Chapter

Potential Therapeutic Options and Perspectives for Alleviation of Endometrial Estrogen Dominance and Progesterone Resistance in Endometriosis

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Abstract

Endometriosis is a chronic disease, influenced by internal and external environment, with long duration from intrauterine life with acme during childbearing, when it is associated to chronic pelvic pains, and infertility/subfertility. DNA hypermethylation of endometrial promoter PRs Hox genes and DNA hypomethylation of promoter ERβ gene is a possible explanation of estrogen dominance, progressive loss of progesterone signaling, followed by progesterone resistance in ectopic, and progesterone attenuance in eutopic endometrium, for failure of hormone therapy (HT), repeated recurrences after surgery, cancers after long time evolution. Animal models, human trials demonstrated progesterone (P4) and progestins influences over progression of disease pathological characteristics, associated to endometrial ER, PR aberrant expressions: ERα loss, and abnormal PRB/PRA ratio. P4 supplementation before mice induced-endometriosis protected from PRs depletion, action that can be translated in women according to the difference of 7 to 12 years between histologic onset and clinical symptoms/signs, parallel to progressive loss of PRs and PR-mediated signaling in ectopic and eutopic endometria. The animal studies have shown that a DNA methylation inhibitor alleviates lesion growth, and induces PRs target gene expression restoration. Continuous/extended contraceptives, dienogest-a new progestin, GnRH agonists/antagonists, aromatase inhibitors, SERM, SPRM, combined molecules are therapeutic options/perspectives aiming restoration endometrial estrogen-progesterone balance, without disease’s cure. HT may be active alone, or surgery associated.

Keywords: endometriosis, estrogen dominance, progesterone resistance/attenuance, timing, progesterone/progestins, DNA methylation inhibition

1. Introduction

Urgent innovations are demanded in endometriosis management, which should start by deeper understanding of disease core features, with disease different phenotypes and idiosyncrasies. Endometriosis, a chronic inflammatory and immune disease, dependent on environment factors, genes- Hox genes.
code (HOXA10/HOXA11), epigenetics, and ovarian steroid hormones, with their receptors and coregulators in the eutopic endometrium and ectopic sites, was demonstrated to have a long duration of progress from intrauterine life, latency in childhood, and acme during reproductive years [1–3] with reappearance in postmenopause when hormone replacement therapy, or even without [4]. This fact permits therapeutic intervention to correct endometrial abnormal functions during menstrual cycle, to prepare decidua receptivity for successful egg implantation in women with minimal or mild to moderate endometriosis, in order to avoid the severe stages, and lack of response to therapy, considering that progesterone resistance is an acquired property of eutopic endometrium [5]. The therapeutic aims are to prevent, or at least to stop the progressive damages induced by ectopic endometriotic lesions in the uterus, entire genital tract, and in extra-uterine sites by the endometrial mesenchymal stromal/stem/progenitor cells with their specific migratory, adhesive, and invasive properties, with their genome changes when are outside the uterus, to influence the self-protected endometriotic lesions [6] and to ensure a normal eutopic endometrial cycle functioning, with normal ovarian steroid hormones receptors distribution during menstrual cycle. The aims can be individually accomplished if therapy is started from early stages of illness, by moving from pure hormonal therapy to drug combination, or novel molecules- SERMs, SPRMs which can escape the disrupted homeostatic mechanisms characteristic for endometriosis [7].

One may propose and discuss a perspective “timing” of endometriosis treatment, to maintain or for an early restoration of endometrial estrogen-progesterone receptors natural balance [8] being recognized a still delay in diagnosis, and treatment [9] delay that permits the evolution of genital damages by progressing endometriosis, and in time the risk of atypical endometriosis, and cancers which is very difficult to be assessed clinically, without specific biochemical markers [10].

The incidence of endometriosis of one case in 10 women of reproductive age [11, 12] or around 17% [13] is increasing when one assesses infertility and chronic pelvic pains - up to 50% [14] respectively up to 60% [15] recently being estimated 176 million women worldwide, making endometriosis as common as diabetes mellitus [16].

There are three distinct forms of disease according to ectopic sites of endometrial-like tissue location (peritoneal, ovarian and rectovaginal endometriosis), each of them being associated with specific symptoms, although dysmenorrhea, and chronic non-menstrual pelvic pains, dyschezia, dyspareunia are the most frequent [17] The literature makes some differences regarding the score of endometriosis, namely the ENZIAN classification/score describes superficial (less than 1 mm), intermediate (2 to 4 mm), deep (greater than or equal to 5 mm), and very deep implants (greater than 10 mm) [18]. The ENZIAN score was proposed of the revised American Society of Reproductive Medicine score (1996) [19], which was demonstrated to be less adequate to clinics, and the ENZIAN score was recently reviewed by experts in imagistic and surgical diagnosis [20]. So endometriosis has a similarity to oncology with 4 stages of evolution, with the difference of missing nuclear atypia [21] but with possible evolution to malignancy in 1% cases [1, 22] or further it may be associated to hematopoietic and breast cancer (HR of 1.3–95% CI 1.1–1.4, in Swidish women) [23] and ovarian cancer, – OR of 1.73, (95% CI: 1.10–2.71) on a pool of women from different continents- Australia, Canada, Denmark, USA, during 10 years (1989–1999) with previous endometriosis [24]. The natural history of endometriosis is uncertain, e.g. it is not known whether superficial peritoneal endometriosis (SPE) can progress to become another subtype, regress spontaneously, or whether disease progression (or lack of treatment) can lead to problems with infertility, and there is poor correlation between pain severity and the amount, location, and subtype of the ectopic lesions [16].
The literature does not mention a conclusive noninvasive diagnostic available tool to allow early detection of endometriosis, the delay in detection being of 7–12 years of latency from symptoms onset to the definitive diagnosis, and even in the best medical centres the full extent of the disease may be unknown [25] and up to 2017 it was not identified any fully validated, symptom-based, patient-reported questionnaire for endometriosis screening in adult women with potential endometriosis [26]. One must think to endometriosis in adolescent girls with family history, abnormal characteristics of menstrual cycles with the aim to avoid the structural and functional abnormalities progression induced in women's endometrial genes - mainly Hox codes for steroid hormones receptors, ligands, and co-regulators.

The next pages will discuss concepts, theories or hypothesis of potential therapeutic options and perspectives for prevention, normalization or restoration the endometrial estrogen-progesterone balance, and their receptors in the epithelium and stroma, because many years there were recommended ovarian suppressing drugs, and surgery.

2. Endometrial phenotypes in endometriosis

The interest for diagnosis and therapeutic success connected to side effects, limitations and failure rate of different classes of medication, and high risks of recurrence after surgery or medication discontinuation imposed the analysis of endometrium phenotypes, proliferative and differentiation in vitro capacities of stromal cells from ectopic lesions of peritoneum, ovaries, and deeply infiltrating endometriosis in comparison to the same structures of normal women, using the new techniques as contrast microscopy, immunocytochemistry, functional bioassays [27] and RT-qPCR of endometrial genes. The study of Burney et al. [28] analyzed eutopic and ectopic endometria comparative to non-ill women, and it was demonstrated a molecular phenotype of attenuated progesterone response within eutopic endometrium, from the dysregulation of numerous genes which are progesterone regulated, and the progesterone resistant phenotype is more frequent in ectopic endometrium comparative to non-ill cases. This condition is associated to a pro-inflammatory phenotype [29, 30] which increases both estrogen dominance, and progesterone resistance. The eutopic endometrium of ill women has an attenuated response to P4 because estrogen-responsive genes are not suppressed in their stromal cells in early secretory phase of menstrual cycle comparative to normal [28, 31, 32]. The British and Italian doctors’ conclusion [29] was that endometriotic cell lines, and stromal eutopic endometrial cells in ill-women are losing the capacities of differentiation, and respectively of decidualisation, aspects that explain cells capacities for proliferation and survival in the ectopic environment, and high infertility/subfertility rates. Deep endometriosis appears to be a special type, because predominance of PR less active isoform (PR-A) over the full length, due to epigenetic abnormalities affecting PR gene transcription, associated to oxidative stress, facts that induce a condition of more resistant to size regression upon medical treatments [7].

3. Concepts on endometrial peculiarities, as a steroid hormones target in endometriosis. Estrogen dominance. Progesterone resistance. Progesterone attenuance

The efficacy of endometriosis therapies may be improved by dissecting the unique molecular properties of eutopic and ectopic endometria compared to normal
endometrium. At first glance, the terms “estrogen dominance” and “progesterone resistance” appear to describe opposite sides of the same coin [6]. Endometrial Progesterone Resistance is described since many years [33] as there are other hormones resistance, being first named as “Pseudocorpus Luteum Insufficiency” [34] meaning that endometrium is not able to respond to bioavailable P4 plasma levels in between normal limits. In non-ill women E2 induces epithelial proliferation to build endometrial thickness during the proliferative phase of the menstrual cycle, then P4 inhibits E2-induced proliferation and allows stromal cells to begin decidu-alization during the secretory phase [35] in order to prepare the stroma to become receptive to blastocyst invasion during “window of receptivity/implantation” in normal, fertile women [36].

3.1 Dysregulation of endometrial steroid hormones receptors in endometriosis

The molecular mechanism triggered by ovarian steroid hormones in endome-trium is well known: steroid hormones link to specific nuclear (ERα, ERβ; PRA, PR-B, with specific ratios between them), cytoplasmic (ERβ), and membrane receptors (non- canonical G protein-coupled estrogen receptor (GPER) [37] and PGRMC-1 and PGRMC-2), which are able to bind to the promoter of the target genes, being up- or down- regulated by their co-activators (SRC-1, SRC-2, SRC-3 for estrogen receptors- ER) or downstream effectors (TGFβ, Dickkopf-1, retinoic acid, c-myc, etc. for progesterone receptors- PRs) [28] and influence endometrial cells proliferation and differentiation [38]. Endometrial repair after menstruation, proliferation and/or differentiation processes are dysregulated in endometriosis when menstrual reflux with endometrial stromal and epithelial glands cells with their exosomes migrate, adhere and invade peritoneal surface and/or other ectopic sites, associated to waves of inflammation (by systemic and local reactive responses to the presence of endometrial debris), neuro- angiogenesis and neuroinflam-mation, and aberant scar formation through fibrosis and adherences (possible a self-protection mechanism for ectopic lesion, as in chronic infections) at every ovarian cycle (different from normal endometrium restoration after each menstrual cycle). All these events are genetic controlled, epigenetic changed by hyper (for PR promoter) or hypomethylation (for ERβ) of DNA steroid hormones receptors promoters [39–41]. When the tightly regulated balance of epithelial- stromal P4 and E2 signaling is lost, P4 resistance and E2 dominance are prone to ensue, as in endometriosis [6]. P4 resistance may lead to both increased lesion growth and a non-receptive endometrium, and actually one speaks also about progesterone attenuated response in eutopic endometrium, which is associated to infertility/ subfertility. Estrogen dominance, progesterone resistance and progesterone attenuance in the ectopic and eutopic endometrium are explained by the dysregulation of estrogen and progesterone receptors their genes (ESR1 and ESR2) [32; a single gene- PRG for PRs] [42] with their micro-RNA (miRNA) –the cells’ critical regulators for development, and physiology, their mis-expression being associated to pathology [43].

When dysregulation condition persists it leads to both increased lesion growth and progression to a superior stage, and a non-receptive endometrium. P4 acts on stromal cells of the normal endometrium, inducing paracrine factor(s), and they induce the expression of the enzyme 17β-hydroxysteroid dehydrogenase type 2 (17β-HSD-2) for metabolization of E2 to estrone (E1), in the epithelial cells, and cell proliferation arrest. Excess E2 with Estrogen Dominance, and Progesterone Resistance are well documented in ectopic endometrium with aberrant levels of ERs and changed ER-α/ ER β ratio, where the genes analyses have proved to become an acquired property, through their migration and exposure process to
peritoneal environment [5]. ER-α and ER-β have essential roles in establishment and progression of ectopic lesions [32]. ERβ is excessively expressed in stromal cells of the ectopic lesions, versus non-ill women, fact due to hypomethylation of ERβ promoter, which contributes to low ER-α expression in ectopic endometrium [41, 44]. ERβ protein network together with SRC-1 coactivator isoform with which it cooperates, may have a cytoplasmic, non-genomic action in endometriosis as Han SJ, Jung SY, Wu SP, et al. discovered [44] (Figure 1).

