2\alpha}$) increases after ovulation in cattle. Developing CL (days 1-5) fail to regress if a single exogenous bolus of PGF$_{2\alpha}$ is given, yet administration of PGF$_{2\alpha}$ at the same dose when the CL is at the mature stage (days 6-15) will induce regression with the cow returning to estrus within 48-72 hours (Rowson et al., 1972; Inskeep, 1973). The cellular mechanisms responsible for this developmental difference are unclear. Lack of receptors (Wiltbank, 1995; Juengel et al., 1998; Wright et al., 2014) or ability to elicit a physiological response to PGF$_{2\alpha}$ (Tsai and Wiltbank, 1998; Choudhary et al., 2005; Sen et al., 2005) does not appear to explain the observed differences. As discussed below, differences in the signal transduction mechanisms associated with FP receptor might explain some of the characteristics of the elicited response to PGF$_{2\alpha}$ in mature versus developing CL.

In luteal cells, PGF$_{2\alpha}$ has affects by binding to its cognate $G_q$ protein-coupled receptor and activating phospholipase C (PLC; Davis et al., 1988). Then, PLC cleaves plasma membrane-bound phosphatidylinositol 4,5-bisphosphate (PIP$_2$) to generate free IP3 (Berridge et al., 1983) and membrane-bound 1,2-diacylglycerol (DAG) (Berridge, 1987). Water-soluble IP3 increases the concentration of cytoplasmic Ca$^{2+}$, and DAG, which in combination with Ca$^{2+}$, activates protein kinase C (PKC). Increases in both cytoplasmic Ca$^{2+}$ and activation of PKC have been demonstrated, in part, to mediate the actions of PGF$_{2\alpha}$ in luteal cells (Wiltbank et al., 1990). Differential expression of genes encoding particular isoforms of PKCs and genes
participating in Ca\textsuperscript{2+} homeostasis have been implicated in acquisition of luteolytic capacity by the bovine CL. In agreement with this suggestion, increased expression of two PKC inhibitors, histidine triad nucleotide binding protein (H1NT1) and tyrosine 2-monooxygenase\tryptophan 5-monooxygenase activation protein zeta polypeptide (YWHAZ), appear to be involved in insensitivity of the early ovine CL (Juengel et al., 1998). Indeed, YWHAZ gene expression was downregulated in mature bovine CL (Goravanahally et al., 2009). Greater concentrations of PGF\textsubscript{2α} are needed to elicit Ca\textsuperscript{2+} responses in steroidogenic cells isolated from developing than from mature CL (Choudary et al., 2005). Additionally, increased gene expression of CAMKK2, a down-stream target of Ca\textsuperscript{2+}, is more relevant in the activated signal transduction pathway in mature than in developing CL (Goravanahally et al., 2009). After an agonist-mediated rise in intracellular Ca\textsuperscript{2+} CAMKK2 was activated via phosphorylation by calcium/calmodulin-dependent kinases (CAMKs) Whether CAMKK2 affects steroidogenesis directly or if it acts upon another intermediary step that impinges on steroidogenesis is not known.

One intermediary step could be activated adenosine monophosphate activated kinase (AMPK), which is activated via phosphorylation by CAMKK2 (Wong, 2009). Typically, activation of AMPK results in a shift from anabolic processes, such as protein synthesis or progesterone production, to catabolic processes. Activation of AMPK may regulate progesterone production by altering cholesterol uptake through high and low-density lipoprotein receptors such as scavenger receptor class B member 1 (SRB-1) and low-density lipoprotein receptor (LDLR). Cholesterol influx is critical in steroidogenic cells because it is the preferred source over \textit{de novo} synthesis in this tissue. Both LDL and HDL provide substrate for progesterone production. During luteal regression in primates, gene expression for lipoprotein receptors SRB-1 and LDLR decreased (Bogan et al., 2009, 2010). Downregulation of LDLR activity in human
fibroblast cells increased expression of acyl CoA cholesterol acyltransferase (ACAT) (Brown et al., 1975), which is responsible for esterification of free cholesterol. These cholesterol esters are stored in lipid droplets and are hormonally regulated in steroidogenic cells of the adrenal, ovary and testis (Behrman and Greep, 1972; Hou et al., 2010). The possibility that PGF$_{2\alpha}$ predicates luteolytic actions of AMPK regulation changes in cholesterol transport has yet to be explored in the bovine CL.

One well known activator of AMPK is metformin, which is the most widely used pharmaceutical for treatment of type II diabetes being prescribed to nearly 120 million people worldwide (Viollet et al., 2012). Metformin activates AMPK by reducing synthesis of ATP through NADH oxidation, which results in a shift in AMP: ATP ratio and phosphorylation of the AMPK complex (Viollet et al., 2012). In addition, metformin has been used to treat polycystic ovary syndrome (PCOS) in women who were overweight or insulin resistant. Metformin directly affects the ovaries and also decreases insulin resistance in PCOS patients. Additionally, metformin has been shown to reduce progesterone and estradiol production in bovine granulosal cells isolated from small antral follicles (Tosca et al., 2007). Reduced steroidogenesis resulting from metformin treatment indicates a potential role in progesterone production in the bovine ovary.

Therefore, the objective of this study was to investigate if AMPK played a role in the signal transduction pathway used by FP receptors in mature bovine CL. The specific aims of this research were to determine changes in progesterone secretion in vivo in response to activation of AMPK via metformin in mature bovine CL; define changes in protein expression of AMPK, phosphorylated AMPK (P-AMPK) and steroidogenic acute regulatory (StAR) resulting from FP activation, and characterize cholesterol transport proteins (LDL, SRB-1, ACAT-1) during
functional regression after activating FP via exogenous PGF$_{2\alpha}$. The overall hypothesis is that luteal developmental differences in signal transduction associated with FP can explain, at least in part, the differences in the ability of PGF$_{2\alpha}$ to induce functional luteal regression in the mature but not the developing bovine CL.

Materials and Methods

Non-lactating beef cows were observed twice daily for estrus at approximately 12 h intervals for 30 min per observation. The day when standing estrus was observed was designated as day 0 (d0). The West Virginia University Animal Care and Use Committee reviewed and approved the protocol for the tissue collection (ACUC # 01-0809).

