1. Introduction

1.1 Progesterone synthesis

Progesterone was independently discovered by different research groups (Butenandt & Westphal 1934, Allen 1935). Allen and collaborators discovered progesterone in 1933, and were the first to determine the molecular weight and partial molecular structure. The name progesterone derives from progestational steroidal ketone (Allen 1935).

Fig. 1. Steroid hormone synthesis. The precursor cholesterol from the maternal circulation is converted to 21 carbon (C21) progestagens. Progestagens can be converted to C21 glucocorticoids, or to C19 androgens. Androgens serve as precursors for C18 estrogens. Source: Wikipedia.
Progesterone belongs to the C21 group of progestagens and is the evolutionary most conserved of the reproductive steroid hormones. The synthesis of progesterone from its precursor cholesterol in the maternal circulation requires only two enzymatic steps to form pregnenolone, which is readily transformed to progesterone. The main source of progesterone in humans is the corpus luteum in the ovary. After conception, the corpus luteum is supported by the secretion of human chorionic gonadotropin (hCG) from the conceptus, and produces progesterone until approximately the 10th gestational week. After a transition period by 7-10 gestational weeks the placenta becomes the major progesterone source, using circulating cholesterol as a substrate, after which maternal serum levels of progesterone increase markedly. Progesterone in serum is to 95-99% bound to corticosteroid binding globulin (CBG) almost as tightly as glucocorticoids (Speroff et al 1994).

Progesterone accompanies and modulates estrogen action. Whereas progesterone is synthesized in the placenta, neither the placenta nor the fetal adrenal glands are capable of producing sufficient quantities of precursors for estrogen synthesis. This observation led to the coining of the unique endocrine system “the maternal-fetal-placental unit” (Diczfaluzy 1969). In early pregnancy, the maternal circulation provides androgen precursors for estrogen synthesis. By 20 gestational weeks the majority of androgen precursors, predominantly dehydroepiandrosterone sulphate (DHEAS), are derived from the fetal adrenals. The fetal compartment is extremely efficient in sulphate conjugation of steroid hormones, protecting the fetus from high steroid concentrations. About 30% of circulating estrogens are loosely bound to albumin, whereas the major amount is tightly bound to sex hormone binding globulin (SHBG) (Speroff et al 1994).

2. Mechanisms of action

2.1 Genomic effects

The nuclear progesterone receptor (nPR) belongs to the steroid supergroup of transcription factor proteins (O’Malley et al 1990). All steroid receptor proteins are composed of a variable N-terminal domain which activates gene transcription and protein-protein interactions, determining the biological response of the steroid, an evolutionary highly conserved DNA-binding domain, a flexible hinge region and a C-terminal ligand-binding domain. The classic genomic mechanism of steroid action involving mRNA and protein synthesis is slow, occurring over hours to days.

The nPR binds to progesterone, and with a much less affinity to cortisol (Sanborn et al 1976). The biological response to progesterone is dependent on the levels and ratios of the nPR isoforms. The nPR isoforms A (nPR-A) (94 kDa) and nPR-B (116 kDa) are transcribed from the same gene, being activated by different promoters. The nPR-B isoform contains an additional 164 amino acids at the N-terminal and activates progesterone responsive genes. The nPR-A isoform is a weaker activator of transcription than nPR-B and can act as an inhibitor of nPR-B and other steroid receptors such as the nuclear estrogen (nER) and nuclear glucocorticoid (nGR) receptors (Vegeto et al 1993, Pieber et al 2001). A third nPR-C isoform has been identified in human myometrium (Condon et al 2006). nPR-C lacks a large segment of the N-terminal and a major part of the DNA-binding domain, and therefore cannot bind to DNA.
Fig. 2. The steroid hormone receptor family. A. Phylogenetic tree of the steroid hormone receptors showing the evolutionary interrelationships between the receptors. B. Sequence homologies of intracellular steroid hormone receptor proteins showing the N-terminal domain (A/B), the DNA-binding domain (DBD, C), the hinge region (D) and the C-terminal ligand binding domain (LBD, E). The human estrogen receptor subtypes (ERα and ERβ), glucocorticoid receptor (GR), progesterone receptor isoforms (PRA and PRB), androgen receptor (AR), and mineralocorticoid receptor (MR) are described. The estrogen receptor α is unique in that it contains an additional C-terminal F domain. Numbers represent the amino acid sequency of the receptors. In Griekspoor A, et al. Nuclear Receptor Signaling (2007) 5, e003.