Eutopic endometrium may prove the same endometrial dominance and progesterone resistance, but after some years of disease evolution, being described cases with initial progesterone attenuation in eutopic endometrium and progesterone resistance in the ectopic sites, usually in deep endometriosis [28]. It is documented a local increased synthesis of estrogen [45, 46] not a systemic high level [47] through the presence of the enzymes (aromatase, and 17β-hydroxysteroid dehydrogenase-1, 17β-HSD-1) [48, 49] and a blunted response to progesterone in eutopic and ectopic endometrium, P4 serum levels being similar to non-ill women [45]. In human and experimental mice the ER-α content may be normal in eutopic endometrium, as in non-ill women, but ER β are increased in both epithelial and stromal endometrial cells of eutopic and ectopic sites, as PRs levels are reduced in eutopic endometrium and PRs are lost in ectopic sites [44]. A recent analyses of ERs in deep endometriosis revealed that ER-α is a subtype for this condition [50] ER-α levels being predictive for symptoms severity, and for responses to treatment [51]. Progesterone attenuation of eutopic endometrium is connected to altered endometrial receptivity in the “window of receptivity/implantation” and to a significant reduction of implantation rate in ill-patients trying in vitro fertilization, due to stromal cells impair decidualization proved by a reduction of nearly 2-fold in IGFBP-1, and of leukemia inhibitory factor (LIF) [52] and to reduction/dysregulation of P4 target genes during the “window of receptivity/implantation”, time when normally, the endometrium is exposed to the highest levels of P4 [53] as it is strikingly down-regulated glycodelin - the prototype progesterone-responsive gene, in eutopic endometrium of ill-women compared to non-ill [53, 54]. LIF low levels are an intrinsic glandular dysfunction, induced by the gland-specific transcription factor Forkhead box A2 (FOXA2) low level [55, 56] also a P4 gene target. There are contradictions on the status of ERs and PRs in ectopic endometrium – an ERs increased expression, or others show reduced expression of both ERα and ERβ.
endometriosis [57–59] and lower levels of PRs [5, 60]. The eutopic endometrium has an attenuate answer to P4, the isoform PR- B is not expressed in patients’ endometrium, being only the isoform PR- A, because progesterone-responsive genes are not deleted in eutopic endometrium in comparison to normal women in the early secretory phase of cycle [28–32], a normal PR-A/PR-B ratio is very important in endometrial function. One may consider that the relative differences between studies regarding the PR isoforms loss in ectopic and eutopic endometrium may be explained by stage of disease, type of lesion and cells, and method of analysis.

Previous studies [32] demonstrated in experimental mice that high levels of ERα are driving proliferation, adhesion and angiogenesis in ectopic tissue, and also modulate inflammation, and ERβ prevents apoptosis and enhance invasion, proliferation, adhesion, and inflammation to stimulate the growth of ectopic lesion, so both isoforms might sinergistically contribute to regulation of proliferation, adhesion, and inflammation in endometriotic lesion. These discrepant findings are explained by the differences in study design, patient selection criteria, cycle stage, and endometriosis type and stage. Similar to these contradictory results on endometrial steroid hormones receptors is the situation with miRNA identification by real-time quantitative reverse transcription-polymerase chain reaction (real time qRT-PCR) in normal endometrium, eutopic and ectopic sites, connected to mi-RNA upregulation (over expression) or down regulation (under expression) in eutopic and in different ectopic areas (peritoneal, ovarian), and some are “mis-expressed” endometriosis [61]. There are differences between authors, with conflicting reports on whether or not miRNA expression was influenced by the menstrual cycle phases, endometrial cell type, miRNA type, level in ectopic/eutopic tissue, and stage of disease.

3.2 High micro-heterogeneity of endometrial steroid hormones receptors signaling in endometriosis

Normal endometrium is containing large quantities of distinct stromal cells with abundant estrogen-induced PRs, which influence glandular epithelial cell proliferation and differentiation, and protect against carcinogenic transformation, when PRA/PRB are in a proper ratio to ensure normal P4 response. PRA and PRB are members of a superfamily of almost 50 ligand-activated nuclear transcription factors [62]. In-vitro studies suggest that the two PR isoforms differ functionally, and that their relative expression in a target cell may determine the nature and magnitude of response to P4. The two isoforms are in comparable levels expressed in proliferative phase, but in the mid- secretory only PR- B is present in the epithelial glands, PR- A is predominantly present in the stroma throughout the cycle. It is a homogeneity in the relative expression in PR-A and PR-B in adjacent cells within the same tissue compartment, and a heterogeneity between glands, observed under some circumstances in the endometrium functionalis, suggesting that PR isoforms down-regulation by P4 is asynchronous, and between the glands of the basalis and functionalis of the endometrium implying region specific responses to hormonal stimuli [63].

A recent European study on deep endometriosis [64] showed a high variability of ERα and PR distribution in the same gland, among distinct glands of the same sample, and among distinct patients receiving the same treatment. Luminal epithelial height variability was primarily due to epithelial cells heterogeneity in a gland, secondarily to the glands randomaly evaluated on the same section, and tertiary to the patient category. The heterogeneity of ERα and PR distribution in the same women could explain why endocrine treatments are unable to cure deep endometriosis. The cause of heterogeneity in endometriotic tissues is difficult to be ascertain, one hypothesis being the DNA methylation of the steroid hormones [65] or of their promoters [39, 40] or
an abnormal proteolysis of steroid hormones chaperons [66]), the co-chaperons are required by PR for signaling uterine cycles and implantation [67].

Mice induced endometriosis demonstrated a high inter-animal variation in the levels of ERs, PRs, in ectopic endometrium vs. controls; with variable levels by almost 100-fold within the same lesion, and with differences between two lesions from the same animal [68] aspect called “micro-heterogeneity”. The changes are tissue intrinsic, and some researchers propose that the variable outcomes in hormonal therapy for endometriosis could be possibly due to heterogeneity or polymorphism in the expression of steroid hormone receptors in the ectopic endometrium [68] or in all endometrial locations- eutopic, and ectopic, being known the heterogeneity of PRs in glands, and the homogeneity of PR isoforms in stroma of normal human endometrium [63] making the magnitude in response, the attenuation, and the resistance in response to P4, independently to the serum levels of P4 [69]. This intrinsic biologic alteration of eutopic endometrium explains the missing differences in endometrial thickness, and histology –respectively luteal differentiation, or epithelial integrin expression at the lower mid-luteal serum progesterone level (as 3–4 ng/ml) in programmed cycles of physiological and subphysiological exogenous progesterone replacement in GnRH agonist-suppressed healthy volunteers. One may consider that these results are an answer to the questions whether the abnormalities of eutopic endometrium in early secretory phase suggestive of attenuated progesterone response in the transition from proliferative to secretory phase are due to lower level of circulation or local bioavailable progesterone or to the changes in endometrial transcriptome, with dysregulation of progesterone- regulated genes.

3.3 Nuclear receptor coregulators in the modulation of progesterone and estrogen signaling in endometriosis

Endocrinologic literature presents different series of nuclear receptors coregulators,- proteins that intervene to modify chromatin structure and regulate large-scale gene transcription programs by forming large complexes with the nuclear receptors of the target cells [70]. In the female genital tract the PRG and ESR1 are critically regulated by a family of regulatory proteins named steroid receptor coactivators (SRCs) - SRC-1, SRC-2, and SRC-3, the first nuclear receptors coregulators are involved in the balance between E2 and P4 in human endometrium. SRC-1 down regulates P4 target genes in the epithelium, and up regulates them in the stroma; SRC-2 is necessary for human endometrial stromal cells decidualisation [71] for P4 signaling and for ESR1 signaling; the transcriptomic analysis revealed that 50% of SRC-2-regulated genes are also regulated by P4 [72] and it is critical for murine uterine function; ablation of SRC-2 induces partial loss of decidualisation and infertility, and both SRCs ablation induces complete loss of decidualisation [73, 74].

4. Potential therapeutic options to maintain/restore endometrial normal estrogen-progesterone signaling in endometriosis

Presently pharmacotherapy mainly with hormones (Hormone Therapy- HT) and surgery are two cornerstones in endometriosis management. Surgery with histological confirmation of ectopic endometrial glands and stroma remains the gold standard for diagnosis [75, 76] surgery being generally reserved for patients who fail medical therapy, or who desire pregnancy, being usually performed by laparoscopy [77, 78]. Surgery is able to eliminate visible endometriotic lesions, without the cure of the disease [79], and majority of drugs are symptomatic, not cytoreductive [80]. Yet it is not known if surgery itself may be incomplete (i.e., microscopic disease)
or if other factors, such as aberrant PRs expression of the eutopic endometrium influences recurrence [60] connected to the eutopic endometrium transcriptome changes compared to non-ill women, indicating abnormalities that predispose to new implants in extrauterine locations after surgical excision of ectopic lesions [81]. Recurrence rate after surgery or medication discontinuation was recorded up to 45% after 5 years [82] and recently one discusses the translation from adjuvant therapy to tertiary endometriosis prevention in postoperative medical care [83]. The moment of medical therapy associated to surgery- before, after, or both before and after surgery is much analyzed, to maximize treatment response, but literature data is still inconclusive [84]. The HT goal is to induce atrophy of endometriotic lesions, even if one missis the ability to predict which medication each individual patient will respond to, being many attempts to find one or more predictive markers [85] to score HT, as it is the immunohistochemical Histo (H) -score on the PRs status, and CYP19A1 expression for the response to progestins, respectively for the need to block estrogen signaling [60]. The high H-score for PRs was proposed to be >80 (associated with 100% positive predictive value), and the low H-score ≤ 5 (associated to 94% negative predictive value), with some authors’ comments regarding H-score usage: the score is useful for ectopic endometrium response to progestin- based therapy, not for eutopic endometrium, because the lack of score’s correlation to eutopic endometrium. It is recommend to score PRs in the excised ectopic tissues, and to determine the reason for progesterone/progestins/COCs failure. When adhesions/fibrosis are predominant it is sure the non-response to HT, but some PRs expression may indicate an insufficient dosage of progestin or noncompliance with therapy (i.e., inability to tolerate side effects), and for cases with low H-score, patients being PR resistant, it is advised to avoid progestins after surgery in favor of GnRH agonists or GnRH antagonists [86, 87], danazol- rarely used actually, or aromatase inhibitors.

Actual medical therapies allow suppression of endometriotic lesions, women require long term treatment [79] to maintain the benefits, to avoid recurrences, being recognized high rate relapse even after surgery. After surgery one must continue HT with contraceptive pills, or IUD [88] with the aim to preserve ovarian follicles reserve, and because ovarian aging, to try spontaneous pregnancy or ART.

Individual genetic characteristics can affect the bioavailability and pharmacodynamics of HT, mainly estrogen dominance and P4 resistance/attenuance, and, hence, explain part of the variability in the therapeutic response, as it is the polymorphism of CYP3A4/5 involved in levonorgestrel metabolism [89].

One can respond to therapeutic wishes by timing progesterone/progestins administration, and by moving from pure hormonal therapy to drug combination, or to novel molecules capable to restore the various homeostatic disrupted mechanisms by disease. A number of potential therapeutics are currently in pre-clinical or early clinical studies, fact which may alter further treatment strategies.

4.1 Progesterone/progestins therapy

Kistner [90] was first who postulated that decidualisation observed during pregnancy might cause necrosis, and the consequent elimination of superficial ectopic implants, and this was the logical reasoning for progesterone recommendation, and use to control endometriosis. During pregnancy on assists at ovarian functions suppression, fact that is mimicised by HT in endometriosis [91]. After Kistner RW first initiave in the years 1960, there were used progestins, which are inducing a pseudo-pregnancy endometrial aspect or endometrial atrophy at microscopy [92]. The progestins are synthetic different derivates, such as C-21 progesterone derivates (medroxyprogesterone acetate, MPA, and depot MPA, and dydrogesterone,
(DYD- a semisynthetic progesterone derivate), or C-19 nortestosterone derivates (levonorgestrel, norethisterone, lynestrenol, desogestrel, and dienogest), or the 19-norprogesterone derivates without androgenic activity; nomegestrol acetate – (NOMAC) has a similar anti-gonadotropic activity compared to the 19-nortestosterone derivates with androgenic activity, norethisterone acetate (NETA) [93] and a separate class with drospirenone, the unique progestin [94]. The COCs with the synthetic ethinyl-estradiol (20, 30 μg/pill) plus one of the previously mentioned progestins represent a very important cathegory in the control of eutopic and ectopic endometria from intrinsic abnormal steroids signaling [95]. One must keep in mind the Japanese opinion [96] that concurrent estrogen action is essential for maximal progestin effect in COC, fact controverted by some studies regarding patients satisfaction on pain alleviation in endometriosis, explained by summerizing endogenous estrogen local levels in ectopic endometrium to that of pills [97] but estrogen priming may be necessary to induce PR in endometriotic lesions [96, 98] as progesterone usually actions on estrogen primed tissue. Progesterone and progestins, alone or in association to a synthetic estrogen are considered first line therapy in endometriosis. The positive response to progestins alone or combined estrogen- progestins may induce only partial improvement, the ectopic sites are not eliminated or some patients do not respond at all [99] being recognized only the quiescence of lesions, at best [100] and when first line therapy fails – in one- third of women because progesterone resistance [79] or it is contraindicated, or it is not tolerated one recommends Gonadotropin-Releasing Hormone (GnRH) agonists, GnRH antagonists, and aromatase inhibitors being used in cases refractory to other therapy. Up to 2021 it was published a single RCT on COCs in endometriosis [101] as Donnez and Dolman [79] mentionned in the last European review. It is described a statistically significant, though modest, improvement in dysmenorrhea with OCPs given for four months compared to placebo, and a lack of any beneficial effect of OCPs on non-menstrual pelvic pain and dyspareunia, being appreciated the studies failure to report data on their efficacy according to lesion phenotype [102, 103].