Experiment 1

Cows (450-700 kg BW) were assigned randomly among treatment groups: day 10 hour 0 (d10-h0; n = 5), day 10 hour 2 (d10-h2; n = 5), day 10 hour 4 (d10-h4; n = 5). Cows received a subcutaneous injection of PGF$_{2\alpha}$ (25 mg, Lutalyse®; Zoetis Florham Park, NJ) or saline (d10-h0) 0, 2, or 4 hours prior to lutectomy. Corpora lutea (CL) were collected via supravaginal incision under epidural anesthesia (Casida, 1959) with administration of 6-9 mL of 2% lidocaine (Butler Company, Columbus, OH). CL were weighed and divided into equal sections for protein isolation and measurement of luteal progesterone. Tissue for protein isolation was snap frozen in liquid nitrogen and stored at -80°C; remaining tissue was transported to the laboratory in ice-cold saline. Tissue for luteal P4 was homogenized with an Omni Tissue Homogenizer (Omni International, Kennesaw, GA) and then frozen at -20°C until assayed for P4 via RIA. Blood samples were collected hourly via caudal venipuncture until CL excision. Samples were allowed to clot and then centrifuged at 3500 x g for 15 minutes, and serum was collected and stored at -20°C. Serum and CL assayed by RIA for P4 as previously described (Sheffel et al., 1982).
Luteal proteins were analyzed using semiquantitative Western blotting as previously described (Sen, 2004). Briefly, protein was isolated from d10-h0, d10-h2 and d10-h4 CL by homogenizing the luteal tissue in a buffer containing 50 mM Tris HCl, 150 mM NaCl, 1 mM EGTA, 1 mM EDTA, 0.1% SDS, 1% Triton-X, protease inhibitors (2 mM phenylmethylsulfonylfluoride, 5 μg/μl leupeptin, 5 μg/μl aprotinin), and phosphatase inhibitors (100 mM sodium fluoride, 2 mM sodium orthovanadate, 10 mM sodium pyrophosphate). Sample protein concentration was determined using a Bio-Rad (Hercules, CA) assay with bovine serum albumin (BSA; Gibco BRL) as a standard. After concentrations were determined, 100 μg/lane of protein was loaded for StAR, AMPK, P-AMPK, SRB-1, and LDLR. Proteins were resolved with a 4-15% gradient polyacrylamide TGX gel (Bio-Rad, Hercules, CA) and transferred to polyvinylidene difluoride (PVDF) membrane (Millipore, Billerica, MA) using a Bio-Rad Mini Trans-Blot Cell (Hercules, CA).

Primary antibodies used included 1) StAR antibody (#bs-3670R, Bioss, Boston, MA) at 1:250 (v/v) 2) AMPK antibody (#2532S, Cell Signaling Technology, Danvers, MA) 1:1000 (v/v) 3) P-AMPK antibody (#2537S, Cell Signaling Technology, Danvers, MA) 1:1000 (v/v) 4) LDL-R antibody (LS-C146979, LSBio, Seattle, WA) 1:500 (v/v) 5) SRB-1 antibody (#ab24603, AbCam, Cambridge, MA) (1:1000) 6) and beta actin (A2228; Sigma-Aldrich, St. Louis, MO) at a dilution of 1:2000 (v/v) and incubated overnight at 4°C with gentle shaking. Secondary antibodies included goat anti-mouse IRDye 680RD (for actin) diluted to 1:10000 and goat anti-rabbit IRDye 800CW (for StAR, AMPK, P-AMPK, LDL-R, SRB-1) diluted to 1:10000 (v/v) (#926-68170, #827-08365, Li-Cor, Lincoln, NE). A two-color detection scheme was used to permit simultaneous probing for target proteins. Band intensity imaging was captured using Odyssey infrared imaging software and quantified through densitometry (ImageJ Bethesda,
Maryland). Signal intensity for a protein of interest was standardized to the corresponding intensity of actin in the same sample. This normalization procedure for semiquantitative estimation of protein was validated by Sen et al. (2004).

**Materials and Methods**

Non-lactating beef cows were observed twice daily for estrus at approximately 12 h intervals for 30 min per observation. The day of standing estrus was designated as day 0 (d0). The West Virginia University Animal Care and Use Committee reviewed and approved the protocol for the tissue collection (ACUC # 12-1008).

**Experiment 2**

Cows were assigned randomly to treatment groups, which received 15 (Met-15, n = 4) or 30 (Met-30, n = 4) mg/kg of BW of metformin (New England Mail Order Pharmacy, Middlebury, VT) every 3 hours for 12 hours on d10 of the estrous cycle. Metformin was pulverized and dissolved into sterile saline solution before centrifugation at 119 RCF for 5 minutes to remove non-active binding ingredients. Supernatant was collected for infusion, the pellet was washed; and the supernatant was collected and added to the total infusion volume. An artificial insemination catheter was passed through the cervix, and metformin was deposited into the uterus by a syringe connected to the catheter by a 200 μl pipette tip. Based on observations from this study, a dose of 30 mg/kg BW of metformin (metformin, n = 7) or saline (control, n = 7) was injected into the jugular vein of cows (n = 7) every 3h for 12 hours. Blood samples were collected one day prior to and hourly for 12h after the first metformin infusion and then one and two days after treatment. Blood samples were allowed to clot, and then centrifuged at 3500 x g for 15 min. Serum was collected and stored at -20°C until assayed by RIA for P4 (Sheffel et al., 1982) and for metformin (Amini et al., 2005). Serum concentrations of insulin were
determined by RIA kit (Insulin Coat-A-Count; Diagnostics Products Corporation; Los Angeles, CA). Signs of estrus were monitored daily for 2 weeks after treatment.

**Statistical Analysis**

Normalized protein concentrations ratios were tested for normal distribution via the Shapiro-Wilk test (Shapiro and Wilk, 1965). A one-tailed Student’s t-test was used for comparison of data sets that had a normal distribution. For data sets that did not have a normal distribution, a one-tailed Wilcoxon two-group test (Mann-Whitney test) was used. Concentrations of serum P4 was examined using ANOVA with repeated measures, and serum concentrations of insulin, as well as luteal P4 content, was examined using ANOVA with Dunnett’s Method post hoc test. Data were analyzed using JMP and SAS software (JMP®, Version Pro 11, SAS Institute Inc., Cary, NC, Copyright ©2013; SAS®, Version 9.3, SAS Institute Inc., Cary, NC, Copyright ©2002-2010). Significance criterion alpha for all tests was 0.05. Data are depicted as the mean ± SEM.

**Results**

**Experiment 1**

A decrease in serum P4 concentration and luteal P4 content at 2 h (2.08 ng/ml vs. 2.96 ng/ml; \( P = 0.054 \)) and 4 h (1.57 ng/ml vs. 2.96 ng/ml; \( P = 0.013 \)) was observed after PGF\(_{2\alpha}\) administration (d10-h0 = 5.05 ng/ml; d10-h2 = 2.03 ng/ml; d10-h4 = 1.74 ng/ml) (Figure 1.1). Changes in protein expression of LDL, SRB-1, AMPK, StAR, and ACAT-1 are shown in Figure 1.2. There was no detection of P-AMPK. A decrease in LDL protein expression was observed at 2 h (0.79 vs. 0.41; \( P = 0.09 \)) and at 4 h after PGF\(_{2\alpha}\) injection (0.79 vs. 0.13; \( P = 0.004 \)) (Figure 1.2-A). An increase in ACAT-1 occurred by 4 h (0.67 vs. 0.21, \( P = 0.010 \)) after PGF injection (Figure 1.2-B). There was a significant quadratic effect of treatment and time (\( P = 0.037 \)) for
AMPK protein expression (Figure 1.2-C). An increase in StAR protein expression was observed 4 h (3.62 vs. 1.25, $P = 0.01$) after PGF$_{2\alpha}$ injection, but not at 2h (1.02 vs. 1.74, Figure 1.2-D). There was also a quadratic effect of treatment and time ($P = 0.05$) for StAR protein expression, with H4 having the highest ratio (Figure 1.2-D). No difference in SRB-1 protein expression occurred at either 2 (1.92 vs. 1.74) or 4 h (2.75 vs. 1.74) after PGF injection (Figure 1.2-E).