2.2 Non-genomic effects
Steroid hormones have been shown to initiate rapid actions, which cannot be explained by the slow genomic mechanisms. Such rapid actions occur within seconds through the activation of intracellular signaling pathways resulting in alterations in ion fluxes and intracellular free calcium concentrations (Blackmore et al 1991), and within minutes through the activation of other second messengers, such as cyclic nucleotides and extracellular-
regulated kinase (ERK) 1 and 2 (Filardo et al 2000). Recently, three new putative membrane progesterone receptors (mPRs), mPRα, mPRβ, and mPRγ were identified in humans (Zhu et al 2003).

Fig. 3. G (guanine nucleotide-binding) protein-coupled transmembrane (TM) receptors communicate signals from hormones and other signaling factors to intracellular messengers. They consist of the Ga and the tightly associated Gβ,γ subunits. Here guanosine-triphosphate (GTP) is hydrolyzed by Ga subunit to guanosine-diphosphate (GDP). Source: CellMosaic, Worcester, MA, US.

Fig. 4. Nuclear and transmembrane-bound progesterone receptors mediating genomic and non-genomic effects. Progesterone (P) activates A. Genomic pathways through nuclear receptor proteins (PR-A, PR-B) resulting in gene activation (slow process) and/or B. Non-genomic pathways via membrane-bound receptors, which activate secondary messengers (fast process).
3. Systemic effects

Serum levels of progesterone increase progressively during human pregnancy and remain high until delivery of the placenta (Caspo et al. 1973, Speroff et al. 1994, Stjernholm et al. 1997). In other species, such as rodents and rabbits, which depend on an active corpus luteum for progesterone synthesis throughout pregnancy, labor is initiated by prostaglandin F₂α (PG-F₂α) from the endometrium, activating prostaglandin F (FP)-receptors in the corpus luteum leading to luteolysis (Sugimoto et al. 1997). These observations led to the concept of a “functional progesterone withdrawal” at parturition in humans (Hertelendy & Zakar 2004).

3.1 The placenta

Placental concentrations of progesterone reach 1-10 μM (Stites & Siiteri 1983, Miyaura & Iwata 2002), whereas serum concentrations reach 100-500 nM until term pregnancy before labor (Stjernholm et al. 1997; Miyaura & Iwata 2002).

3.2 The vascular system

Increased levels of prostacyclin (PGI₂) are considered to be a factor behind the physiological angiotensin resistance observed in normal pregnancy (Friedman 1988). A progesterone induced mechanism behind this refractoriness to angiotensin has been suggested (Everett et al. 1978, Rupnow et al. 2002). Nitric oxide (NO) and protein kinase C (PKC) pathways are involved in the regulation of vascular tone during pregnancy (Kublickienė et al. 1997, Chang et al. 2008).

3.3 The respiratory system

The pulmonary function is not impaired by pregnancy, but the tidal volume, minute ventilator volume and minute oxygen uptake increase with advancing gestation. This pregnancy-induced respiratory alkalosis is partially compensated for by increased renal excretion of bicarbonate. As a consequence, maternal arterial pH is increased to 7.46. The increased respiratory effort and decrease in PCO₂ has been related to progesterone and to a lesser degree to estrogen (Wolfe et al. 1998, Jensen et al. 2005).