Patients present variable responses to progesterone/progestins, according to their pharmacokinetics, to their capacities to reacts to medication (genes, receptors for E2, P4), to metabolize progesterone/progestins connected to their levels of N-Acetyl transferases, associated to drug kinetics control by acetylating and converting active forms into inactive metabolites, to their cytochrome P450 3A (CYP3A) system- one of the most known pathway in steroid hormones metabolism, and P450 3A polymorphism is not precisely known in endometrial pathology [89].

4.1.1 “Old” hormone therapies still active in endometriosis

During the long time of progestins usage it was reported a good rate of patient adherence and satisfaction [104, 105]. There are few direct comparisons between different types of progestins in endometriosis clinical studies, with the scarce evidence suggesting that when given by the same route and the same regimen, the effectiveness may be similar [7]. By binding to PRs, progesterone/the synthetic compounds, administrated orally or by non conventional routes (vaginal [106], intrauterine [107]) induce inhibition of estrogen synthesis through down regulation of ERs [64] and of steroidogenic enzymes in endometrial stromal cells, either by up regulation of the oxidative 17β-HSD type 2, which transforms E2 to E1 [108], being triggered by retinoic acid [109] or by reduction of 17β-HSD type 1 expression, and activity, – as was proved specially for MPA, DYD, dienogest [108, 110] which induces the inhibition of aromatase expression with a low local estrogen production in immortalized human endometrial epithelial cells [111] with inhibition of eutopic and endometriotic cells proliferation, and reducing cells survival by apoptosis.
induction [112] with limitation of local angiogenesis and neurogenesis (proved with the CD31- a neurovascularisation marker), and linking all these mechanisms, the attenuation of the immune-inflammatory response (proved to be more accentuated on new progestins as drosipirenone by its effects on inflammatory cytokines - VEGF, and nerve growth factor) [113] so progesterone/progestins change the morphology of endometrium- ectopic and eutopic with reduction of lesion sizes, but with net differences in their response to these drugs, as experimental and clinical studies demonstrated [7, 114], stages III- IV are not responding. Longitudinal assessment of endometrial morphology has demonstrated that the histology aspects are changing in time: initially one registers secretory differentiation, and after several cycles through ERs down- regulation one registers atrophy with tubular glands, weak secretory vacuolation, and stroma low cellular density [115]. Usually one recommends a HT for long term duration, short term therapy being insufficient to obtain the wanted effects, even with dienogest, — which in a short term therapy demonstrated a high frequency of decidualisation and a tendency of inflammation reduction, but it was not able to induce differences comparable to non- treated cases in terms of necrosis, glandular atrophy and angiogenesis [116] (Table 1).

4.1.2 Timing progesterone/progestins for preventing progesterone attenuance and progesterone resistance in endometriosis

The discussion on administration of progesterone from early ages of reproductive period, just to permit P4 to stop endometriosis progression, inflammation, and angiogenesis, by maintainance of ERs, PRs normal ratios in epithelial glands and stroma, is supported by Li et al. study [114] at University of Illinois at Urbana-Champaign, (USA). Endometriosis was induced in two immunocompetent mice groups, differently maintained afterwards: one group with E2, and the second group with E2 and P4 subcutaneously, at every 4 days beginning at 4 days before lesion induction (pre- P4 treatment). The endometriosis-like islands were very quickly developed, with different numbers, aspects and sizes according to steroids administrated- after E2 alone there were yellow, more numerous, larger, with abundant blood vessels and extensive adhesions; after E2 plus P4 were white, smaller, non-vascular, with loose attachment. The microscopy (H&E, special biomarkers, IHC for ER, PR, their genes) have shown marked differences regarding mitotic activity (Ki-67 reduction), glandular secretion, endothelial cells and angiogenesis (CD31 increased when was added P4 before induced endometriosis). Another peculiar aspect in the treated group with E2 plus P4 are the inflammatory response changes - first an increase, and later a reduction of inflammatory cells, and changes of their type in endometriosis-like tissues, with similarities between groups in the first 16 days, and a dramatically changed aspect of inflammation after 24 days from induced endometriosis, such as suppressed production of pro-inflammatory cytokines, and infiltration of immune cells in ectopic sites. The authors debated the P4 action to alleviate lesion outgrowth and to maintain ERα and PR expression

- P4 restricts expansion of the ectopic lesions by inhibiting endometrial cell proliferation, and neovascularization
- P4 suppresses E2-dependent inflammatory responses in the ectopic lesions
- P4 maintains ERα/PR-mediated signaling; their loss in the ectopic lesions leads to resistance to P4 therapy if the treatment is postinduction of lesion

Table 1. Progesterone alleviates endometriosis, induced in the peritoneal cavities of female immunocompetent mouse, maintained with estrogen if administered before illness induction (from Li et al. [114]).
when P4 is administrated before lesion induction. The use of RT-qPCR permitted
the endometrial genes assessment in eutopic and ectopic endometria, showing a
progressive down-regulation of miRNA expression corresponding to ESR1, PRs
and PR-stromal targets Hand2 and Hoxa-10 with time of disease progression, and
in contrast a gradually increase of ESR2 miRNA, parallel to increased expression of
ESR1, PRs, and HOXA 10 in ectopic lesions, while Hand2 expression remained sup-
pressed when P4 was administrated before lesion induction (PreP4). The P4 inhibi-
tory effect was not observed when P4 was started at 4 days after endometriosis
induction (Post-P4). The study results clearly indicate that the loss of PR-mediated
signaling components is a major causal factor for the P4 resistance existing in mice
with endometriosis. The conclusion is that progesterone supplementation from
ey early moments of disease, when PRs are still present can preserve steroid respon-
siveness and ameliorate E2-dependent disease progression, but when at later stage,
P4 supplementation has minimum effects, because of PRs absence. The hope is that
if PRs women’s loss is progressive after disease onset, the add of P4/progestins may
action with the wanted effects.

4.1.3 Not all progestins are equal in restoring estrogen-progesterone balance in
endometriosis. New trends-dienogest

If there were questions if progestins are all equally capable of acting on endo-
metriosis lesions to induce apoptosis, or to inhibit cell proliferation, adhesion,
invasiveness, angiogenesis/neuroangiogenesis, and inflammation, actually one
knows their different actions and effects, according to their biochemical structure,
and cross effects on glucocorticoid, mineralocorticoid and androgen receptors
[117]. In the last 20 years, starting with the Japanese experiments [118] on rats
induced- endometriosis, the progestin dienogest (DNG) was much analyzed in
Japan, USA, Europe (The European Clinical Study Program) [119] in order to avoid
progesterone resistance or progesterone attenuation, by reducing estrogen deleteri-
ous effects on endometrium- normal, eutopic and ectopic increasing PRs expres-
sion and decreasing proinflammatory cytokines, and the necessity of “add back”
therapy imposed by GhRha agonists [120, 121] as it will be further discussed. It was
demonstrated that DNG 2 mg/day (once/day in Europe, and 1 mg twice/day in
Japan) [120] after 24–52 weeks of administration reduces lesions size, [122] without
changing bleeding pattern [123] and with a good score regarding chronic pelvic
pains [120]. Because it was proved an in vitro dose-dependent inhibition of human
endometrial stomal cells proliferation together with morphological and functional
changes [124], an Italian clinical study on 20 cases of endometriosis [125] had
evaluated doses of 20 mg/day effects, for 24 weeks, with no comparative study-
group, and showed no clinically significant effects on hemostasis, haematologic
parameters, thyroid and adrenal, liver functions, glucose and lipid metabolism, or
electrolyte balance, with maintainance of mean high-density lipoprotein-3 choles-
terol from the baseline.

The morphological studies reported inhibition of endometrial cells prolifera-
tion, by down-regulation of ESR2/ESR1 ratio [126] and aromatase expression [127]
with local estrogen synthesis reduction, and the inhibition of human endometrial
stomal cells proliferation, together with functional changes, as it is prolactin syn-
thesis- a typical marker for decidualisation [124] and the association of increased
apoptosis in endometriotic lesions [128]. DNG increases the PR-B/PR-A ratio in
ovarian endometriosis [126] and down-regulates the expression of CYP19A1, and
inflammatory, and neuroangiogenesis factors through PR- A and B- isoforms [129].
Some studies have revealed that DNG reverses some alterations of the immune sys-
tem by increasing natural killer cells in the peritoneal fluid, and spleen, parallel to
the decrease of peritoneal fluid cellular content, and lower peritoneal macrophages synthesis of inflammatory cytokine IL-1β [118], and inhibits IL-1β release by the endometriotic epithelial cells [130].

4.1.4 Extended regimen of continuous combined hormonal contraceptives in endometriosis

The continuous regimen or extended cycle vs. cyclic use of combined hormonal contraceptives was first proposed by Loudon [131] in a family planning clinic from Edinburgh (UK) by skipping the tablet-free interval of 7 or 4 days. The initial proposed non-traditional regimen was with 84 active pill with ethyl estradiol 50 μg plus lynestrenol 250 μg/day and 7 days free, and it obtained a great adherence in women suffering of endometriosis, according to suppression of withdrawal bleeding, and inducing atrophy in ectopic and eutopic endometrium, as it is recognized by Cochrane Database Syst Rev., 2014 [132]. The regimen was recommended with levonorgestrel – LNG (90 or 100 μg/day), drosiprenone- DRS (3 mg/day) [133, 134] desogestrel (150 μg/day) [135], NETA (1000 μg/day) [136], and recently dienogest (2 mg/day) [137] as progestins, associated to 20 or 30 μg/day of ethinyl estradiol. The regimen of ethinyl estradiol 20 μg plus LNG 90 μg/day was reported for 364 days at Eastern Virginia Medical School, Norfolk (USA) [138] and in Italy during 6 months with ethinyl estradiol 20 μg plus LNG 100 μg/day [139], or during 3 to 6 months in Germany, Frankfurt University [140, 141], and recently the Italian prospective open label study [137] analyzed ethinyl estradiol 30 μg/day plus dienogest 2 mg/day in a continuous regimen vs. 21/7 to assess quality of life and sexual function in women with endometriosis. The pharmaceutical industry created some drugs combining ethinyl estradiol 30 μg/LNG 150 mg/day for 84 days plus 7 days placebo, or the combination of ethinyl estradiol 30 μg/LNG 150 mg/day for 84 days plus ethinyl estradiol 10 μg/day for 10 days, or of ethinyl estradiol 20 μg/LNG 150 mg/day for 84 days plus ethinyl estradiol 10 μg/day for 10 days [142] after the concept called “tricycle regimen” or “tricycling”, first named so in Sweden in 1993 very easy to be followed, and very well appreciated by users [135].

The literature describes some attempts to shorten free hormones interval for the reduction of hormone withdrawal symptoms, as pelvic pains, headaches [143] and this regimen used LNG, norethindrone (1 mg/day), desogestrel, and norgestimate with ethinyl estradiol 35 μg or less/day. All these attempts were followed by better results of pelvic pain control, but after a longer duration of administration (as after 6 months of norethindrone).

One must consider the tricycle regimen of contraceptives in adolescents suffering of menorrhagia, dysmenorrhea, or with family history of endometriosis, in order to reduce the concerns of a future progress of an eventually later diagnosed endometriosis, existing already findings on this category of women, at the Department of Adolescent Medicine of the University of Pitsburg (USA), with a significant ovarian suppression, endometrial atrophy, no metabolic change, and adolescents safety [144]. At this study one must add a recent one [145] sustaining that women who used the new generation of COC or only progestins have lower circulatory inflammatory biomarkers [IL-6, sTNFR2 (soluble tumor necrosis factor α receptor 2), and in a lower degree C-reactive protein] similar to the effect of a higher number of lifetime ovulatory years. The presence of progesterone after ovulation, the new generation of COCs, and the effects of progestins can be the explanation for the benefits in reducing systemic inflammation; being considered that chronic inflammation reduces ovulation rate in premenopausal women, and ovulation and menstruation increase local inflammation.
There are recent reviews on the comparison of flexible/extended COCs to cyclic COC use in endometriosis, published in 2018 in USA (on dysmenorrhea, non-pelvic pains, dyspareunia [146]), or in Europe [147] and the systematic review (2019) of only 8 RTC published between 1934 and 2018 (of a total 743 studies) on dysmenorrhea [148] and in 2021 [79]. It is shown a statistically significant, though modest, improvement in dysmenorrhea - a reduction of 50% when COCs were given for four months compared to placebo [147] and a shorter duration of only 4 days of the pains, with conflicting results on interference with daily activity, pain severity, and pain recurence [148].

4.1.5 Different routes of administration for progesterone/progestins to increase positive effects in alleviation progesterone attenuation/resistance and estrogen dominance

Since many years one discusses non-conventional routes for drug administration in order to avoid the second liver passage, and to increase bioavailable active substance where is necessary. Vaginal route for progesterone and medicated intrauterine devices are most analyzed for their beneficial endometrial effects, without or with less systemic effects of progesterone/progestins.

4.1.5.1 Vaginal route for micronized progesterone

Natural Progesterone has far more anti-inflammatory properties, fewer side effects, is very versatile in how it can be used, and the micronization of P4 is very important, and the oral micronized progesterone capsules were re-directed to be used vaginally [149]. Micronized P4 has a more selective effect on PRs, and results in less interaction with androgenic and mineral-corticoid receptors compared with progestins.