**Experiment 2**

The preliminary experiment revealed that even though concentrations of metformin were detectable in blood after intrauterine administration, which is shown by the representative HPLC metformin determination from a 30-mg/kg bw cow in Figure 1.3; intravenous administration was simpler. There was a decrease in serum concentrations of P4 ($P = 0.001$) with 30 mg (1.01 ± 0.08 ng/ml) of metformin compared to 15 mg (1.33 ± 0.08 ng/ml, Figure 1.3). However, days to estrus was not different between the groups (x = 21 d ± 1). Figure 1.4 illustrates that in the there was no difference in P4 between metformin (0.784 ± 0.15 ng/ml) and saline (0.788 ± 0.29 ng/ml). Additionally, there was no difference in insulin concentrations between saline (4.49 ± 0.154 µg/L) and metformin (4.48 ± 0.154 µg/L) treatments ($P > 0.05$; Figure 1.5), or time to return to estrus (x = 11 d ± 1).

**Discussion**

These results corroborate that activation of the FP receptor by exogenous PGF$_{2\alpha}$ causes a decrease in serum and luteal P4 concentrations (Juengel et al., 1993; Wiltbank et al., 1995; Tsai and Wiltbank 1998; Atli et al., 2012). Similarly, both serum concentrations and luteal P4 content decreased by 4 hours after PGF$_{2\alpha}$ administration. These decreases of both serum and luteal P4 were observed as early as 2 h after PGF$_{2\alpha}$, yet no changes in serum P4 were observed at 1 h after PGF$_{2\alpha}$ (data not shown). These data established a time frame for the process of functional
regression after exogenous injection of PGF$_{2\alpha}$.

During this functional regression, cholesterol transport through LDLR was lower at 2 and 4 h after PGF$_{2\alpha}$, indicating that substrate transport by these receptors was critical during reduction of P4 production. Lower LDLR would lead to decreased cholesterol, the substrate for P4 production. Decreases in LDL receptors reduce 3-hydroxy-3-methylglutaryl coenzyme A reductase (HMG-CoA reductase), catalyzing a rate-limiting step in cholesterol production (Siperstein, 1970), and increasing ACAT-1 (Brown et al., 1975). Cells adjust to the number of LDL receptors in order to produce sufficient cholesterol for the metabolic need of the cell without over accumulation (Brown and Goldstein, 1975). This function is key because cells are able to keep the concentration of unesterified cholesterol in membranes constant, while requirements and exogenous supply are fluctuating constantly (Goldstein and Brown, 2009). The observed increases of ACAT-1 indicate an attempt by the luteal tissue to compensate for decreases in cholesterol.

No change in SRB-1 in mature CL was observed after PGF$_{2\alpha}$ administration. During spontaneous luteolysis in primate CL, changes in gene expression or protein concentrations of SRB-1 were not different between mid-late to functional late CL but were decreased from functional to functionally regressed late stage CL (Bogan et al., 2009, 2010). Functional late CL were defined as having been collected on day 14-16 of the menstrual cycle, and functionally regressed late CL were a subset of those CL from animals with serum P4 values less than 0.5 ng/ml (Bogan et al., 2009). Based on the criteria used by Bogan et al. (2009, 2010), the time points used in the current study after PGF$_{2\alpha}$ was given, could have reflected functional regression; whereas, the results reported by Bogan et al. reflected structural regression.

Expression of AMPK mRNA has been measured in whole bovine ovaries, small and
large follicles, oocytes, and CL. Additionally, AMPK-α subunit protein expression was detected in small and large follicles and their granulosal cells (Tosca et al., 2007). Previous data indicated a developmental difference in AMPK gene expression between developing and mature CL, with increased expression in mature bovine CL (Goravanhally, unpublished results). Therefore activation of AMPK would have a greater effect in the mature than developing CL. Activation of AMPK using metformin decreased P4 production in bovine granulosal (Tosca et al., 2007) and luteal cells (Hou et al., 2009). Thus, elevated concentrations of AMPK protein observed 2 h after FP activation may serve as a mechanism by which both serum and luteal P4 concentrations are decreased. At 4 h after FP activation, P4 concentrations decreased such that AMPK activation was not required to mediate further decreases in steroid production. No detection of P-AMPK occurred at either 2 or 4 hours. This may be a result of rapid AMPK phosphorylation. Park et al. (2002) reported AMPK phosphorylation within 5 seconds in skeletal muscle of rats. Therefore, time points chosen after PGF2α may have been too late to observe any change in P-AMPK.

Metformin was delivered successfully into the uterine lumen of the cows, and detection of its presence in serum was achieved using HPLC. This delivery method was chosen originally due to the concept that counter-current exchange in the utero-ovarian vein (McCracken et al., 1999) would aid in delivery of metformin to the CL. Despite the fact that metformin was detectable in blood, passing the AI rod through a cow in the luteal phase was technically difficult. Differences in P4 concentration were observed between 15 and 30 (mg/kg of BW) doses of metformin during the preliminary study. The 30 mg dose was chosen based on studies in horses, in which metformin was being used as treatment for insulin resistance (Hustace et al., 2009; Rendle et al., 2013). This observation combined with the lack of visible negative side
effects of metformin on cows, indicated that a 30 mg/kg of BW dose would be appropriate for IV administration in subsequent experiments. However, at this dose there was no difference in P4 concentrations between the metformin- and saline-treated cows, and no difference in their time to return to estrus. Additionally, no changes in insulin concentrations after metformin were observed. This was surprising because this same dose caused a significant decrease in insulin concentrations in horses (Rendle et al., 2013). A greater dose of metformin might have elicited a response in both insulin and P4 production. This would be difficult to test due to the cost prohibitive nature of administering a drug that is used in the human market to treat patients with PCOS (Tang et al., 2010) and diabetes (Viollet et al, 2012).

In summary, activation of the FP receptor caused a decrease in luteal and serum P4 concentrations as soon as two hours after exogenous administration PGF$_2$α. Alterations in cholesterol transport accounted for this reduction, at least in part, in P4 production. Changes in AMPK protein during this same time period indicate that it may be the downstream target of FP activation, which controls cholesterol trafficking.
Serum progesterone (ng/ml) = 2.9742 - 0.3399*Time from PGF$_{2\alpha}$ (Hour)

Luteal progesterone (ng/ml) = 4.498 - 0.7984*Time from PGF$_{2\alpha}$ (Hour)

Figure 1.1: Bivariate fit of serum (A) and luteal (B) P4 (ng/ml) by time from PGF$_{2\alpha}$ (hour). Significant declines in serum and luteal progesterone occurred at both 2 and 4 hours after PGF$_{2\alpha}$.
Figure 1.2: Semiquantitative analysis of the densitometric data derived from Western blots using protein samples isolated mature bovine CL 0, 2, and 4 hours after PGF$_{2\alpha}$. The y-axis shows the ratio of the optical density (o.d.) for the protein of interest corrected by the detected o.d. for its corresponding actin. The data are given as mean ± SEM; values with differing letters denote statistically significant differences (P < 0.001).
Figure 1.3: A-Serum progesterone values for 15 or 30 (mg/kg of bw) IU of metformin. Serum progesterone was significantly lower 12 hours after starting treatment ($P < 0.001$) when cows received 30 mg/kg bw than when cows received 15 mg/kg bw. B-Representative HPLC graph of metformin concentrations in a cow receiving 30/mg kg of bw.
Figure 1.4: Serum progesterone concentrations pooled across all experiments showed that there was no difference in metformin-treated (n = 7) versus saline-control (n = 7) cows.
Figure 1.5: Serum insulin concentrations one day prior to and during metformin treatment. Cows were given 30 mg/kg bw of metformin IV or sterile saline for control cows every 3 hours over 12 hours, during the day of treatment. No significant difference of treatment or time was seen in insulin concentrations.
Chapter 3: Adenosine monophosphate activated protein kinase activation in vitro modulated progesterone secretion in mature but not developing bovine CL

