Endometriosis and ovarian cancer risk, an epigenetic connection

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Abstract: Endometriosis is a gynecological disorder that affects 176 million women worldwide and 1 in 10 females in the United States. Endometriosis most often affects women of child-bearing age, with most going undiagnosed. Endometriosis also shares many characteristics common to invasive cancer and has been known to be associated with epithelial ovarian cancer. Ovarian cancer is the 11th most common cancer among women and over 22,000 new cases will be diagnosed within the next year. Women most commonly diagnosed with this cancer are between the ages of 55–64 years, outside the range of the age of women affected with endometriosis. While no known cause of either disease has been established, epigenetic regulation is thought to play a major role in both. This review focuses on epigenetic changes that occur within each individual disease as well as those that are similar in both, suggesting a possible etiological link between the two diseases.

Keywords: Endometriosis-associated carcinoma; long non-coding RNA; epigenetic drugs

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Introduction

Endometriosis is an elusive gynecological disorder defined by the growth and presence of endometrial tissue (lesions) outside of the uterus (1). This ectopic endometrial tissue responds to hormones and drugs in the same manner as its eutopic counterpart, and its continued growth is dependent on the hormone estrogen (2-4). Approximately 10–15% of women of child-bearing age suffer from endometriosis. Of the 1 in 10 women who suffer from it, 20–25% show no signs or symptoms, while others may experience symptoms such as infertility, chronic pelvic pain, or both (5-7). Currently, there are three main theories on how endometriosis originates (8). The most widely accepted is Sampson's theory, based on the retrograde shedding of endometrial cells into the abdominal cavity during menstruation, which in turn promotes a chronic inflammatory response (9). The inflammatory response in endometriosis has been studied extensively including in our laboratory (10-20). The second is the coelomic metaplasia theory in which the mesothelium differentiates in situ into endometrial tissue (21). The third is the embryonic origin theory in which the lesions are already present throughout the development and maturation in women (22). Other theories include stem cells (23), genetics, and autoimmune disease (24,25). In addition to the lesions, affected women have an increased volume of peritoneal fluid (PF), that comprises of inflammatory cells, pain-inducing molecules, growth factors, cytokines/chemokines, etc. (10,11,19,26). Our studies and others have shown a dynamic role for the PF in the etiology of endometriosis and the symptoms

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associated with this disease (10,11,19,27-29). While endometriotic lesions are not considered malignant, they do have cancer-like properties such as invasion of local organs, abnormal growth, and genetic damage (30,31).

Ovarian cancer is diagnosed annually in nearly a quarter of a million women globally. It is responsible for 140,000 deaths each year making it the 4th largest cause of cancer death in women and represents 1.3% of all new cancer cases. Epithelial ovarian cancer (EOC) contributes to 90% of the diagnosis. There are currently nine different types of EOC: high-grade serous, low-grade serous, mucinous, ovarian endometrioid invasive, clear-cell, undifferentiated or unclassified, primary peritoneal, Brenner tumors, and borderline tumors. With EOC, the early stages show no symptoms and there are no screening processes currently available. If diagnosed in the later stages, the cancer has usually spread to the peritoneal cavity and surgery is typically no longer an option. The 5-year survival rate for late stage diagnosis is 25% versus 90% for early stage (32).

Both endometriosis and ovarian cancer have many etiological similarities. One glaring commonality between the two is that there is no definitive known cause for either of these diseases. Epidemiological studies have shown that the prevalence of ovarian cancer in women with endometriosis is higher than sporadic ovarian cancer. Among the EOC subtypes, clear-cell, endometrioid invasive ovarian cancer, and low-grade serous are the most commonly seen in women with endometriosis (33,34). A pooled analysis of 13 case-control studies of ovarian cancer in women with self-reported endometriosis consisting of 13,226 controls and 7,911 women with invasive ovarian cancer showed that endometriosis was associated with an increased risk of clear-cell with an odds ratio (OR) of 3.05, low-grade serous with an OR of 2.11, and endometrioid invasive ovarian cancer with an OR of 2.04 (35). In another study, 2.9% of women with endometriosis went on to develop ovarian cancer and the rate increased in women who had a longer history of endometriosis (36,37).