The vaginal route for P4 was proposed since many years ago [150] but the new hypothesis regarding the higher endometrial P4 levels than that obtained after intravenous administration was presented by Cicinelli and de Ziegler [151]. This phenomenon of preferential uterine distribution after vaginal administration was named “first uterine pass effect” [38] or “uterine specificity of vaginal progesterone” [152]. The high uterine level of P4 vaginally administered is explained by various putative modes of transport including direct diffusion through tissue, intracervical aspiration, absorption into the venous or lymphatic circulatory systems and countercurrent vascular exchange with diffusion from utero-vaginal veins/lymphvessels to arteries. While the serum P4 concentration is often low or “sub-physiological”, endometrial effects show in most cases clear and complete secretory changes, or pseudodecidual aspects, or atrophy after long duration of therapy [153]. In USA at Brigham and Women’s Hospital [154] one recommends vaginal gel, cream, or suppositories, in a higher dosage for endometriosis control than that for replacement therapy recommended in menopause, such as 300 mg twice a day for minimum 3 months, then 300 mg at bed-time.

4.1.5.2 Medicated intrauterine systems to restore estrogen-progesterone balance

The direct application of progesterone or a progestin (usually LNG), or of a SPRM (ulipristal acetate was proposed) was another potential option to control estrogen dominance in eutopic endometrium, for avoidance irregular effects of systemic progestins, considering a consistent inhibition of ERs expression in eutopic endometrium [107, 155] with pain reduction, or to avoid pain relapse in
postoperative period, and a total patients’ satisfaction when compared to their attention for daily pills intake [156].

4.2 New therapeutic strategies to restore estrogen-progesterone endometrial balance in endometriosis. Selective estrogen receptor modulators (SERMs). Selective progesterone receptor modulators (SPRMs)

Medical literature presents under the name of selective estrogen receptors modulators (SERMs) a relative new category of drugs aimed to target down-regulation of E2 signaling, by a direct binding to ERs in a tissue specific manner [91]. The studies on rat induced endometriosis [157] have shown the reduction of endometriotic lesions by down-regulation of ESR1 and cell proliferation by bazedoxifene or raloxifene, but in a RCT on administration of raloxifene (180 mg/day) after laparoscopically apparent complete excision of ectopic lesions, it was recorded a shortly time for chronic pain relapse with raloxifene vs. placebo, but without recurrence of endometriotic lesions at the second laparoscopy [158] and these results have reduced the enthusiasm for first and second generation of SERMs in endometriosis. The third generation of SERMs administration - basedoxifene, reduced glands size and number of ectopic sites in mice [159].

The selective progesterone receptor modulators (SPRMs) are known since long time, with the antiprogestin representative RU 436 or mifepristone. SPRMs were proposed to treat unresponsive cases to progestin treatment, due to progesterone resistance dilemma [160] connected to their direct interaction to progesterone receptors, in order to reduce estrogen- induced cells proliferation and prosta-glandins production [161]. Two old multicentres trials on mifepristone showed its efficacy in the endometriosis chronic pain control, with lesions size reduction, although results are mixed [162, 163].

Ulipristal acetate suppressed ectopic endometrium induced in mice, with slow reappearance after discontinuation [100]. After many discussions on the promising evidence of inhibiting human endometrial cell proliferation in vivo [164] and on the so-called progesterone receptor modulator-associated endometrial changes (PAEC) recorded after 6 months of therapy, which is reversible after ulipristal discontinuation [164] the liver life threatening complications induced the discontinuation of the study on endometriosis effects [165].

Per global these so called “new” classes of molecules with selective receptors of steroid hormones modulation objectives did not covered the expectations in endometriosis, according to women pathologic condition progressively induced on their steroid hormones receptors dysregulation/loss.

4.3 New potential therapeutic perspectives: A combination therapy using agonist of ERβ and the SRC-1 isoform as the next generation of endometriosis therapy to restore estrogen-progesterone signaling balance

Long duration of systemic hypoestrogenism may affect brain, heart, and bones, in young women, and for non responders to first line therapy mentioned above, long term medication with GnRH agonists/ antagonists or aromatase inhibitors cannot be recommended, fact that imposed more researches to try to improve estrogen dominance/progesterone resistance, to stop ectopic tissue growth and disease stage progression with chronic pelvic pains, and infertility. These aims were objectives at the Baylor College of Medicine, Houston (USA) where it was proposed a combination therapy using an agonist of ER β and the steroid receptor coactivator-1 (SRC 1) isoform as the next generation of endometriosis therapy – Han et al. [44] (Figure 2).
The combination of ER$\beta$-selective agonist and SRC-1 is based on the significant suppressive effect of PHTPP on ectopic lesion growth by inhibiting ER$\beta$ activity in mice induced endometriosis [166] associated to minimum side effects on ER$\alpha$, without influences on eutopic endometrium or negative influences on mice fertility. SRC-1 is considered a PR co-activator, and indirectly through it, ER$\beta$ may impair PR-mediated signaling in the ectopic lesions, because actually it is no evidence available to show that PR could potentially interact directly with ER$\beta$ [167]. These drivers combination allows marked suppression of ectopic lesion growth compared with either individual agents alone, both demonstrating essential roles in early stages of disease pathogenesis [168] specially on apoptosis modulation and inflammation reduction.

4.4 Potential therapeutic options for estrogen dominance control

Several medical treatments for endometriosis directly aim to reduce E2 production or action in order to mitigate E2 dominant conditions, and actually one discusses GhRh agonists, and the new class of GhRh antagonists. These therapies are efficient in cases wishing to conceive, and for the control of chronic pelvic pains- menstrual and non-menstrual [169] but associated hypoestrogenism is cause of many health concerns, requiring hormone “add-back” for long term use [91]. Numerous studies included in the Cochrane Data Base Systematic Review (2003) [170] revealed since long time that these effects limit the duration of treatment, which cannot be administered without “add back”/hormone-replacement therapy [171] and treatment cannot be dose-adjusted to alleviate the side effects - Practice Committee of the American Society for Reproductive Medicine (2014) [172].

The recent history of HT for estrogen dominance and progesterone resistance in endometriosis shows that the “old” injectable depot MPA, which can decrease ESR1 and ESR2 while increasing PR-A and PR-B in the eutopic and ectopic endometria [173] has equivalent efficacy to leuprolide- a GnRH agonist, in reducing pain, but with less adverse hypoestrogenic effects on bone density [174].

4.4.1 GnRH agonists. GnRH antagonists

Injectable GnRH agonists (leuprolide, diferelin, nafarelin) are normally second-line treatments. Which decrease hormone levels by down-regulating the pituitary through negative feedback mechanisms [91] and indirectly they favor an endometrial complete silenced hormonal milieu, on the bases of estrogen dominance concept. GnRH agonists have been shown to be effective in reducing endometriosis-related pain [175] with adverse effects – hot flushes, bone mineral density loss, or coronary heart disease, headaches due to a hypoestrogenic state, requiring hormone
“add-back” [91]. It was reported a mean percent decrease from baseline in bone mineral density at the lumbar spine of 3.2% at 6 months, and of 6.3% at 12 months after leuprolide alone, and these side effects were associated with rates of discontinuation by 6% because hot flushed, and 8% because of emotional changes [171].

GnRH antagonists, known since long time [176] were in last 10 years under investigation for endometriosis treatment in USA, and Europe according to their capacity to downregulate gonadotropins, without flare-ups like GnRH agonists because they rapidly and directly compete for GnRH receptors [177] and the oral administration is another benefit. Their mechanism of action is different to that of GnRH agonists. After an initial stimulatory phase desensitize GnRH receptors in the pituitary, causing a subsequently depletion of pituitary gonadotropins and full suppression of E2 to levels that are equivalent to those associated with bilateral oophorectomy [178].

Based on Barbieri RL (1992)” estrogen threshold hypothesis” [179] the 2 large multicenters RCTs (Australia, Brazil, Canada, New Zeeland, Poland, USA) [177] have concluded that the new oral Elagolix, a nonpeptide GnRH antagonist, in 2 different doses (one with Elagolix 150 mg/day, and the other RCT with Elagolix 100 mg twice/day) significantly lower scores for dysmenorrhea and non-menstrual pelvic pain than placebo after 3 and respectively 6 months of treatment, and dyspareunia at 3 months, with significantly better results in cases on 200 mg/day with respect to the use of rescue analgesic agents at 3 months and 6 months, dyspareunia at 3 months, and rescue opioid use at 3 months less than did those receiving placebo. The dysmenorrhea control was better than that of non-menstrual pelvic pains, dysmenorrhea is mostly dependent on cyclic changes in ovarian hormones, whereas the mechanism of non-menstrual pelvic pain are considered more complex [180]. The results allowed the researchers to consider that complete estrogen supression may not be needed to controll endometriosis-associated pain, and estrogen may be adjusted to a level that is adequate to control pain with minimum hypoestrogenic effects (hot flushes, bone mineral density, lipid levels), more frequently claimed in the higher dosage, similar to those induced by GnRH agonists. Elagolix was associated with an antiproliferative effect at each dose, and with endometrial atrophy at higher dose, which was consistent with decreases in endometrial thickness. AI have significant adverse effects like irregular bleeding, and joint pains [185].

4.4.2 Aromatase inhibitors

Aromatase inhibitors (AI), anastrozole and letrozole - the most known derivates, were destinated for control of local E2 levels in many pathologies, and in endometriosis they induce regression of peritoneal lesion size, in a higher degree than MPA [182] by reducing androgen aromatization into estrogens, both in adipose tissue and within endometriotic sites. These drugs are inhibiting endometrial progenitor cells migration to ectopic sites [183], and by increasing apoptosis, and diminishing endometrial VEGF and PGE2 in a mouse model [184] they reduce chronic pelvic pains. AI have significant adverse effects like irregular bleeding, and joint pains [185].
AI action is explained by the promotion of pituitary gonadotropin hormones release, with consequent ovarian stimulation; therefore, they must be combined with a progestin or other method of gonadotropin inhibition to treat endometriosis in premenopausal women [186]. There was compared the association of letrozole (2,5 mg/day) to NETA (2,5 mg/day) vs. NETA (2,5 mg/day) alone, and the Italian open-label study proved a better control for pain and dyspareunia of the drugs association in rectovaginal endometriosis [187] with a better sizes reduction of rectovaginal nodules [188] and of endometrotic ovarian cysts [189]. The effect of the combination of a progestin to an AI is not lasting after termination of HT, so Reis et al. [7] in their published review on progesterone ligands consider letrozole as a second therapeutic-line for selected patients, who fail to respond to first line HT- progesterone/progestins.

5. Potential therapeutic perspectives to restore the estrogen-progesterone receptors balance in endometriosis

The dysregulation of ER and PR in endometrial glandular epithelium and stroma is for sure a pathological mechanism involved in eutopic and ectopic endometrium in women suffering of endometriosis, a disease with onset during embryofetal life, and long time duration, sometimes up to death. The prevention and the attempts to maintain or restore the ER\textsubscript{α}/ER\textsubscript{β}, and PR-A/PR-B through their normal genes is one of new potential contemporary therapeutic options.

5.1 Genetic/epigenetic interventions in dysregulated endometrial progesterone responses in endometriosis

Progesterone resistance in endometriosis has genetic causes as PRs gene polymorphisms, altered microRNA expression, and epigenetic changes of PRs and their targets. A consequence of impaired progesterone action is that hormonal therapy is rendered ineffective for a subset of women with endometriosis. Environmental toxins as dioxin may play a role in the genesis of endometriosis by permitting an inflammatory milieu.

5.1.1 Progesterone responsive genes methylation: a cause of a dysregulated progesterone response

The studies at Yale University, New Haven, Connecticut (USA) using RT-qPCR for assessment of endometrial genes in mice induced endometriosis compared to normal mice [190] have demonstrated the silencing or inhibition of P4 target HOX genes (HOXA 10/HOXA 11, known also as genes of receptivity) by promoter hypermethylation - an epigenetic mechanism, which favorizes lack of menstrual cycle variation of PRs distribution, which is proper in normal menstruated women, and this is appreciated as a partial explanation for the refractoriness of some endometriotic lesions to progesterone/progestin therapy [7, 191]. Altered PRs expression or diminished activity may lead to attenuated or dysregulated P4 response in ectopic endometrium, and decreased expression of P4 responsive genes including HOX genes in the eutopic endometrium. Cakmak and Taylor [191] and Lee et al. [190] concluded that normal endometrium placed in an ectopic location, in order to create experimental endometriosis led to characteristic changes in gene expression of eutopic endometrium, fact that was previously observed regrading stromal stem cells migration to ectopic sites, without this genetic/epigenetic explanation, and it was controverted. These data were suggestive for the existence of a signal
conduction pathway from endometriosis that alters endometrial gene expression through altered Pgr signaling and epigenetic programming. The relatively permanent nature of methylation may explain the widespread failure of HT [7]. One discusses about promoters methylation, being controversies if both PRs promoters are methylated, discovering only PR-B methylation, not PR-A [39] or both PRs promoters [192] or DNA methylation of the CG-islands in PR promoter, and its gene HOXA 10 [114] as there are controversies regarding ERs genes (ESR-1 and ESR-2), being hypomethylated CpG island at the ESR2 promoter region, involved in primary mechanism responsible for differential expression of ESR2, discovered to be increased in ectopic and eutopic endometria [40] while other study from Brazil [192] denies their methylation in eutopic and ectopic endometria.