Introduction

The mechanisms governing the action of prostaglandin F 2 alpha (PGF₂α), are incompletely known. Nevertheless, this mechanism involves activation of the phospholipase C-Ca²⁺ pathway (Davis et al., 1987). In luteal cells, PGF₂α has affects by binding to its cognate G₉ protein-coupled receptor and activating phospholipase C (PLC; Davis et al., 1988). The PLC cleaves plasma membrane-bound phosphatidylinositol 4,5-bisphosphate (PIP₂) to generate free IP3 (Berridge et al., 1983) and membrane-bound 1,2-diacylglycerol (DAG; Berridge, 1987), which increases Ca²⁺ and activates protein kinase C (PKC), respectively. Increases in both concentrations of cytoplasmic Ca²⁺ and activation of PKC have been demonstrated to mediate intracellular actions of PGF₂α in luteal cells (Wiltbank et al., 1990). Differential expression of genes associated with Ca²⁺ homeostasis plays a role in the development of luteolytic capacity of bovine CL. Greater concentrations of PGF₂α are needed to elicit Ca²⁺ responses in steroidogenic cells isolated from developing than from mature CL (Choudary et al., 2005). Additionally, gene expression of CAMKK2, a down-stream target of Ca²⁺, is increased in mature CL (Goravanahally et al., 2009). The serine/threonine protein kinase CAMKK2 has been reported to mediate the action of intracellular Ca²⁺ via phosphorylation of calcium/calmodulin-dependent kinases (CAMKs). Whether CAMKK2 affects steroidogenesis or rather acts upon another intermediary to mediate its effects is not known. One target of CAMKK2 is adenosine monophosphate activated protein kinase AMPK (Wong, 2009). The kinase CAMKK2 phosphorylates AMPK, but this particular component of this signal transduction pathway has not
been investigated in the bovine CL.

In multiple cell types, AMPK is an important regulatory protein for energy balance. In all eukaryotic cells it is a heterotrimeric complex composed of one catalytic subunit α (with two isoforms: PRKAA1 and PRKAA2) and two regulatory subunits β (with two isoforms: PRKAB1 and PRKAB2) and γ (with three isoforms: PRKAG1, PRKAG2, and PRKG3; Wong et al., 2009). Increases in cytoplasmic Ca\(^{2+}\) lead to activation of CAMKK2, which is capable of phosphorylating AMPK at Thr\(^{172}\). Binding of AMP to the γ subunit activates AMPK by inducing conformational changes in the complex that promote phosphorylation of Thr\(^{172}\). Typically, activation of AMPK results in a shift from anabolic processes (Hardie, 2008), such as synthesis of proteins that are required for progesterone (P4) production, to catabolic processes. Recently, activated AMPK decreased P4 secretion in murine, bovine, and primary F1 galline granulosal cells (Tosca et al., 2005; Tosca et al., 2006; Tosca et al., 2007). All subunits of AMPK and their isoforms were present in whole ovaries, CL, small and large follicles as well as granulosal cells in the cow (Tosca et al., 2007). AMPK is activated in response to the antidiabetic drug metformin (Viollet et al., 2012), and by 5-Aminoimidazole-4-carboxamide-1-β-4-ribofuranoside (AICAR), a cell-permeable allosteric AMPK activator (Corton et al., 1995). Dorsomorphin dihydrochloride (DM) is a potent, selective, cell-permeable and reversible AMPK inhibitor (Liu et al., 2014). It inhibited AMPK induced either by AICAR (Tang et al., 2011) or metformin (Isakovic et al., 2007). If CAMKK2 targets AMPK in bovine CL, increased gene expression of CAMKK2 during this developmental stage may lead to increased AMPK activation, and, therefore, become an additional mechanism to mediate effects of PGF\(_{2α}\).

The goal of this project was to investigate alternative downstream components activated by the rise in cytosolic [Ca\(^{2+}\)] initiated when the FP receptor is activated in bovine luteal tissue.
The objectives of this research were to determine the changes in P4 secretion in the presence of AMPK activators or inhibitor in developing and mature bovine CL in vitro. Developmental differences in components of signal transduction associated with AMPK activation in developing and mature CL may explain, at least in part, the developmental differences in the ability of PGF$_{2\alpha}$ to induce functional luteal regression.

Materials and Methods

Animal Handling and Surgical Procedures
Non-lactating beef cows were observed twice daily for estrus at approximately 12 h intervals for 30 min per observation. The day when standing estrus was observed was designated as day 0 (d0). The West Virginia University Animal Care and Use Committee reviewed and approved the protocol for the tissue collection (ACUC # 01-0809). Ovaries (d4; n = 5) or CL (d10; n = 5) were collected via supravaginal incision under epidural anesthesia (Casida, 1959) and transported in ice-cold saline to the laboratory. Connective tissue was removed prior to obtaining weights and slicing for treatment.

Luteal Slice Incubation Procedure
Luteal slices were incubated for 30 min with 500 µl of minimum essential media (MEM, Life Technologies, Grand Island, NY) and pharmaceutical agents before hormones were added. Media were removed, and fresh media added back along with treatments for 2 hr incubation at 37°C, shaking at 200 rpm. Progesterone was assayed by radioimmunoassay (RIA) as previously described (Sheffel et al., 1982).

Experiment 1
Luteal slices were treated with MEM (control), PGF$_2\alpha$ (1.0 µg/ml), DM (10.9 µM/L, Tocris, Bristol, UK), or DM (10.9 µM/L) and PGF$_{2\alpha}$ (1.0 µg/ml). Media were collected and frozen at -
20°C until assayed for P4.

**Experiment 2**

Luteal slices were treated with MEM (control), PGF$_{2\alpha}$ (PGF$_{2\alpha}$, 1.0 μg/ml), metformin (Met; 10, 5, 1, 0.5, 0.1 mM; New England Mail Order Pharmacy, Middlebury, VT), or AICAR (7.5, 0.75, 0.075 mM, Tocris, Bristol, UK). After incubation, slices were homogenized using an Omni Tissue Homogenizer (Omni International, Kennesaw, GA) in incubation media and frozen at -20°C until assayed for P4.

**Statistical Analysis**

Progesterone values were analyzed using ANOVA followed by Dunnett’s Method for means comparisons to control. T-Test was used to compare the value of progesterone at the highest dose of metformin and AICAR with control values in both D4 and D10. Data were analyzed using JMP and SAS software (JMP®, Version Pro 11, SAS Institute Inc., Cary, NC, Copyright ©2013; SAS®, Version 9.3, SAS Institute Inc., Cary, NC, Copyright ©2002-2010). Significance criterion alpha for all tests was 0.05. Data are depicted as the mean ± SEM.

**Results**

**Experiment 1**

No significant effects of PGF2α (0.94 ± 0.13; $P = 0.96$), DM (1.921 ± 1.3, $P = 0.39$), or PGF$_{2\alpha}$ and DM (1.11 ± 0.53; $P = 0.91$) were observed on P4 production when compared to basal values (1.00 ± 0) in d4 CL. In d10 CL, PGF$_{2\alpha}$ increased P4 production (2.4 ± 0.35 vs 1.00 ± 0; $P = 0.0003$), while DM treatment alone (1.34 ± 0.11 vs. 1.00 ± 0) had no effect ($P = 0.24$).