4. The uterus

4.1 The decidua at implantation

Successful maintenance of pregnancy depends on maternal tolerance of the fetal semi-allograft (Szekeres-Bartho 2002). Progesterone, cortisol and prolactin have strong immunomodulatory effects leading to immunotolerance during pregnancy (Stites & Siiteri 1983, Speroff et al. 1994). The human decidua is adjacent to the myometrium, the fetal trophoblasts of the placenta and to the fetal membranes. Natural killer (NK) cells is the predominant immune cell in the decidua before implantation and in early pregnancy, constituting 70% of decidual immune cells, followed by macrophages constituting about 10% of total decidual cells, dendritic cells (DC) and T lymphocytes. The local endocrine environment regulates the recruitment of monocytes into the uterus, and the subsequent differentiation of monocytes into macrophages with specific phenotypes promoting immunotolerance or inflammation (Stout et al. 2004). Colony-stimulating factor (CSF)-1, macrophage migration inhibitory factor (MIF), monocyte chemoattractant protein (MCP)-1 and regulated on activation, normal T cell expressed, and secreted (RANTES) have been
suggested as factors involved in the recruitment and modulation of decidual macrophages, and are synthesized by decidual stromal cells, NK-cells and trophoblasts at the maternal-fetal interface (Wood et al, 1997; Lockwood et al 2006). Resident decidual macrophages appear to express immunosuppressive actions that favor the maintenance of pregnancy. In contrast, monocytes/macrophages migrating into the lower uterine segment prior to parturition are involved in the inflammatory process associated with cervical ripening and labor initiation (Nagamatsu et al 2010). A switch in decidual type 1 (Th1) to type 2 (Th2) T cell dominance in the fetal-placental interface has been suggested to play a crucial role in the establishment of pregnancy (Wegmann et al 1993). Human Th1 T-cells are the main effectors of host defence and Th1-type cytokines produce proinflammatory responses. The Th-1 response involves interferon (IFN)-γ, interleukin-2 (IL-2), tumor necrosis factor (TNF)-α, and the generation of cell-mediated immunity. On the contrary, human Th2 T cells inhibit macrophage functions. A Th2 response involves IL-4, IL-5, anti-inflammatory IL-10, IL-10, IL-13, and the stimulation of humoral immunity (Abbas et al 1996, Weiner et al 2001).

Fig. 5. Endocrine and immune cross-talk in the fetal-maternal interface at implantation. CL= corpus luteum, HCG= human chorionic gonadotropin. In Fujiwara H. Molecular Human Reproduction (2009) 15, 335–343.

Progesterone at concentrations higher than in serum but comparable to those in the maternal-fetal interface induces differentiation of T cells along the Th2 pathway (Stites & Siiteri 1983; Piccinni et al 1995, Miyaura & Iwata 2002). Glucocorticoids and 1,25-dihydroxy Vitamin D increase IL-4 (Rook et al 1994), whereas dihydrotestosterone decreases IL-4 and IL-5 production (Vacca et al 1990).
The progesterone induced protective immune environment in the decidua during early pregnancy includes production of the immunomodulatory progesterone-induced blocking factor (PIBF) protein by decidual cells (Szekeres-Bartho et al 1985, Piccinni et al 1995). The presence of nPRs in immune cells has been debated. nPRs in the thymus are necessary for progesterone induced involution of the thymus during pregnancy (Tibbetts et al 1999). Most studies have showed an absence of nPRs in lymphocytes from nonpregnant women (Szekeres-Bartho et al 1990, Mansour et al 1994, Bamberger et al 1999). Recently, transcripts for mPRα and mPRβ but not mPRγ, were detected in human peripheral blood leukocytes and T lymphocytes. Progesterone activated an inhibitory G-protein (Gi), suggesting that mPRs are coupled to Gi. These results suggest a potential novel mechanism for progesterone’s immunoregulatory function through activation of mPRs (Dosio et al 2008).

The establishment of human pregnancy is associated with an adequate synthesis of leukemia inhibitory factor (LIF), and macrophage colony-stimulating factor (M-CSF) producing T-cells. Progesterone at concentrations comparable to those in the maternal-fetal interface induces LIF and M-CSF (Piccinni 2010).