5.1.2 DNA methylation inhibitor: a new therapeutic perspective/challenge to restore PR-mediated signaling molecules in endometriosis

The women's ectopic and eutopic endometrial progressive loss of PRs and ERα/PR-mediated signaling in the developing ectopic lesions during 7 to 12 years from histologic disease's onset to clinical symptoms/signs permits to health care providers to intervene in the epigenetic regulation therapeutic control.

The animal studies at the University of Illinois at Urbana-Champaign (USA) [114] have shown that a DNA methyltransferase (DNMT) inhibitor, such as Decitabine (DAC, 5-aza-2′-deoxycytidine), an analogue of deoxycytidine that can incorporate into DNA strands and cause DNA demethylation [193] compared to vehicle administration in immunocompetent mice induced endometriosis, treated also with E2 had alleviated lesion growth, and increased expressions of PR protein, and miRNA corresponding to Esr1, Pgr, Hand2, and Hoxa10, (P4 target), but not for Ccl5 and Ptgs2 in the eutopic endometrium and ectopic sites. The results of the study enable the authors to review the involvement of estrogens and progesterone in eutopic and ectopic endometrium, the epigenetic regulation including DNA methylation, the fact that ERα and ERβ (ESR1 and ESR2), PRs (PRA and B), as well as PR-targets Hoxa10, Gata2/6, and Hand2 are susceptible to DNA methyltransferases, leading to an aberrant expression of these molecules in endometriosis.

Early studies showed that the promoter sequences of ERα and ERβ (ESR1 and ESR2), PRs (PRA and B), as well as PR-targets Hoxa10, Gata2/6, and Hand2 are susceptible to DNA methyl transferases, leading to an aberrant expression of these molecules in endometrial diseases [194] and loss of PR-mediated signaling during disease progression contributes to the increased susceptibility to P4 resistance in ill women. Li et al. [114] accepted the proposed role of inflammation to provoke widespread changes in the genes and chromatin landscape of lesions, because their analyses showed that DNA methylation in PRs and Hoxa10 promoters was enhanced in the ectopic lesions in comparison to the normal endometrium, and inhibition of genome-wide DNA methylation in female mice restrained lesion expansion and partially restored target gene expression.

6. Conclusions

Endometriosis is a chronic disease, with possible onset in embryonic life, latency in childhood, reactivation from the menarche, with clinical symptoms and signs difficult to be early accurate assessed during reproductive years. Endometriosis has a progressive evolution, which drives through tissue intrinsic properties to increase lesions size and number, and to functional damages of original eutopic endometrium. The cascade of dysregulations in chromatin machinery is difficult
to be stopped by hormone therapy and surgery (associated in different strategies), when the abnormal signaling of steroid hormones has started: endometrial estrogen dominance and progesterone attenuate response up to progesterone resistance, because their genes epigenetic disorders. The 7–12 years delay in diagnosis and proper therapy drives to accentuation of chronic pelvic pains, dyspareunia, dyschezia, infertility/subfertility.

The best therapy is to avoid the dysregulation of steroid hormones signaling, induced initially in ectopic and later in eutopic endometrial stroma and glands, to maintain or to restore the estrogen/progesterone receptors natural balance, which may be possible when an early intervention, from the menarche. One may discuss about timing in starting HT in adolescent girls with family history of endometriosis, precocious dysmenorrheal, menorrhagia, abundant blood and clots loss, favorising menstrual blood reflux.

First line therapy is progesterone/progestins, working in prevention of estrogen dominance and progesterone dysregulated signals, and can maintain their positive effects, when it is early administrated, as animal models have shown. One must recommend this first line therapy, for a long time duration, with a single drug (micronized progesterone, or dienogest, a new progestin) or combined with a synthetic/natural estrogen in COCs, in cyclic or in continuous/extended regimens up to response failure, with the advice to avoid drug discontinuation. Vaginal route for micronized progesterone, IUD with progesterone/levonorgestrel may work better for alleviation of estrogen dominance and progesterone resistance.

GnRH agonists/antagonists- second line therapy, are active in correcting the steroid imbalance, but less than our expectances, even with the new oral molecules (as elagolix), or combination of molecules [GnRH antagonists (relugolix or elagolix) plus an estrogen, or plus an estrogen and a progestin], which were proposed to avoid or to reduce add back therapy. The most recent studied molecule of elagolix may be used no longer than 24 months, because a longer administration may impose add back therapy.

Aromatase inhibitors are also second line therapy, recommended in selected patients for refractory endometriosis, with chronic pelvic pains. Letrozole combination to norethindrone acetate is better than the progestin alone in correcting estrogen dominance, and progesterone resistance.

The new therapeutic perspectives regarding the combination of ER agonists with co-activators, and DNA methylation inhibitors are still in studies at high level technology laboratories, not for current medical use.

**Conflict of interest**

None.
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References

[1] Signorile PG, Baldi F, Bussani R, Baldi A, et al – Embryologic origin of endometriosis: analysis of 101 human female fetuses. J Cell Physiol, 2012; 227 (4):1653-6; doi: 10.1002/jcp.22888.

[2] Crispi S, Piccolo MT, D’Avino A, Signorile PG, et al – Transcriptional profiling of endometriosis tissues identifies genes related to organogenesis defects Journal of Cellular Physiology; 2013; 226(9): 1927- 1937 DOI: 10.1002/jcp.24358

[3] Brosens I, Gordts S, Benagiano G– Endometriosis in adolescents is a hidden, progressive and severe disease that deserves attention, not just compassion. Hum Reprod.; 2013; 28: 2026-2031.

[4] Secosan C, Balulescu L, Brasoveanu S, Balint O, Pirtea P, Dorin G, et al– Endometriosis in menopause- renewed attention on a controversial disease Diagnostics (Basel), 2020, 10:134. doi 103390/diagnostics10030134

[5] McKinnon B, Mueller M, Montgomery G– Progesterone resistance in endometriosis: an acquired property? Trends EndocrinolMetab. 2018 Aug;29(8):535-548. doi: 10.1016/j.tem.2018.05.006

[6] Al-Sabbagh M, Lam EW, Brosens JJ– Mechanisms of endometrial progesterone resistance. Mol Cell Endocrinol. 2012; 358: 208-215

[7] Reis MF, Coutinho ML, Vannuccini S, Batteux F, Chapron Ch, Petraglia F – Progesterone receptor ligands for the treatment of endometriosis: the mechanisms behind therapeutic success and failure Human Reproduction Update, July-August 2020, 26 (4) 565-585, https://doi.org/10.1093/humupd/dmaa009

[8] Grümmér R, Traub O, Winterhager E – Gap junction connexin genes cx26 and cx43 are differentially regulated by ovarian steroid hormones in rat endometrium. Endocrinology 1999; 140:2509-2516

[9] Warren LA, Shih A, Renteira SM, Sechkin T, Gregersen PK, et al – Analysis of menstrual effluent: diagnostic potential for endometriosis. Mol Med, 2018; 24 (1), 1; doi: 10.1186/s10020-018-0009-6.

[10] Kondi-Pafiti A, Spanidou-Carvouni H, Papadias K, et al. – Malignant neoplasms arising in endometriosis: clinicopathological study of 14 cases. Clin Exp Obstet Gynecol. 2004;31(4):302-304.

[11] Wheeler JM – Epidemiology of endometriosis-associated infertility. J Reprod Med; 1989; 34 41 –46

[12] Giudice LC, Kao LC – Endometriosis. Lancet. 2004;364(9447):1789-1799

[13] Du H, Taylor SH – The Role of Hox Genes in Female Reproductive Tract Development, Adult Function, and Fertility. Cold Spring Harb Perspect Med 2015 Nov , 6 (1), a023002. doi: 10.1101/cshperspect.a023002.

[14] Olive DL, Pritts EA – Treatment of endometriosis. N. Engl J Med;2001; 345: 266-275.

[15] Giudice LC – Clinical practice. Endometriosis. N Engl J Med. 2010; 362(25):2389-2398

[16] Johnson NP, Hummelshoj L – World Endometriosis Society Montpellier Consortium. Consensus on current management of endometriosis. Hum Reprod 2013;28:1552-68.

[17] Patel B, Elguero S, Thakore S, Dahoud W, Bedaiwy M, Mesiano S
- Role of nuclear progesterone receptor isoforms in uterine pathophysiology. Hum. Reprod Update. 2015; 21: 155–73

[18] Tuttles F, Keckstein J, Ulrich U, Possover M, Tinnerberg HP, et al – ENZIAN-Score, a classification of deep infiltrating endometriosis. 2005, Zentralt Gynakol, 127(5):275-81

[19] Haas D, Shielb O, Shamiyeh, Oppelt P – The rASRM score and the Enzian classification for endometriosis. Their stengths and weaknesses. Acta Obstet Gynecol. Scand 2012 oct, 92 (1). doi: 10.1111/aogs.12026

[20] Keckstein J, Saridogan E, Ulrich AU, Sillem M, Hudelist G, et al– The ENZIAN classification; A comprehensive non-invasive description system for endometriosis. Acta Obstetricia et Gynecologica Scandinavica, 2021, 100(7), 1165-1175 doi. Org/10.1111/aogs,14099

[21] Starzinski-Powitz A, Zeitvogel A, Schreiner A, Baumann R – In search of pathogenic mechanisms in endometriosis: the challenge for molecular cell biology. Curr Mol Med; 2001; 1 (6): 655-64.

[22] Forte A, Cipollaro M, Galderisi U – Genetic, epigenetic and stem cell alterations in endometriosis: new insights and potential therapeutic perspectives Clin Sci (Lond); 2014; 126(2):123-38. doi: 10.1042/ CS20130099.

[23] Brinton LA, Gridley G, Persson I, Baron J, Bergqvist A– Cancer risk after a hospital discharge diagnosis of endometriosis. Am J Obstet Gynecol. 1997 Mar;176(3):572-9. doi: 10.1016/ s0002-9378(97)70550-7.

[24] Ness RB, Cramer DW, Goodman MT, Wu HA, et al – Infertility, fertility drugs, and ovarian cancer: a pooled analysis of case-control studies. Am J Epidemiol. 2002 Feb 1;155(3):217-24. doi: 10.1093/aje/155.3.217.

[25] Sakr S, Naqvi H, Komm B, Taylor SH– Endometriosis Impairs Bone Marrow-Derived Stem Cell Recruitment to the Uterus Whereas Bazedoxifene Treatment Leads to Endometriosis Regression and Improved Uterine Stem Cell Engraftment. Endocrinology; 2014 Apr; 155(4): 1489-1497

[26] Surrey E, Carter MC, Soliman MA, Khan S, Snabes CM, et al – Patient-completed or symptom-based screening tools for endometriosis: a scoping review. Arch Gynecol Obstet. 2017 Aug; 296 (2):153-165. doi: 10.1007/ s00404-017-4406-9.

[27] Klemmt PA, Carver JG, Koninckx P, Mardon HJ, et al – Endometrial cells from women with endometriosis have increased adhesion and proliferative capacity in response to extracellular matrix components: towards a mechanistic model for endometriosis progression. Hum. Reprod. 2007; 22(12):3139-3147; doi: 10.1210/jc.2009-0207

[28] Burney OR, Talbi S, Hamilton EA, Chi V, Giudice CL, et al – Gene expression analysis of endometrium reveals progesterone resistance and candidate susceptibility genes in women with endometriosis. Endocrinology. 2007;148:3814-3826; doi:10.1210/en.2006-1692

[29] Jones JP C, Inuwa MI, Nardo GL, Litta P, Fazleabas TA– Eutopic Endometrium From Women With Endometriosis Shows Altered Ultrastructure and Glycosylation Compared to That From Healthy Controls— A Pilot Observational Study. Reprod. Sci 2009 Jun, 16 (6):559- 572 doi: 10.1177/1933719093328285

[30] Patel BG, Rudnicki M, Yu J, Shu Y, Taylor RN – Progesterone resistance in endometriosis: Origins, consequences and interventions. Acta Obstet. Gynecol. Scand. 2017;96:623-632. doi: 10.1111/aogs.13156
[31] Attia GR, Zeitoun K, Edwards D, Johns A, Carr BR, Bulun SE – Progesterone receptor isoform A but not B is expressed in endometriosis. Journal of Clinical Endocrinology and Metabolism. 2000; 85(8):2897-2902

[32] Burns KA, Rodriguez KF, Hewitt SC, Korach KS, et al – Role of estrogen receptor signaling required for endometriosis-like lesion establishment in a mouse model. Endocrinology. 2012; 153; 3960-3971. doi:10.1210/en.2012-1294

[33] Chrousos GP, MacLusky NJ, Brandon DD, Lipsett MB, et al – Progesterone resistance. Adv Exp Med Biol. 1986;196:317-328. doi: 10.1007/978-1-4684-5101-6_21

[34] Keller DW, Wiest WG, Strckler RC, et al – Pseudocorpus Luteum Insufficiency: A Local Defect of Progesterone Action on Endometrial Stroma. J. Clin. Endocrinol. Metab. 1979:48; (1):127-132

[35] Gellersen B, Brosens IA, Brosens JJ – Decidualization of the human endometrium: mechanisms, functions, and clinical perspectives. Semin Reprod Med; 2007; 25: 445-453

[36] Vasquez YM, DeMayo FJ – Role of nuclear receptors in blastocyst implantation. Semin.Cell.Dev.Biol. 2013;24:724-735. doi: 10.1016/j.semcdb.2013.08.004

[37] Plante BJ, Lessey BA, Taylor RN, Wang W, Bagchi MK, Yuan, et al – G protein-coupled estrogen receptor (GPER) expression in normal and abnormal endometrium. Reprod Sci. 2012; 19(7):684-693. doi: 10.1177/1933719111431000

[38] Groothuis PG, Dassen HH, Romano A – Estrogen and the Endometrium lessons learned from gene expression profiling in rodents and human. Hum Reprod. Update; 2007;13:405-417

[39] Wu Y, Strawn E, Basir Z, Halverson G, Guo SW – Promoter hypermethylation of progesterone receptor isoform B (PR-B) in endometriosis. Epigenetics. 2006;1:106-111. doi: 10.4161/epi.1.2.2766

[40] Xue Q, Lin Z, Cheng YH, Huang CC, Marsh E, Bulun SE, et al – Promoter methylation regulates estrogen receptor 2 in human endometrium and endometriosis. Biol Reprod. 2007;77(4):681-687

[41] Bulun SE, Monsavais D, Pavone ME, Attar E, et al – Role of estrogen receptor-beta in endometriosis. Seminars in reproductive medicine. 2012; 30(1): 39-45.