Stimulatory effect of PGF$_{2\alpha}$ was eliminated when DM was added, such that P4 values did not differ from controls (1.85 ± 0.54 vs 1.00 ± 0; $P = 0.31$) but were different from PGF$_{2\alpha}$ alone (1.34 ± 0.11 vs 2.421 ± 0.35; $P = 0.0037$).
Experiment 2

No significant effect of metformin ($P = 0.91$) or AICAR ($P = 0.51$) at any concentration was observed in D4 CL. A decrease in basal P4 was observed with both metformin (0.39 ± 0.20; 10 mM; $P = 0.006$) and AICAR (0.40 ± 0.15; 7.5 mM; $P = 0.0117$) in D10 CL when compared to control (0.99 ± 0.32).

Discussion

These results indicate that AMPK is involved in the signal transduction mechanism of the mature but not developing bovine CL. No significant changes were observed in P4 production when d4 luteal tissue was treated with DM, metformin, or AICAR. However, d10 CL responded to all AMPK activators with a significant inhibition in P4 production, and the AMPK inhibitor eliminated this effect. All subunits of AMPK are present in the bovine ovary (Tosca et al., 2007), but gene expression of these subunits might change throughout development of the CL. In fact, Goravanahally (unpublished results) demonstrated that, with the exception of AMPKγ1 and AMPKγ2, mRNA expression of AMPK subunits were increased in the mature CL ($P < 0.05$). Moreover, an additional observation supporting the interpretation that the effects of increases in cytosolic calcium are conveyed via CAMKK2-AMPK was that a CAMKK2 inhibitor, STO-609, blocked the inhibitory effects of PGF$_{2\alpha}$ in dispersed mature bovine luteal cells (Goravanahally, unpublished results). STO-609 also inhibited PGF$_{2\alpha}$-induced phosphorylated state of AMPK (P-AMPK) in mature but not developing bovine CL. Collectively, these data demonstrate that the signal transduction pathway involving CAMKK2 and AMPK regulates luteal function in the mature, but not the developing CL.

Tosca et al. (2005, 2006, 2007) have provided evidence in the rat, hen, and cow that
AMPK is involved in steroid production in granulosal cells. It is not surprising that AMPK plays a role in steroidogenesis of luteal cells, as this population is derived from granulosal cells. In agreement with previous results, these data reveal that metformin and AICAR decreased basal P4 production. The effects of DM were developmentally different; DM had no effect on P4 production in developing CL, but blocked the stimulatory effects of PGF$_{2\alpha}$ in mature CL. However, lower doses of metformin (5, 1, 0.5, 0.1 mM) or AICAR (0.75, 0.075 mM) did not affect P4 production. Doses of both AMPK activators were based on previous reports, in which a significant decline in basal P4 production was seen *in vitro* (Tosca et al., 2005; Tosca et al., 2007). However, CL tissue from this study was incubated for 2 hours before collection, compared to the range of 3 to 48 hours in the previous study with bovine granulosal cells (Tosca et al., 2007). Differences in P4 were significant at 2 hours of incubation, but longer incubation times with lower doses may have decreased steroid production.

In conclusion, DM eliminated the effect of PGF$_{2\alpha}$ on mature but not developing bovine CL. Higher doses of metformin and AICAR were capable of decreasing P4 in mature but not in developing bovine CL. This developmental difference in signal transduction may play a significant role in the ability of mature CL to regress in response to a single luteolytic dose of PGF$_{2\alpha}$. 

49
Figure 2.1-DM reduced the stimulatory effect of PGF$_{2\alpha}$ on P4 secretion in mature (D10) but neither PGF$_{2\alpha}$ nor DM alone or in combination with PGF$_{2\alpha}$ or LH were affected in the developing CL (D4). Luteal slices were incubated for 30 minutes in MEM and DM without hormones after which all media were removed, and fresh MEM, DM and hormones were added back. Media were collected and frozen until analyzed for progesterone with RIA. Different letters indicate significant differences at (P < 0.05).
Figure 2.2: Neither PGF$_{2\alpha}$ nor its activators, metformin and AICAR, had any significant effect on basal progesterone production in developing (D4) bovine CL. Luteal slices were incubated for 30 minutes in MEM and metformin or AICAR without hormones after which all media were removed, and fresh MEM, metformin or AICAR and hormones were added back. Slices were homogenized in the media and then collected and frozen until analyzed for progesterone with RIA. Different letters indicate significant differences at (P < 0.05).
Figure 2.3: At the greatest concentrations tested, both metformin and AICAR significantly (P < 0.05) decreased basal progesterone production in mature (D10) bovine CL. Luteal slices were incubated for 30 minutes in MEM and metformin or AICAR without hormones after which all media were removed, and fresh MEM, metformin or AICAR and hormones were added back. Slices were homogenized in the media and then collected and frozen until analyzed for progesterone with RIA. Different letters indicate significant differences at (P < 0.05).
Chapter 4: Primary infertility affects progesterone production by luteinized human mural granulosal cells in vitro

Introduction

Follicles, comprised of the oocyte, granulosal and thecal cells, are the functional unit of the ovary. The somatic cells in the outermost vascular portion of the follicle include thecal and myoepithelial cells. The avascular component of the follicle is delimited by a basement membrane upon which the other somatic cells of the follicle, the granulosal epithelial layer and the germ cell reside. Two populations of granulosal cells have been identified: mural and cumulus granulosal cells. Both populations of cells are organized as a stratified epithelium with the basal layer of mural cells contacting the basement membrane adjacent to the thecal cells and the cumulus cells surrounding the oocyte (Buccione et al., 1990). Mural granulosal cells express receptors for FSH and LH as well as the enzymes 3β-HSD and P450scc (Zoller and Weisz, 1979). Expression of these two enzymes is representative of a luteal characteristic in these cells, while cumulus granulosal cells remain with the oocyte even after ovulation (Russell and Salustri, 2006). Once ovulation has occurred, mural granulosal cells lose their ability to proliferate, undergo epithelial-mesenchymal transition and differentiate into large luteal cells (LLC) of the corpus luteum (CL), which secrete high amounts of basal progesterone (P4; Chaffkin et al., 1992). Interestingly, P₄ receptors (PGR) cannot be detected with immunohistochemical analysis in human granulosal cells at any follicular stage but they are detected readily in human luteal tissue (Suzuki et al., 1994). The fact that PGR is absent in preovulatory follicles but present in periovulatory follicles and in the CL after a surge of gonadotropins supports the hypothesis that LH induces PGR expression (Hild-Petito et al., 1988). Thecal cells become small luteal cells (SLC) once luteinization has occurred.
In the CL of primates, granulosal and thecal cells remain as two distinct populations and are separated by the remnants of the follicular basement membrane (Guraya, 1971). In contrast, the basement membrane separating these two cell populations is remodeled, and thecal cells, which become SLC, invade the granulosal cells to form a heterogeneous mixture of cells in the CL of nonprimate species (Niswender, 2000). Rescue of the CL during the fertile cycle by human chorionic gonadotropin (hCG) is essential for maintenance of pregnancy once the oocyte is fertilized. Infertility is a problem that affects millions of women and their partners across the United States. Causes of infertility range from ovulatory dysfunction to male related infertility due to abnormal sperm motility or morphology. Insufficient luteal rescue can account for pregnancy loss in couples conceiving naturally as well as those that have undergone time consuming and expensive procedures such as intrauterine insemination or in vitro embryo production.