4.2 The myometrium
The corpus uteri is a muscular organ with about 70% smooth muscle cells surrounded by extracellular matrix (Danforth 1954). Progesterone is holding the uterine myometrium in a quiescent state, “a progesterone block”, during pregnancy by suppressing the propagation of electrical activity between the excitable myocyte membranes (Csapo 1956, Csapo et al 1973). The genomic and nongenomic pathways co-operate to maintain myometrial relaxation. At parturition, a functional progesterone withdrawal occurs by increased expression of the nPR-A and/or nPR-C to nPR-B ratios and changes in nPR co-regulator levels which result in repression of the nPR-B transcriptional activity. The diminished progesterone influence leads to an estrogen dominance (Mesiano et al 2002). Prostaglandins have been shown to induce an increased nPR-A/nPR-B ratio through the protein kinase C (PKC) pathway in human myometrial cells (Madsen et al 2004). Proinflammatory IL-1β up-regulates nPR-C in human myometrial cells, leading to diminished activation of nPR-B (Condon et al 2006). The increased expression of specific membrane-associated PRs (mPRs) at parturition augments contractility by decreasing intracellular cyclic adenosine monophosphate (cAMP) and altering intracellular Ca²⁺ levels. (Pieber et al 2001, Mesiano et al 2002, Madsen et al 2004, Mesiano 2007).

5. The cervix uteri
The cervix uteri is up to 85% composed by connective tissue, which is dominated by collagen fibers. Fibroblasts, smooth muscle cells, T and B lymphocytes, leukocytes and Langerhans cells are scattered within the tissue (Danforth & Evanston 1954, Schwalm & Dubrauszky 1966, White et al 1997). Cervical remodeling is a prerequisite for cervical effacement and dilatation prior to labor and is characterized by increased levels of vascular adhesion molecules (VCAM), diapedesis and activation of neutrophils, monocytes/macrophages, T lymphocytes, mast cells, eosinophils, the release of proinflammatory cytokines such as IL-1β and the strong chemotactor IL-8, and increased tissue concentrations of metalloproteinase enzymes (MMPs) (Junquiera et al 1980, Liggins 1981, Uldbjerg et al 1983, Bokström et al 1997, Sennström et al 2000, Stygar et al 2002, Winkler et al 2003). At parturition, a functional progesterone withdrawal occurs in the cervix uteri with decreased total nPR and an increased nPR-A/nPR-B ratio.
These endocrine and inflammatory events are followed by an up to 30-50% decreased collagen concentration, and an altered proteoglycan composition with a decreased density of the small proteoglycan decorin and an increased density of the large proteoglycan Versican. These events result in dispersed collagen fibrils clinically recognized as cervical effacement and dilatation (Uldbjerg et al 1983, Ekman et al 1986, Norman et al 1993, Stjernholm et al 1997). Evidence suggest that progesterone effects on the cervix uteri are even more pronounced than its effects on the myometrium (Romero 2007).

Fig. 6. Cervical effacement and dilatation before labor onset.

6. Parturition

6.1 Animal studies
Classical experiments in sheep demonstrated that parturition in this species is initiated by activation of the fetal hypothalamic pituitary adrenal (HPA) axis leading to increased fetal cortisol secretion and induction of placental P450 enzymes (17α-hydroxylase and 17-20-lyase activities), which favor the conversion of C21 to C18 steroids (Liggins 1974, Anderson et al 1975).

6.2 Human parturition
Progesterone is the main progestational hormone in humans, whereas the HPA axis has a modulatory function (Hertelendy & Zakar 2004). Prostaglandins (PGs) from the E and F series are considered to be the main promoters of cervical ripening and myometrial contractility, and the influence of PG-E₂ in promotion of cervical maturation and uterine vasodilatation has been
suggested as the primary functions of PGs in human parturition (Hertelendy & Zakar 2004). Human decidual macrophages synthetize PGs (Norwitz 1991). Mechanical stretch of the lower uterine segment, proinflammatory cytokines such as IL-1β and the peptide hormone oxytocin induce PG synthesis (Molnar et al 1999, Allport et al 2001, Leguizamon et al 2001). Successful treatment with PG-E2 for cervical priming before labor induction, allowing for resulting in cervical effacement and dilatation allowing for parturition was associated with diminished cervical progesterone and androgen receptor concentrations (Vladic Stjernholm, 2009).