[42] Hewitt SC, Korach KS – Estrogen Receptors: New Directions in the New Millennium. Endocr. Rev. 2018;39:664-675. doi: 10.1210/er.2018-00087

[43] Mendell JT – MicroRNAs critical regulators of development, cellular physiology and malignancy. Cell Cycle. 2005; 4(9):1179-1184

[44] Han SJ, Jung SY, Wu SP, Hawkins SM, Tsai MJ, et al – Estrogen Receptor beta Modulates Apoptosis Complexes and the Inflammasome to Drive the Pathogenesis of Endometriosis. Cell. 2015;163:960-974. doi: 10.1016/j.cell.2015.10.034

[45] Bulun SE, Cheng YH, Pavone ME, Xue Q, Kim JJ, et al – Estrogen receptor-β, estrogen receptor-α, and progesterone resistance in endometriosis. Semin Reprod Med. 2010;28 (01):036-043; doi: 10.1055/s-0029-1242991

[46] Qi QM, Guo SW, Liu XS – Estrogen biosynthesis and its regulation in endometriosis. Reprod Dev Med. 2017;1(1):55. doi: 10.4103/2096-2924.210698

[47] Simmen RC, Kelley AS – Reversal of fortune: estrogen receptor-β in
endometriosis. J Mol. Endocrinol. 2016; 57(2): F23–F27. doi: 10.1530/JME-16-0080.

[48] Acien P, Velasco I, Gutierrez M, Martinez-Beltran M – Aromatase expression in endometriotic tissues and its relationship to clinical and analytical findings. Fertility and sterility. 2007; 88(1):32-8.

[49] Ferrero S, Remorgida V, Maganza C, Venturini P, et al – Aromatase and endometriosis: estrogens play a role. Annals of the New York Academy of Sciences. 2014; 1317:17-23.

[50] Shafrir AL, Missmer SA – Towards subtypes- deep endometriosis oestrogen receptor α expression Nat. Rev. Endocrinol. 2020 Oct; 16(10):541-542. doi: 10.1038/s41574-020-0394-0

[51] Pluchino N, Mamillapalli R, Wenger J-M, Ramyead L, Taylor SH, et al – Estrogen receptor-α immunoreactivity predicts symptom severity and pain recurrence in deep endometriosis. Fertil Steril., 2020 Jun;113(6):1224-1231.e1. doi: 10.1016/j.fertnstert.2020.01.036

[52] Dimitriadis E, Stoikos C, Stafford-Bell M, Clark I, Salamonsen LA, et al – Interleukin-11, IL-11 receptor alpha and leukemia inhibitory factor are dysregulated in endometrium of infertile women with endometriosis during the implantation window. J. Reprod. Immunol. 2006; 69;53-64. doi: 10.1016/j.jri.2005.07.004.

[53] Kao LC, Germeyer A, Tulac S, Lobo S, Giudice LC, et al – Expression profiling of endometrium from women with endometriosis reveals candidate genes for disease-based implantation failure and infertility. Endocrinology. 2003;144(7):2870-2881

[54] Osteen KG, Bruner-Tran KL, Eisenberg E – Reduced progesterone action during endometrial maturation: a potential risk factor for the development of endometriosis. Fertil Steril. 2005; 83(3): 529-537.

[55] Arici A, Engin O, Attar E, Olive DL – Modulation of leukemia inhibitory factor gene expression and protein biosynthesis in human endometrium. J. Clin. Endocrinol. Metab. 1995; 80:1908-1915.

[56] Kelleher AM, Behura SK, Burns GW, Young SL, DeMayo FJ, Spencer TE – Integrative analysis of the forkhead box A2 (FOXA2) cistrome for the human endometrium. FASEB J. 2019 Jul; 33(7):8543-8554

[57] Nisolle M, Casanas-Roux F, Wyns C, Donnez J, et al – Immunoistochemical analysis of estrogen and progesterone receptors in endometrium and peritoneal endometriosis: a new quantitative method. Fertil. Steril. 1994;62(4):751-759. doi: 10.1016/S0015-0282(16)57000-9.

[58] Shao R, Cao S, Wang X, Feng Y, Billig H – The elusive and controversial roles of estrogen and progesterone receptors in human endometriosis. Am J Transl Res. 2014; 6(2):104.

[59] Colón-Caraballo M, García M, Mendoza A, Flores I – Human endometriosis tissue microarray reveals site-specific expression of estrogen receptors, progesterone receptor, and Ki67. ApplImmunohistochemMolMorphol. 2019;27(7):491-500. doi: 10.1097/PAI.0000000000000663

[60] Flores VA, Vanhie A, Dang T, Taylor HS – Progesterone receptor status predicts response to progestin therapy in endometriosis. J Clin Endocrinol Metab. 2018;103 (12): 4561-4568. doi: 10.1210/jc.2018-01227.

[61] Nothnick WB – MicroRNAs and Endometriosis: Distinguishing Drivers from Passengers in Disease Pathogenesis.Semin Reprod Med; 2017; 35(2):173-180. doi: 10.1055/s-0037-1599089
[62] McEwan IJ – Nuclear receptors: one big family. Methods in Molecular Biology. 2009; 505 3-18. Doi. org/10.1007/978-1-60327-375-0_1

[63] Mote PA, Balleine RL, McGowan EM, Clarke CL – Heterogeneity of progesterone receptors A and B expression in human endometrial glands and stroma. Hum Reprod. 2000 Aug;15Suppl 3:48-56. doi: 10.1093/humrep/15.suppl_3.48.

[64] Brichant G, Nervo P, Albert A, Munaut C, Foidart JM, Nisolle M – Heterogeneity of estrogen receptor α and progesterone receptor distribution in lesions of deep infiltrating endometriosis of untreated women or during exposure to various hormonal treatments. Gynecol Endocrinol. 2018; 34(8):651-655

[65] Lee B, Du H, Taylor HS – Experimental murine endometriosis induces DNA methylation and altered gene expression in eutopic endometrium. Biol Reprod. 2009; 80(1):79-85

[66] Han SJ, O’Malley BW – The dynamics of nuclear receptors and nuclear receptor coregulators in the pathogenesis of endometriosis. Hum Reprod Update. 2014 Jul-Aug; 20(4):467-84.

[67] Tranguch S, Smith FD, Dey KS – Progesterone receptor requires a co-chaperone for signalling in uterine biology and implantation. Reprod. Biomed. Online. 2007; 14 (Spec No 1): 39-48

[68] Mishra A, Galvankar M, Singh N, Modi D – Spatial and temporal changes in the expression of steroid hormone receptors in mouse model of endometriosis. J. Assist Reprod. Genet, 2020 May;37(5):1069-1081. doi: 10.1007/s10815-020-01725-6

[69] Usadi RS, Lessey BA, Kowalik AI, Meyer WR, Fritz MA – The effects of low luteal phase progesterone concentrations on the histologic and functional characteristics of the endometrium. J. Soc. Gynecol. Investig. 2003; 10 (Suppl 2):386A

[70] Szwarc MM, Lydon JP, O’Malley BW – Steroid receptor coactivators as therapeutic targets in the female reproductive system. J. Steroid Biochem. Mol. Biol. 2015; 154:32-38. doi: 10.1016/j.jsbmb.2015.06.010

[71] Han SJ, Jeong J, Demayo FJ, Xu J, Tsai SY, O’Malley BW, et al – Dynamic cell type specificity of SRC-1 coactivator in modulating uterine progesterone receptor function in mice. Mol. Cell. Biol; 2005;25:8150-8165. doi: 10.1128/MCB.25.18.8150-8165.2005

[72] Kommagani R, Szwarc MM, Kovanci E, Gibbons WE, Lydon JP, et al – Acceleration of the glycolytic flux by steroid receptor coactivator-2 is essential for endometrial decidualization. PLoS Genet. 2013;9:e1003900. doi: 10.1371/journal.pgen.1003900

[73] Mukherjee A, Soyal SM, Fernandez-Valdivia R, Gehin M, O’Malley BW, et al – Steroid receptor coactivator 2 is critical for progesterone-dependent uterine function and mammary morphogenesis in the mouse. Mol. Cell. Biol. 2006;26:6571-6583. doi: 10.1128/MCB.00654-06

[74] Jeong JW, Lee KY, Han SJ, Aronow BJ, Lydon JP, DeMayo FJ et al – The p160 steroid receptor coactivator 2, SRC-2, regulates murine endometrial function and regulates progesterone-independent and -dependent gene expression. Endocrinology. 2007; 148:4238-4250. doi: 10.1210/en.2007-0122

[75] Wykes CB, Clark TJ, Khan KS – Accuracy of laparoscopy in the diagnosis of endometriosis: a systematic quantitative review. BJOG. 2004; 111
[76] Hori Y, Committee SG – Diagnostic laparoscopy guidelines: this guideline was prepared by the SAGES guidelines committee and reviewed and approved by the Board of Governors of the society of American gastrointestinal and endoscopic surgeons (SAGES), November 2007. Surg Endosc. 2008;22(5):1353-1383. doi: 10.1007/s00464-008-9759-5.

[77] Wykes CB, Clark TJ, Chakravati S, et al – Efficacy of laparoscopic excision of visually diagnosed peritoneal endometriosis in the treatment of chronic pelvic pain. Eur J Obstet Gynecol Reprod Biol. 2006;125(1):129-133. doi: 10.1016/j.ejogrb.2005.08.008.

[78] Burney RO, Giudice LC – Pathogenesis and pathophysiology of endometriosis. Fertil Steril. 2012;98(3):511-519. doi: 10.1016/j.fertnstert.2012.06.029.

[79] Donnez J, Dolman M-M – Endometriosis and Medical Therapy: From Progestogens to Progesterone Resistance to GnRH Antagonists: A Review. Journal of Clinical Medicine 2021, 10, 1085. https://doi.org/10.3390/jcm10051085

[80] Vercellini P, Viganò P, Somigliana E, Fedele L – Endometriosis: pathogenesis and treatment. Nature reviews Endocrinology. 2014;10(5):261-75. doi: 10.1038/nrendo.2013.233

[81] Zhao L, Gu C, Ye M, Han M, et al – Identification of global transcriptome abnormalities and potential biomarkers in eutopic endometria of women with endometriosis: a preliminary study. Biomed Rep. 2017; 6(6):654-662. doi: 10.3892/br.2017.902

[82] Evers J, Dunselman G, Land J, Bouckser F– Is there a solution for recurrent endometriosis? Br J Clin Pract Suppl 1991, 72, 45-53

[83] Somigliana E, Vercellini P, Viganò P, Benaglia L, Busnelli A, Fedele L– Postoperative medical therapy after surgical treatment of endometriosis: from adjuvant therapy to tertiary prevention. J Minim Invasive Gynecol. 2014;21(3):328-334.

[84] Chen I, Velth V, Choudry AJ, Murji A, Maas J, et al – Pre- and postsurgical medical therapy for endometriosis surgery, 2020 Nov 18;11(11): CD003678. doi: 10.1002/14651858.CD003678.pub3.

[85] Hou Z, Mamillapalli R, Taylor HS– Predictive biomarkers may allow precision therapy of endometriosis. J. Endometr. Pelvic. Pa. 2017; 9:279-285. doi: 10.5301/jeppd.5000311

[86] Vercellini P, Viganò P, Somigliana E, Fedele L– Endometriosis: pathogenesis and treatment. Nat. Rev. Endocrinol. 2014;10(5):261-275.

[87] Taylor HS, Giudice LC, Lessey BA, Abrao MS, Chwalisz K, et al – Treatment of endometriosis-associated pain with elagolix, an oral GnRH antagonist. N Engl J Med. 2017; 377(1): 28-40.