Human granulosal cells that have been recovered during oocyte retrieval in patients undergoing in vitro fertilization spontaneously luteinize in culture, thus these cells have been used as a model to study granulosal-lutein cells of the CL (Stewart and Vandevooort, 1997). The degree of luteinization of these cells has not been clearly defined, and studies using them do not take into account the heterogeneity of a granulosal cell population or the fertility status of the patient from whom they were isolated. Using patient cause of infertility as an experimental variable offers a unique opportunity to study how infertility affects P4 production by luteinized granulosal cells when subjected to hormonal treatments.

In multiple species, PGF$_{2\alpha}$ elicits both inhibitory and stimulatory P4 responses in luteal cells in vitro. Most investigators use a non-physiological dose of PGF$_{2\alpha}$ when treating luteal cells in vitro. The fact that PGF$_{2\alpha}$ at high concentrations is able to activate the PGE$_2$ receptor
could, in part, explain increases in P4 production (Rao, 1974). Additionally, the entire population of cells that would normally be present in addition to luteinized granulosal cells is not present in all in vitro experiments. Inhibitory action of PGF$_{2\alpha}$ on human luteinized granulosal cells may be enhanced by presence of endothelial and/or immune cell types (Liptak et al., 2005).

In multiple cell types, AMPK is an important regulatory protein for energy balance. Typically, activation of AMPK results in a shift from anabolic processes (Hardie, 2008), such as synthesis of proteins that are required for progesterone production, to catabolic processes. The antidiabetic drug metformin activates AMPK (Viollet et al., 2012), and by 5-Aminomidazole-4-carboxamide-1-β-4-ribofuranoside (AICAR), a cell-permeable allosteric AMPK activator (Corton et al., 1995). Dorsomorphin dihydrochloride (DM) is a potent, selective, cell-permeable and reversible AMPK inhibitor (Liu et al., 2014), and DM inhibited AMPK induced either by AICAR (Tang et al., 2011) or metformin (Isakovic et al., 2007). Metformin is the most widely used drug for the treatment of type II diabetes, being prescribed to nearly 120 million people worldwide (Viollet et al., 2012). Recently, metformin was used to treat women experiencing PCOS who were overweight or insulin resistant. Metformin’s ability to increase insulin sensitivity in individuals made it an ideal candidate for the treatment of PCOS. Metformin increased insulin sensitivity in patients with PCOS, and it increased ovulation rate, improved menstrual cyclicity, and reduced serum androgen (Tang et al., 2010). Metformin appears to work in PCOS patients by both direct ovarian effects and secondarily through decreasing insulin resistance. In bovine granulosal cells isolated from small antral follicles metformin has been shown to have effects on steroidogenesis (Tosca et al., 2007). Metformin (10 mM) reduced both progesterone and estradiol production as well as key enzymes (HSD3β, CYP11A1, and STAR) with or without FSH and/or IGF1 in the media (Tosca et al. 2007).
The goal of this project was to investigate the use of human mural granulosal and cumulus cells to study potential downstream targets activated by the known intracellular mediators, AMPK and Ca\(^{2+}\), of luteolytic actions of PGF\(_{2\alpha}\) actions in bovine luteal cells. Additionally, it was hypothesized that P4 stimulatory and inhibitory responses to hCG and PGF\(_{2\alpha}\), respectively, would differ among infertility types. The objectives of this research were to define culture conditions and responses of luteinized human granulosal cells derived from patients with various infertility causes to luteotropic and luteolytic signals and to determine effects of increasing intracellular Ca\(^{2+}\) from low to high values on P4 production. The proposed research had one central hypothesis: luteinized mural granulosal cells can be used to study how infertility influences luteal efficiency and the effects of increasing intracellular Ca\(^{2+}\) on P4 production in order to determine if providing the known intracellular mediator of the best known luteolytic factor, PGF\(_{2\alpha}\), could directly induce a reduction of P4, which is the hallmark event underlying luteal regression.

**Materials and Methods**

**Patients**

Mural and cumulus granulosal cells from patients affected by different causes of infertility were provided by the WVU Center for Reproductive Medicine in Morgantown, WV after patient’s ovaries were stimulated for transvaginal oocyte collection for *in vitro* fertilization (IVF). Patients ranged in age from 25-40 years of age, had a BMI range of 18.9 to 44.3, and presented with primary infertility types of male factor (male, n = 5), endometriosis (endo, n = 6), tubal factor (tub, n = 12), and unexplained/other (other, n = 3). Daily or twice daily injections of recombinant follicle stimulating hormone (FSH) began on d2 or d3 of the cycle and continued until ovulation was induced with hCG. Patients returned at least every other day for ultrasonic
monitoring of follicular growth beginning four days after stimulation. To prevent premature ovulation, GnRH agonist injections began when follicles reached at least 14 mm in diameter. Ovulation was induced with hCG injection once follicles were \( \geq 18 \text{ mm in diameter} \), followed by oocyte collection 34 h later.

**Isolation and culture of primary human GCs**

Follicular aspirates from each patient were centrifuged at 10,000 rpm for 5 min at room temperature. Supernatant was discarded, and the pellet was resuspended in 500 µl of DMEM/F12 (Invitrogen, Grand Island, NY) with 20% (v/v) fetal bovine serum, insulin/transferrin/selenium B (1000; 555; 0.67 mg/L respectively; Gibco, Grand Island, NY), gentamicin/amphotericin (10 µg/ml-0.25µg/ml; Gibco, Grand Island, NY), and penicillin/streptomycin (100 I.U./ml, 100 µg/ml; Life Technologies, Grand Island, NY). Viable cell number was determined using trypan blue dye exclusion and hemocytometer. Cells were plated at 10,000 cells per well in 96-well flat bottom culture plates with 300 µl DMEM/F12 media (NUNC, Scientific Laboratory Supplies, Wilford, Nottingham, UK) at 37°C and 5% CO\(_2\) in a humidified incubator to permit luteinization and cell attachment. Media was changed every 24 h whereby150 µl of spent media was replaced with fresh DMEM/F12 with serum. Four days after collection, cells were transferred to serum free DMEM/F12 media, and treatments were administered. Every treatment was tested in triplicate for each patient and media pooled for P4 by radioimmunoassay (RIA) as previously described (Sheffel et al., 1982).

**Experiment 1**

Cells were treated with PGF\(_2\alpha\) (0.1 µg/ml), human hCG (5 IU/ml), or a combination of both PGF\(_2\alpha\) and hCG. Cells were cultured further for 24 h after which medium was removed and frozen at -20°C until assayed for P4. Primary cause of infertility (male factor, endometriosis,
tubal factor, and unknown), age (< 30, 30-35, 36-40 years of age), BMI (normal = 18.5-24.9, overweight = 25-29.9, obese > 30), and pregnancy outcome (non-pregnant, and pregnant) were examined separately due to the variability among patients in these categories.