7. The puerperium

After delivery of the placenta, serum concentrations of estrogen and progesterone decrease within hours, and the puerperum (puer: infant, pario: give birth) is a hypoestrogenic and hypoprogestagenic state. The high progesterone level during pregnancy inhibits lactation. The fall in progesterone levels after delivery is one factor that stimulates milk production (Tucker 1979).

8. Progestin and progesterone treatment

Natural progesterone and synthetic progestins do both exert a progestogenic effect, defined as the decidualizing effect on estrogen-primed rabbit endometium (Elton 1966, Schindler et al 2003).

8.1 Progestins and progesterone for preventing miscarriage

In clinical practice, progestin treatment was practised since the 1950s as luteal phase support to prevent miscarriage during the first trimester of pregnancy. The amount of data from well-controlled clinical trials is limited. Further studies are required to establish the optimal treatment situation as well as type and dose of progestin (LeVine et al 1964, Daya & Gunby 2004).

8.2 Progestins and progesterone for preventing premature birth

Since the 1960s studies on treatment with synthetic progestins for preventing premature childbirth have reported beneficial effects. Human pregnancy lasts 40 gestational weeks and birth between 22 and 37 weeks is defined as premature (WHO 1977). The highly active progesterone ester 17α-hydroxyprogestrone caproate has a long duration allowing for

Fig. 7. The synthetic progestin 17α-hydroxyprogestrone caproate.
intramuscular administration 1-3 times weekly. It has been administered to risk groups with previous recurrent abortions or previous premature births and to patients with premature contractions and short cervices (Johnsson et al 1976, Meis et al 2003, Dodd et al 2006). Natural progesterone has been administered as of vaginal gel to such risk groups and in situations with premature contractions and short cervices (daFonseca et al 2003, deFranco et al 2007, O’Brien et al 2007). Reduced incidence of premature birth before 32, 34 and 37 gestational weeks and improved neonatal outcome were reported (Brent 2005). Further studies are required to establish the optimal dose and type of agent as well as long term effects on the newborn.

9. Summary

Progesterone is the evolutionary most conserved of the reproductive steroid hormones. It is the main progestational hormone in humans, and its strong immunomodulatory effects are important for the physiological immunotolerance at implantation. After a transition period by 7-10 gestational weeks the placenta becomes the major progesterone source. Placental concentrations of progesterone reach 1-10 µM and serum concentrations 100-500 nM until term pregnancy. Progesterone exerts its effects through genomic nuclear receptor mediated and non-genomic transmembrane receptor mediated processes, keeping the myometrium in a quiescent state and stabilizing the cervix uteri during pregnancy. A functional progesterone withdrawal occurs at human parturition with a diminished total receptor density and altered isoform ratios. In clinical practice, progestin treatment has been given as luteal phase support to prevent miscarriage during the first trimester of pregnancy. Treatment with synthetic progestins and natural progesterone has been shown to reduce the incidence of premature birth. Further studies are required to establish the optimal dose and type of agent as well as long term effects on the newborn.

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11. References


Sex Hormones not only regulate reproductive function, but they also play a prominent role in the biology and physiology of several organs/tissues and in the pathophysiology of several diseases. During the last two decades, the information on the mechanisms of action of sex hormones, such as estrogens and androgens, has rapidly evolved from the conventional nuclear receptor dependent mechanisms to include additional non-nuclear, non-genomic and receptor-independent mechanisms. This highlights the need to update the current knowledge on sex hormones and their mode of action. Increasing evidence that exogenous/epigenetic factors can influence sex hormone production and action highlights the need to update our knowledge on the mechanisms involved. This book provides a systematic and updated overview of the male/female sex-hormones and their impact in the biology and physiology of various organs. Additionally, the book discusses their positive and negative association with the pathophysiology of various diseases (e.g. osteoporosis, cardiovascular-disease, hypogonadism, reproduction, cancer) and their therapeutic potential.

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