[88] Morelli M, Sacchinelli A, Venturella R, Mocciaro R, Zullo F – Postoperative administration of dienogest plus estradiol valerate versus levonorgestrel-releasing intrauterine device for prevention of pain relapse and disease recurrence in endometriosis patients. J. Obstet. Gynaecol. Res. 2013; 39:985-990

[89] Moreno I, Quiñones L, Catalán J, Miranda C, Roco A, Gaete K, et al– Influence of CYP3A4/5 polymorphisms in the pharmacokinetics of levonorgestrel: a pilot study. Biomedica. 2012;32:570-577

[90] Kistner RW – The use of newer progestins in the treatment of
endometriosis. Am J Obstet Gynecol 1958; 75: 264-278

[91] Ferrero S, Evangelisti G, Barra F – Current and emerging treatment options for endometriosis. Expert. Opin. Pharmacother. 2018; 19 (10):1109-1125. doi: 10.1080/14656566.2018.1494154

[92] Andrews MC, Andrews WC, Strauss AF – Effects of progestin-induced pseudopregnancy on endometriosis: clinical and microscopic studies Am J Obstet Gynecol; 1959; 78: 776-785

[93] Couzinet B, Le Strat N, Brailly S, Schaison G – Comparative effects of cyproterone acetate or a long-acting gonadotropin-releasing hormone agonist in polycystic ovarian disease. J. Clin. Endocrinol. Metab. 1996, 63:1031-1035

[94] Krattenmacher R – Drospirenone: pharmacology and pharmacokinetics of a unique progestogen. Contraception. 2000; 62: 29-38

[95] Gezer A, Oral E – Progestin therapy in endometriosis. Womens Health (Lond) 2015;11: 643-652

[96] Bono Y, Kyo S, Kiyono T, Mizumoto Y, Fujiwara H, et al – Concurrent estrogen action was essential for maximal progestin effect in oral contraceptives. Fertil Steril 2014; 101; 1337-1343

[97] Casper RF – Progestin-only pills may be a better first-line treatment for endometriosis than combined estrogen-progestin contraceptive pills Fertil Steril 2017; 107: 533-536

[98] Vercellini P, Ottolini F, Frattaruolo MP, Buggio L, Somigliana E, et al – Is shifting to a progestin worthwhile when estrogen–progestins are inefficacious for endometriosis-associated pain? Reprod Sci; 2018; 25:674-682

[99] Barra F, Grandi G, Tantari M, Scala C, Ferrero S, et al – A comprehensive review of hormonal and biological therapies for endometriosis: latest developments. Expert. Opin. Biol. Ther. 2019; 19:343-360

[100] Liang B, Wu L, Xu H, Cheung CW, Wang CC, et al – Efficacy, safety and recurrence of new progestins and selective progesterone receptor modulator for the treatment of endometriosis: a comparison study in mice. Reprod. Biol. Endocrinol. 2018, 16: 32

[101] Harada T, Momoeda M, Taketani Y, Hoshiai H, Terakawa N – Low-dose oral contraceptive pill for dysmenorrhea associated with endometriosis: A placebo-controlled, double-blind, randomized trial. Fertil. Steril. 2008, 90, 1583-1588.

[102] Vercellini P, Frattaruolo MP, Buggio L – Toward minimally disruptive management of symptomatic endometriosis: Reducing low-value care and the burden of treatment. Expert Rev. Pharmacoecon. Outcomes Res. 2017, 18, 1-4.

[103] Vercellini P – Are combined hormonal contraceptives the neglected treatment for symptomatic endometriosis? Fertil. Steril. 2018, 110

[104] Andrews WC – Hormone therapy of endometriosis -estrogen, androgens, progestin estrogens. Excerpta Med, 1976; 368:46 –59

[105] Vercellini P, Donati A, Ottolini F, Frassinetti A, Somigliana E, et al – A stepped-care approach to symptomatic endometriosis management: a participatory research initiative. Fertil. Steril 2018; 109: 1086-1096

[106] De Lignieres B, Dennerstein L, Backstrom T – Influence of route of administration on progesterone metabolism Maturitas. 1995; 21: 251-257
[107] Engemise SL, Willets JM, Taylor AH, Emembolu JO, Konje JC – Changes in glandular and stromal estrogen and progesterone receptor isoform expression in eutopic and ectopic endometrium following treatment with the levonorgestrel-releasing intrauterine system. Eur. J. Obstet. Gynecol. Reprod. Biol. 2011; 157; 101-106

[108] Beranič N, Rižner TL– Effects of progestins on local estradiol biosynthesis and action in the Z-12 endometriotic epithelial cell line. J. Steroid. Biochem. Mol. Biol. 2012; 132′: 303 – 310

[109] Bulun SE, Yilmaz BD, Sison C, Miyazaki K, Wei J, et al– Endometriosis. Endocr. Rev. 2019; 40, 1048-1079.

[110] Mori T, Ito F, Matsushima H, Takaoka O, Koshiba A, Kitawaki J, et al– Dienogest reduces HSD17β1 expression and activity in endometriosis. J Endocrinol. 2015: 225:69-76

[111] Miyashita M, Koga K, Takamura M, Izumi G, Nagai M, Osuga Y, et al – Dienogest reduces proliferation, aromatase expression and angiogenesis, and increases apoptosis in human endometriosis. Gynecol. Endocrinol, 2014: 30: 644-648

[112] Minami T, Kosugi K, Suganuma I, Yamanaka K, Kitawaki J, et al – Antiproliferative and apoptotic effects of norethisterone on endometriotic stromal cells in vitro. Eur. J. Obstet. Gynecol. Reprod. Biol., 2013;166:76-80

[113] Makabe T, Koga K, Miyashita M, Takeuchi A, Sue F, Harada M, et al – Drospirenone reduces inflammatory cytokines, vascular endothelial growth factor (VEGF) and nerve growth factor (NGF) expression in human endometriotic stromal cells. J. Reprod. Immunol. 2017; 119:44-48

[114] Li Y, Adur MK, Kannan A, Davila J, Zhao Y, LI Q, et al – Progesterone Alleviates Endometriosis via Inhibition of Uterine Cell Proliferation, Inflammation and Angiogenesis in an Immunocompetent Mouse Model. PloS One, 2016 Oct 24, 11(10), e0163347, DOI: 10.1371/journal.pone.0163347

[115] Dinh A, Sriprasert I, Williams AR, ArcherDF – A review of the endometrial histologic effects of progestins and progesterone receptor modulators in reproductive age women Contraception 2015; 91:360-367

[116] Mabrouk M, Paradisi R, Arena A, Del Forno S, Seracchioli R, et al – Short-term histopathological effects of dienogest therapy on ovarian endometriomas: in vivo, nonrandomized, controlled trial Gynecol. Endocrinol 2018; 34: 399– 403

[117] Stanczyk FZ, Hapgood JP, Winer S, Mishell DR Jr – Progestogens used in postmenopausal hormone therapy: differences in their pharmacological properties, intracellular actions, and clinical effects. Endocr Rev.2013; 34:171-208

[118] Katsuki Y, Takano Y, Futamura Y, Shibutani Y, Nozawa S, et al – Effects of dienogest, a synthetic steroid, on experimental endometriosis in rats Eur. J. Endocrinol. 1998: 138:216-226

[119] Strowtzki T, Faustmann T, Gerlinger C, Schumacher U, Ahlers C Seitz C – Safety and tolerability of dienogest in endometriosis: Pooled analysis from the European clinical study program. International Journal of Women’s Health. 7 (default):393; DOI: 10.2147/WH.S77202

[120] Angioni S, Tinelli R, Coelice V, Socolov R– New trends of progestins treatment of endometriosis Gynecological Endocrinology. August 2014 30(11): 769-773 DOI: 10.3109/09513590.2014.950646
[121] Bedaiwy AM, Allaire C, Afraraj S – Long term medical management of endometriosis with dienogest and with a gonadotropin releasing hormone agonist and add back therapy Fertility and Sterility, 2017, 107, 3, 537-548 doi: 10.1016/j.fertnstert.2016.12.024

[122] Andres M, Alves Lopez L, Baracat Chada E, Podgaec S – Dienogest in the treatment of endometriosis: systematic review. Archives of Gynecology and Obstetrics, March 2015, 292(3); DOI: 10.1007/800404-015-3681-6

[123] Strowitzki T, Faustmann T, Gerlinger C, Seitz C – Dienogest in the treatment of endometriosis-associated pelvic pain: A 12-week, randomized, double-blind, placebo-controlled study. Eur. J. Obstet. Gynecol. Reprod. Biol. 2010;151:193-198. doi: 10.1016/j.ejogr.2010.04.002

[124] Okada H, Nakajima T, Yoshimura T, Yasuda K, Kanzaki H – The inhibitory effect of dienogest, a synthetic steroid, on the growth of human endometrial stromal cells in vitro. Mol. Hum. Reprod. 2001: 7: 341-347

[125] Artini PG, Volpe A, Angioni S, et al – A comparative randomized study of three different progesterone support of the luteal phase following IVF/ET program. J Endocrinol. Invest 1995; 18:57-62.

[126] Hayashi A, Tanabe A, Kawabe S, Hayashi M, Ohmichi M, et al – Dienogest increases the progesterone receptor isoform B/A ratio in patients with ovarian endometriosis. J Ovarian Res2012, 5:31; doi: 10.1186/1757-2215-5-31

[127] Shimizu Y, Mita S, Takeuchi T, Notsu T, Kyo S, et al – Dienogest, a synthetic progestin, inhibits prostaglandin E2 production and aromatase expression by human endometrial epithelial cells in a spheroid culture system. Steroids. 2011;76:60-67. doi: 10.1016/j.steroids.2010.08.010.

[128] Miyashita M, Koga K, Takamura M, Izumi G, Osuga Y, et al – Dienogest reduces proliferation, aromatase expression and angiogenesis, and increases apoptosis in human endometriosis. Gynecol. Endocrinol, 2014: 30: 644-648

[129] Ichioka M, Mita S, Shimizu Y, Kyo S, et al – Dienogest, a synthetic progestin, down-regulates expression of CYP19A1 and inflammatory and neuroangiogenesis factors through progesterone receptor isoforms A and B in endometriotic cells. J. Steroid. Biochem. Mol. Biol, 2015;147:103– 110

[130] Mita S, Nakakuki M, Ichioka M, Shimizu Y, Kyo S, et al – Dienogest inhibits C-C motif chemokine ligand 20 expression in human endometriotic epithelial cells. Eur J Obstet. Gynecol. Reprod. Biol. 2017; 214:65–70

[131] Loudon NB – A three month trial of a regimen of oral contraceptives that reduces the frequency of menstruation. Br. Med. J.; 1977:2: 487-490

[132] Edelman A, Micks E, Gallo MF, Jensen JT, Grimes DA – Continuous or extended cycle vs. cyclic use of combined hormonal contraceptives for contraception. Cochrane Database Syst Rev’2014 Jul 29;7: CD004695. doi: 10.1002/14651858.CD004695.pub3.

[133] Foidart JM, Sulak PJ, Schllschnidt A, et al. The use of an oral contraceptive containing etynyl estradiol and drospirenone in an extended regimen over 136 days. Contraception, 2006; 73: 34-40

[134] Harada T, Kosaka S, Elliesien J, Yasuda M, Momoeda M, et al, – Ethinyl estradiol 20 μg/drospirenone 3 mg in a flexible extended regimen for the management of
endometriosis-associated pelvic pain: a randomized controlled trial. Fertil. Steril 2017; 108:798–805

[135] Cackrimsnidou AC, Hellberg D, Nilsson S, Waldemström U, Sikström B, et al – Long interval treatment regimen with a desogestrel-containing oral contraceptive. Contraception, 1993 Sep; 48 (3), 205-16. doi:10.1016/0010-7824(93)80141-h

[136] Edelman A, Koontz SL, Nichols MD et al – Continuous oral contraceptives. Are bleeding patterns depending on the hormones given? Obstet. Gynecol; 2006, 107:657-65

[137] Caruso S, Iraci M, Cianci S, Fava V, Cianci A, et al – Comparative, open-label prospective study on the quality of life and sexual function of women affected by endometriosis-associated pelvic pain on 2mg dienogest/30μg ethinyl estradiol continuous or 21/7 regimen oral contraceptive. J. Endocrinol. Invest. 2016; 39, 923-931

[138] Archer DF, Jensen JT, Johnson JV, Borisute H, Constantine GD, et al – Evaluation of a continuous regimen of levonorgestrel/ethinyl estradiol: phase 3 study results. Contraception. 74(6):439-45

[139] Benagiano G, Carrara S, Fillippi V – Safety, efficacy and patient satisfaction with continuous daily administration of levonorgestrel/ethinylestradiol oral contraceptives. Patient Prefer Adherence: 3: 131-43.

[140] Wiegratz J, Kuhl H – Long-cycle treatment with oral contraceptives. Drugs: 64 (21): 2447-62

[141] Kuhl H, Birkhäuser M, Müeck A, Neulen J, Brandle W, et al – Long-cycle treatment in oral contraception. Ther Umsch; 66 (2):101-8. doi: 10.1024/0040-5930.66.2.101

[142] Read MC – New regimens with combined oral contraceptive pills-moving away from traditional contraceptive 21/7 cycle. The European J. of Contraception & Reproductive Health Care, 2010, 12 (suppl 2)

[143] Slak PJ, Scow RG, Preece C, et al – Hormone withdrawal symptoms in oral contraceptives users. Obstetr. Gynecol, 2000, 95: 261-6

[144] Sucato GS, Gerschultz KL – Extended cycle hormonal contraception in adolescents Curr. Opin. Obstetr. Gynecol; 2005 Oct;17(5): 461-5.