**Experiment 2**

Cells were treated with an intracellular calcium buffer 1,2-bis (o-aminophenoxy) ethane-N,N,N',N'-tetraacetic acid, BAPTA (500, 50, 5, 0.5 µmol, Tocris, Bristol, UK) or a calcium ionophore A23187 (1, 0.1, 0.01, 0.001, 0.0001, 0.00001, 0.000001 µmol, Tocris, Bristol, UK) to determine effects of altering intracellular Ca^{2+} on P4 production in luteinized human granulosal cells. Treatment concentrations of BAPTA and A23187 and incubation times were based on previous studies with bovine luteal steroidogenic cells and preliminary studies testing 1, 3, and 24 h incubation times in mural granulosal cells, respectively. The medium for the control group contained 0.1% dimethylsulfoxide (DMSO, Pierce Rockport, IL), the solvent used for BAPTA and A23187. Cells were treated with BAPTA or A23187 for 1 hour, and medium was removed and frozen at -20°C until assayed for P4.

**Experiment 3**

Under the same culture conditions, mural granulosal cells were treated with DM (109, 10.9, 1.09, 0.109 µm/L, Tocris, Bristol, UK) or AICAR (7.5, 0.75, 0.075, 0.0075 mM, Tocris, Bristol, UK) to evaluate the roles of inhibiting/stimulating AMPK, respectively, in luteinized mural granulosal cell P4 production. Cells were cultured for further 24 h and medium was removed and frozen at -20°C until assayed for P4.

**Experiment 4**

Due to stimulation of P4 that was observed when cells were treated with PGF_{2α} it was hypothesized that this increase was due to promiscuous binding to another PG receptor, such as
PGE₂, which is known to increase P₄. To test this hypothesis, mural granulosal cells were cultured with media alone (control), PGF₂α, PGF₂α antagonist (AL 8810; 1.0 µmol, Cayman Chemical, Ann Arbor, MI), and PGF₂α and AL 8810. Preliminary dose response studies for AL 8810 at 10, 1, and 0.1 µmol were conducted in order to find an optimum dose for experiment 4. The medium for the control group contained 0.1% dimethylsulfoxide (DMSO; Pierce, Rockport, IL), the solvent used for AL 8810. Cells were preincubated with AL 8810 for 30 min, treatments were administered and cells cultured for 24 hours. Medium was collected and frozen at -20°C until assayed for P₄. Additionally, the effect of PGF₂α and hCG on P₄ production in cumulus cells was tested.

**Statistical Analysis**

Concentrations of P₄ were tested for normal distribution via the Shapiro-Wilk test. All data were transformed using natural logarithm. The one-tailed Student t-test was used to examine P₄ production in A23187 experiments. All other experiments were analyzed using ANOVA and Dunnett’s post hoc test. Data were analyzed using JMP and SAS software (JMP®, Version Pro 11, SAS Institute Inc., Cary, NC, Copyright ©2013; SAS®, Version 9.3, SAS Institute Inc., Cary, NC, Copyright ©2002-2010). Differences were considered significant when \( P < 0.05 \). Data in all graphs are depicted as the mean ± SEM.

**Results**

**Experiment 1**

There was an increase in P₄ production in cells treated with hCG who had been collected from patients that presented with male infertility factor compared to control (\( P = 0.0045 \); Figure 3.1-A), but not in patients who presented with tubal, endometriosis, or diminished ovarian reserve factor (\( P > 0.05 \); Figure 3.1-B,C,D). No significant differences in P₄ were seen when
granulosal cells were treated with PGF$_{2\alpha}$ across all infertility types ($P > 0.05$). Additionally, hCG elicited a significant increase in P4 production in patients in the age range of 30-35 (n = 15; $P = 0.008$) and 36-40 (n = 6; $P = 0.04$), but not in patients in the ranges of 25-30 (n = 4; $P = 0.73$) (Figure 3.2). Patients with normal BMI (18.5-24.9, n = 16) had a significant increase in P4 production when treated with hCG ($P < 0.0001$), but granulosal cells of patients who were categorized as overweight (BMI of 25-29.9, n = 6) or obese (BMI >30, n = 3) did not have a significant increase in P4 production when stimulated with hCG ($P > 0.05$) (Figure 3.3). Figure 3.4 demonstrates that granulosal cells from patients who became pregnant to the IVF cycle had significantly greater P4 production when stimulated with hCG (3.23 ng/ml; $P = 0.0018$ vs. control) than those who did not become pregnant (1.57 ng/ml; $P > 0.05$ vs. control).

**Experiment 2**

A significant reduction in P4 production was found with all concentrations of AICAR except the lowest (0.0075) when compared to control ($P < 0.001$). However, no difference was observed when cells were treated with DM at any concentration ($P > 0.05$).

**Experiment 3**

There was a trend ($P = 0.062$) for PGF$_{2\alpha}$ treatment to increase P4 production in mural granulosal cells (Figure 3.6-A). The PGF$_{2\alpha}$ receptor antagonist, AL 8810, did not block this stimulation. However, there was a significant difference between controls and PGF$_{2\alpha}$ and AL 8810 in combination ($P = 0.024$). Cumulus cells responded to hCG stimulation with a significant increase in P4 production ($P = 0.045$), but no difference in P4 was observed when cumulus cells were treated with PGF$_{2\alpha}$ or PGF$_{2\alpha}$ and the PG antagonist (Figure 3.6-B).

**Experiment 4**

Progesterone production was increased in mural granulosal cells by A23187 at 0.01 µmol
(P = 0.041), and a trend (P = 0.096) was observed at 0.1 and 1.0 µmol concentrations tested in comparison to control concentrations. There was a cubic trend for the effect of BAPTA concentration on P4 production (P = 0.093) in mural granulosal cells.

Discussion

Data from experiment 1 supported the conclusion that types of infertility affect the ability of luteinized mural granulosal cells to respond to the luteotropic factor, hCG, in vitro. Knowing the cause of a patient’s infertility prior to cell culture experiments offers a unique model to study how infertility affects P4 production by luteinized granulosal cells subjected to hormonal treatments. Failure of these cells to respond to hCG in vivo could lead to luteal insufficiency or even early regression of the corpus luteum, resulting in pregnancy loss. Implementing different post-implantation hormonal treatments based on specific causes of infertility could lead to increased pregnancy rates while keeping the number of multiple gestations low.

Interestingly, mural cells of women who did not become pregnant had significantly lower P4 production when stimulated with hCG, when compared to the mural cells of pregnant women. Failure of these cells to secrete high levels of P4 in vivo is similar to a condition known as luteal phase defect (LPD). In LPD is a brief elevation in P4 secretion (typically < 11 days) after ovulation, and is reported to occur in up to 20 percent of women (Balasch and Vanrell, 1987). This short luteal phase may be due a defect in the process of luteinization or progesterone production. However, it was postulated by Strott et al. (1970) that luteal phase defect occurs as a result of low preovulatory FSH levels. This leads to abnormal follicular development and subsequent inadequate luteinization and/or luteal function. This condition occurs despite normal LH stimulation at the time of the ovulatory surge (Strott et al., 1970). This could be the case in the cells from patients who did not become pregnant, where inadequate FSH levels resulted in
the mural granulosal cells being unable to respond to hCG. These experiments were performed four days after oocyte retrieval, which is later than when embryos are normally transferred at the WVU CRM. It would be interesting to stimulate cells sooner to determine if they could be used as a predicator of pregnancy success prior to embryo implantation. If these cells could be used as a diagnostic tool, it could save patients and embryologists substantial time and monetary resources.

It was not surprising that P4 production by mural granulosal cells was affected by patient age. The ability to conceive naturally, which decreases as women age; naturally decreases by 11% from 30-34, and another 12% from ages 35-40 (Leridon, 2004). Decreased fertility may be explained partially by luteal insufficiency. The fact that luteinized mural granulosal cells did not respond to hCG with increases in P4 leads to the implication that luteal cells derived from these follicles would not respond sufficiently. Surprisingly, the age group that did not have an increase in P4 production was the youngest age group. Types of primary infertility were represented well in this group, but the small population size (n = 4) may explain this anomaly. In contrast, it was encouraging to see that mural cells from older patients responded to luteotropic hCG.

Patients with BMIs in the overweight or obese range did not respond to hCG with increased P4 production. The interrelationship between nutritional status and reproduction is complicated, and the mechanisms connecting the two are not well defined. However, insulin, glucose, IGF-1, and leptin all impinge on follicular development directly or influence gonadotropins at the hypothalamic level. As BMI and adiposity increase so do leptin, and insulin resistance. It will be critical to ascertain if the mural cells of overweight and obese patients have insufficient substrate to produce P4 when stimulated or if they are resistant despite
adequate substrate. Women with PCOS are often obese as well as insulin resistant, and are prescribed insulin-sensitizing medicines such as metformin (Tang et al., 2010).

Metformin works, in part, by activating AMPK, a key regulator of energy homeostasis that has direct effects on P4 production in bovine (Tosca et al., 2007), galline (Tosca et al., 2007) and rodent (Tosca et al., 2005) granulosal cells. Using another AMPK activator, AICAR, a significant decrease in P4 was seen at all concentrations tested. A decrease in P4 production has been shown in mature but not developing bovine CL, as mentioned earlier in this dissertation. This decrease in P4 could be explained by a decrease in key enzymes such as 3βHSD, P450scc, or StAR, all of which decreased in rat granulosal cells (Tosca et al., 2005). In contrast, no difference in P4 production was seen when mural cells were treated with an AMPK inhibitor, DM. The concentrations of DM chosen for experiment 2 were based on studies in bovine CL, in which DM significantly decreased P4 production. Possibly, a greater concentration of DM was needed to elicit a change in P4 production in human mural granulosal cells.

A stimulatory effect of PGF$_{2\alpha}$ has been observed consistently over time and has been attributed by some to the use of non-physiological doses of PGF$_{2\alpha}$ in vitro, which activate other prostaglandin receptors (Rao et al., 1974). Progesterone stimulation was observed in mural granulosal cells but not in cumulus granulosal cells (Figure 6). A specific PGF$_{2\alpha}$ receptor antagonist was unable to eliminate this effect in mural cells, indicating that the stimulation of P4 production was mediated by another prostaglandin receptor. Possibly, PGF$_{2\alpha}$ is binding to PGE$_2$ receptors and increasing cAMP, which leads to an increase in P4. The ability of cumulus cells to respond to hCG with an increased P4 indicates that, even after ovulation, hCG is capable of signaling the oocyte. Human cumulus oocyte complexes express LH receptors (LHR) and up-regulate LHR expression during the ovulatory process even though LHR expression in cumulus
cells is much lower than in mural granulosal cells (Jeppesen et al., 2012).

It is commonly accepted that increases in intracellular Ca\(^{+2}\) in response to FP activation cause a decrease in P4 production in luteal cells. A quadratic effect on P4 production was seen when the calcium buffering capacity of the cells was increased by BAPTA. Increases in intracellular Ca\(^{+2}\) were needed to support P4 production in bovine luteal cells (Alila et al., 1990). Concentrations of BAPTA at 5 and 50 µmol were needed before a decrease in P4 production was observed in experiment 4 (Figure 7). A threshold of intracellular Ca\(^{+2}\), even beyond the highest concentration in experiment 4, may be needed to decrease P4 production. Lower concentrations of BAPTA had either no effect or stimulated P4 production (Figure 7). This result supports a biphasic role for Ca\(^{+2}\) in P4 production in human mural granulosal cells. By pharmacologically increasing intracellular Ca\(^{+2}\) with an ionophore, A23187, basal P4 production was increased. This is contradictory to the findings of Goravanahally et al. (2009), in which A23187 did not affect basal P4 production in cultured bovine luteal cells. Differences between species or the stage of luteinization could explain this difference. Luteinized granulosal cells and luteal cells share several characteristics; however, they are not identical.

In summary, human mural granulosal cells can serve as a valuable model for understanding how increases in intracellular Ca\(^{+2}\) and its downstream targets affect P4 production in steroidogenic cells. Additionally, P4 production by mural granulosal cells differed among primary infertility type, age, BMI, and pregnancy outcome and may be used to study each of these factors individually. Understanding similarities and differences between previously mentioned factors has the potential to improve IVF protocols and to decrease the incidence of luteal-insufficiency-induced miscarriages.
Figure 3.1: Progesterone production by granulosal cells from patients with male factor (A), endometriosis (B), diminished ovarian reserve (C), and tubal (D) primary infertility.
Figure 3.2: Progesterone production in mural granulosomal cells was affected by age. Cells from patients who ranged in age from 25-30 (n=4) did not respond to hCG with a significant increase in P4 production (A). However, cells from other age groups (31-35, n = 15 and 36-40, n = 6) had significant increases in P4 when stimulated with hCG (ages 31-35 and ages 36-40 in B and C, respectively).
Figure 3.3: Progesterone production in mural granulosal cells was affected by BMI. Cells from patients with (A) normal BMI (18.5-24.9, n = 16) had a significant increase in P4 production when treated with hCG (P < 0.0001). Cells of patients who were categorized as (B) overweight (BMI of 25-29.9, n = 6) or (C) obese (BMI >30, n = 3) did not have an increase in P4 production when stimulated with hCG (P > 0.05).
Figure 3.4: Progesterone production in mural granulosal cells was affected by pregnancy status post IVF. Luteinized mural granulosal cells from patients who became pregnant (B) to the IVF cycle produced significantly more P4 when stimulated with hCG than cells from those who did not become pregnant (A) (3.23 vs. 1.57 ng/ml; P < 0.05).
Figure 3.5: The AMPK activator, AICAR, inhibited P4 production by human luteinized mural granulosal cells at all dosages tested (A). In B, the AMPK inhibitor, DM, did not affect basal P4 production.
Figure 3.6: PGF$_{2\alpha}$ stimulated P4 production in mural granulosal cells (A) but not in cumulus granulosal cells (B). Adding a PG receptor antagonist, AL 8810, did not block stimulation in mural cells and had no effect in cumulus cells (A). P4 production in cumulus cells was stimulated by hCG, shown in B.
Figure 3.7: BAPTA had a biphasic effect on P4 production in luteinized mural granulosal cells. Data are presented as natural log of the ratio of BAPTA-treated to control values.

\[
\ln (P4/C) = 0.2802551 + 0.0045803 \times \ln (\text{BAPTA concentration}+0.0001) - 0.0097173 \times (\ln (\text{BAPTA concentration}+0.0001) - 3.589)^2
\]
Figure 3.8: Increases in intracellular calcium increased P4 production in mural granulosal cells. A23187 increased P4 concentrations at 0.01, 0.1, and 1μmol concentrations (A, B, and C respectively).
Chapter 5: Literature Cited


Pate, J.L. (1988a) Regulation of prostaglandin synthesis by progesterone in the bovine corpus luteum. *Prostaglandins* **36** (3), 303-315