[145] Huang T, Shafrir LA, Eliasssen A-H, Rexrode KM, Tworoger SS – Estimated Number of Lifetime Ovulatory Years and Its Determinants in Relation to Levels of Circulating Inflammatory Biomarkers. Am. J. Epidemiol; 2020 Jul; 189 (7);660-70 doi: 10.1093/aje/kwz264

[146] Jensen TG, Schiaff W, Gordon K – Use if combined hormonal contraceptives for the treatment of endometriosis related pain: a systematic review of evidence. Fertil Steril 2018 Jul, 110 (1):137-152

[147] Vercellini P, Buggio L, Frattaruolo MP, Borghi A, Somigliana E, et al – Medical treatment of endometriosis-related pain. Best Pract. Res. Clin. Obstet. Gynaecol. 2018; 51:68-91. doi: 10.1016/j. bpo.2018.01.015.

[148] Damm T, Lamvu G, Carrillo J, Ouyang C, Feranec J – Continuous vs cyclic combined hormonal contraceptives for treatment of dysmenorrhea: a systematic review. Contracept X, 2019 Jan; 24:1; 100002

[149] Cicinelli E, de Ziegler D, Alfonso R, Nicoletti R, Bellavia M, Colafiglio G – Endometrial effects, bleeding control, and compliance with a new postmenopausal hormone therapy regimen based on transdermal estradiol
gel, and every-other-day vaginal progesterone in capsules: a 3-year pilot study. Fertil. Steril. 2005; 83:1859-63.

[150] Price JH, Ismail H, Gorwill RH, Sarda IR – Effect of the suppository base on progesterone delivery from the vagina. Fertil. Steril. 1983, 39,490–493.

[151] Cicinelli E, de Ziegler D, Alfonso R, Nicoletti R, Bellavia M, Colafiglio G – Endometrial effects, bleeding control, and compliance with a new postmenopausal hormone therapy regimen based on transdermal estradiol gel, and every-other-day vaginal progesterone in capsules: a 3-year pilot study. Fertil. Steril. 2005; 83: 1859-63

[152] Cicinelli E, Schonauer LM, Galantino P, Matteo MG, Cassetta R, Pinto V – Mechanisms of uterine specificity of vaginal progesterone. Human Reproduction, 2000; 15, (Suppl. 1), 159-165

[153] Deligdish L – Hormonal ptology of The Endometrium. Modern Pathology 2000, vol 13, 285-294

[154] Davids Landau M, Church K – Should You Opt for Natural Progesterone Treatments for Endometriosis? Brigham and Women's Hospital. website, last updated february 2018

[155] Gomes MKO, Rosa-E-Silva JC, Garcia SB, De Sá Rosa-E-Silva ACJ, Ferriani RA, et al – Effects of the levonorgestrel-releasing intrauterine system on cell proliferation, Fas expression and steroid receptors in endometriosis lesions and normal endometrium Hum Reprod 2009;24:2736-2745

[156] Morelli M, Sacchinelli A, Venturella R, Moccioaro R, Zullo F – Postoperative administration of dienogest plus estradiol valerate versus levonorgestrel-releasing intrauterine device for prevention of pain relapse and disease recurrence in endometriosis patients. J. Obstet. Gynaecol. Res. 2013; 39:985-990

[157] Yao Z, Shen X, Capodanno I, Donnelly M, Fenyl-Melody J, Vakerich K, et al – Validation of rat endometriosis model by using raloxifene as a positive control for the evaluation of novel SERM compounds. J. Invest. Surg. 2005;18:177-183. doi: 10.1080/08941930591004412.

[158] Stratton P, Sinaii N, Segars J, Koziol D, Nieman LK, et al – Return of chronic pelvic pain from endometriosis after raloxifene treatment: A randomized controlled trial. Obstet. Gynecol. 2008;111:88-96. doi: 10.1097/01.AOG.0000297307.35024.b5.

[159] Kulak J, Jr, Fischer C, Komm B, Taylor HS – Treatment with bazedoxifene, a selective estrogen receptor modulator, causes regression of endometriosis in a mouse model. Endocrinology. 2011;152: 3226-3232. doi: 10.1210/en.2010-1010.

[160] Vercellini P, Cortesi I, Crosignani PG – Progestins for symptomatic endometriosis: A critical analysis of the evidence. Fertil. Steril. 1997;68:393-401. doi: 10.1016/S0015-0282(97)00193-3.

[161] Tosti C, Biscione A, Morgante G, Bifulco G, Luisi S, Petraglia F – Hormonal therapy for endometriosis: From molecular research to bedside. Eur. J. Obstet. Gynecol. Reprod. Biol. 2017;209:61-66. doi: 10.1016/j.ejogrb.2016.05.032.

[162] Kettel LM, Murphy AA, Morales AJ, Yen SS – Clinical efficacy of the antiprogestrone RU486 in the treatment of endometriosis and uterine fibroids. Hum. Reprod. 1994;9(Suppl. 1):116-120. doi: 10.1093/humrep/9.supp1_1.116.

[163] Kettel LM, Murphy AA, Morales AJ, Ulmann A, Baulieu EE, Yen
SS – Treatment of endometriosis with the antiprogesterone mifepristone (RU486). Fertil. Steril. 1996; 65:23-28. doi: 10.1016/S0015-0282(16)58022-4.

[164] Whitaker LH, Murray AA, Matthews R, Shaw G, Critchley HO, et al – Selective progesterone receptor modulator (SPRM) ulipristal acetate (UPA) and its effects on the human endometrium. Hum Reprod 2017; 32:531-543

[165] Fu J, Song H, Zhou M, Zhu H, Wang Y, Chen H, et al – Progesterone receptor modulators for endometriosis. Cochrane Database Syst Rev. 2017;(7)

[166] Compton DR, Sheng S, Carlton KE, Rebacz NA, Katzenellenbogen JA, et al – Pyrazolo[1,5-alpyrimidines] estrogen receptor ligands possessing estrogen receptor beta antagonist activity. Journal of medicinal chemistry, 2004, 47; 5872-5893

[167] Zheng QM, Mao HL, Zhao YJ, Zhao J, Wei X, Liu PS – Risk of endometrial polyps in women with endometriosis: a meta-analysis. Reproductive biology and endocrinology: 2015;13(1):103. doi: 10.1186/x12958-015-0092-2

[168] Han SJ, Hawkins SM, Begum K, Jung SY, O’Malley BW, et al – A new isoform of steroid receptor coactivator-1 is crucial for pathogenic progression of endometriosis. Nat. Med. 2012;18:1102-1111. doi: 10.1038/nm.2826

[169] Prentice A, Deary AJ, Goldbeck-Wood S, Farquhar C, Smith SK – Gonadotrophin-releasing hormone analogues for pain associated with endometriosis. Cochrane Database Syst Rev 2000: CD000346-CD000346

[170] Sagsveen M, Farmer JE, Prentice A, Breeze A – Gonadotrophin-releasing hormone analogues for endometriosis: bone mineral density. Cochrane Database Syst Rev 2003: CD001297-CD001297

[171] Hornstein MD, Surrey ES, Weisberg GW, Casino LA – Leuprolide acetate depot and hormonal add-back in endometriosis: a 12-month study. Obstet Gynecol 1998; 91: 16-24

[172] Practice Committee of the American Society for Reproductive Medicine – Treatment of pelvic pain associated with endometriosis: a committee opinion. Fertil Steril 2014; 101: 927-935

[173] Sroyraya M, Songkoomkrong S, Changklungmoa N, Poljaroen J, Sobhon P, et al – Differential expressions of estrogen and progesterone receptors in endometria and cyst walls of ovarian endometrioma from women with endometriosis and their responses to depot-medroxyprogesterone acetate treatment. Mol. Cell Probes. 2018; 40:27-36. doi: 10.1016/j.mcp.2018.07.001.

[174] Crosignani PG, Luciano A, Ray A, Bergqvist A – Subcutaneous depot medroxyprogesterone acetate versus leuprolide acetate in the treatment of endometriosis-associated pain. Hum. Reprod. 2006; 21:248-256. doi: 10.1093/humrep/dei290

[175] Brown J, Pan A, Hart RJ – Gonadotrophin-releasing hormone analogues for pain associated with endometriosis. Cochrane Database Syst Rev 2010: CD008475-CD008475

[176] Cetel NS, Rivier J, Vale W, Yen SS – The dynamics of gonadotropin inhibition in women induced by an antagonistic analog of gonadotropin-releasing hormone. J. Clin. Endocrinol. Metab. 1983; 57:62-65. doi: 10.1210/jcem-57-1-62.

[177] Taylor HS, Giudice LC, Lessey BA, Abrao MS, Kotarski J, Watts NB, et al.– Treatment of Endometriosis-Associated Pain with Elagolix, an Oral GnRH Antagonist. N. Engl. J. Med. 2017;
[178] Maggi R, Cariboni AM, Marelli MM, et al – GnRH and GnRH receptors in the pathophysiology of the human female reproductive system. Hum. Reprod. Update: 2016; 22:358-381

[179] Barbieri RL – Endometriosis and the threshold theory. J Reprod Med, 1992, 43 (3) suppl,287-92

[180] Stratton P, Berkley KJ – Chronic pelvic pain and endometriosis: translational evidence of the relationship and implications. Hum Reprod Update 2011; 17: 327-346

[181] Ng J, Chwalisz K, Carter DC, Klein CE – Dose-dependent suppression of gonadotropins and ovarian hormones by elagolix in healthy premenopausal women. J Clin. Endocrinol. Metab. 2017;102:1683-1691

[182] Amsterdam LL, Gentry W, Jobanputra S, Bulun SE – Anastrozole and oral contraceptives: a novel treatment for endometriosis Ferti.l Steril 2005;84:300-304

[183] Ersoy GS, Zolbin MM, Cosar E, Mamillapalli R, Taylor HS– Medical therapies for endometriosis differentially inhibit stem cell recruitment Reprod Sci 2017;24:818-823

[184] Bilotas M, Meresman G, Stella I, Sueldo C, Baranao RI – Effect of aromatase inhibitors on ectopic endometrial growth and peritoneal environment in a mouse model of endometriosis. Fertil. Steril. 2010; 93:2513-2518. doi: 10.1016/j. fertnstert.2009.08.058.

[185] Verma A, Konje JC – Successful treatment of refractory endometriosis-related chronic pelvic pain with aromatase inhibitors in premenopausal patients. Eur. J. Obstet. Gynecol. Reprod. Biol. 2009;143:112-115. doi: 10.1016/j.ejogrb.2008.12.002

[186] Ailawadi RK, Jobanputra S, Kataria M, Gurates B, Bulun SE – Treatment of endometriosis and chronic pelvic pain with letrozole and norethindrone acetate: a pilot study. Fertil. Steril 2004; 81:290-296

[187] Ferrero S, Camerini G, Seracchioli R, Ragni N, Venturini PL, Remorgida V – Letrozole combined with norethisterone acetate compared with norethisterone acetate alone in the treatment of pain symptoms caused by endometriosis. Hum Reprod; 2009; 24: 3033-3041

[188] Ferrero S, Leone Roberti Maggiore U, Scala C, Remorgida V, et al – Changes in the size of rectovaginal endometriotic nodules infiltrating the rectum during hormonal therapies, Arch Gynecol Obstet; 2013;287:447-453

[189] Ferrero S, Remorgida V, Venturini PL, Leone Roberti Maggiore U– Norethisterone acetate versus norethisterone acetate combined with letrozole for the treatment of ovarian endometriotic cysts: a patient preference study Eur. J. Obstet. Gynecol. Reprod. Biol; 2014; 174:117-122

[190] Lee B, Du H, Taylor SH (2009) –Experimental murine endometriosis induces DNA methylation and altered gene expression in eutopic endometrium. Biology of Reproduction, 2009, 80 (1), 79-85

[191] Cakmak H, Taylor HS (2010) – Molecular mechanisms of treatment resistance in endometriosis: the role of progesterone-hox gene interactions. Semin Reprod Med; 2010 28 1 69 74

[192] Meyer J, Zimbardi D, Podgaec S, Amorim RL, Abrao MS, Rainho CA– DNA methylation patterns of steroid receptor genes ESR1, ESR2 and PGR in deep endometriosis compromising the rectum. International journal of molecular medicine. 2014;33(4):897-904. doi:10.3892.ijmm.2014.1637
[193] Slack A, Cervoni N, Pinard M, Szyf M – Feedback regulation of DNA methyltransferase gene expression by methylation. European journal of biochemistry / FEBS. 1999;264(1):191-9

[194] Dyson MT, Roqueiro D, Monsivais D, Ercan CM, Pavone ME, Bulun SE, et al – Genome-wide DNA methylation analysis predicts an epigenetic switch for GATA factor expression in endometriosis. PLoS Genet. 2014 Mar; 10(3):e1004158