These two diseases also share similar etiologies such as: damaged ovarian epithelium, elevated gonadotropins, androgen excess with progesterone deficiency, retrograde menstruation, oxidative stress and chronic inflammation (38). Recently there are several supporting evidence for the role of epigenetics in each of these diseases and possibly linking them. Figure 1 summarizes some of the commonalities that are seen in both endometriosis and ovarian cancer. In this review, we will only highlight the epigenetic mechanisms that they share. Some of these mechanisms have been linked to the transformation of endometriosis to its associated carcinomas, however others have been shown to be expressed in each of these diseases independently.

**Epigenetic mechanisms**

Epigenetics refers to the stable inheritance of phenotypes of cells and organisms without changes in DNA sequence or DNA content. This could involve DNA methylation, histone modifications, or changes in microRNAs (Figure 2). The changes and modifications that occur are reversible and dynamic and can be modified by environmental or lifestyle factors throughout an organism’s entire lifespan (39,40).

The most common epigenetic mechanism which regulates many cellular processes, including embryonic development, transcription, chromatin structure and stability (41) is DNA methylation. This involves the covalent transfer of a methyl group to the C-5 position of the cytosine ring of DNA by DNA methyltransferases (DNMTs) (42). The donor molecule then dissociates, and the methylated DNA is released. DNMTs initiate and maintain the DNA methylation patterns, which are then regulated by methyl-CpG binding proteins (MDBs) that recognize and read the methylation marks created by the DNMTs (41). Typically, when a CpG island becomes methylated in cancer cells, loss of gene expression or silencing occurs due to the methylation of the islands inhibiting specific transcription factors from binding, as well as the recruitment of MDBs and their repressive chromatin remodeling functions (41). The methylation of CpGs is irreversible and can be passed down to daughter cells (43,44).

The second common type of epigenetic change is histone modifications. These modifications are commonly referred to as post-translational modifications. They occur on the N-terminal histone tails that extend from the nucleosomes, which then affects the chromatin structure, and the recruitment of remodeling enzymes. One of the widely investigated histone modification is histone methylation, which occurs mainly on lysines and arginines that are present on the histone tails, but does not change the charge of the tail itself. These changes can be reversed by demethylation by lysine demethylase 1 (LSD1). The second most common modification is histone acetylation which occurs on the lysines of the tails and is regulated by histone acetyltransferases (HATs) and histone deacetylases (HDACs). Acetylation of lysine in the histones results in removal of its positive charge thus weakening its interaction with the DNA. The third modification is histone
phosphorylation of serines, threonines and tyrosines, which adds a negative charge and changes the conformation of the histone on the nucleosome structure (45,46). Chromatin remodeling has been shown to have a higher level of complexity and diversity than DNA methylation. The dynamic changes occurring affect the chromatin structure which influences the gene expression. Gene expression can also be affected by chemical modifications of histone proteins themselves (47).

One of the key regulatory complexes involved in histone modifications is the Polycomb repressive complex 2 (PRC2). A major subunit of the PRC2 complex is Enhancer of Zeste Homologue (EZH2), a histone methyltransferase (48-50). EZH2 plays a key role in transcriptional repression through chromatin remodeling, thus regulating gene expression as well as maintaining cell identity and oncogenesis (51). EZH2 activates H3K27 through trimethylation, H3K27me3 (52). This trimethylation causes gene silencing in cells. Studies have shown elevated EZH2 expression levels in various cancers (49,53,54).

MicroRNAs (miRNAs) are an emerging third field of study in epigenetics which play a role in regulating gene expression. miRNAs are single-stranded non-coding RNAs, usually consisting of 22 nucleotides, that perform gene regulation without changing the DNA sequence. MiRNAs function in RNA silencing by two mechanisms. The first is via mRNA degradation where there is complete complementarity between the miRNA and the target mRNA nucleotide sequences. A single miRNA may be complementary to one or more mRNA, meaning that the gene silencing is not very specific (55-57). The second is by preventing the mRNA from being transcribed if the miRNA sequence is not completely complimentary. Several miRNAs control the expression of important epigenetic regulators which include DNMTs, HDACs and PRC2 complex, thus forming an epigenetic-miRNA regulatory circuit and feedback network (58,59).

**Epigenetics and endometriosis**

![Figure 1: Etiological pathways that are commonly shared between endometriosis and ovarian cancer.](image)
With no well-defined etiology known for endometriosis, epigenetics may be involved in its pathogenesis. Figure 3A summarizes some of the recently reported studies that support the role of the three epigenetic processes in endometriosis. There are evidences that show an over expression of the three DNMTs (DNMT 1, 3A, and 3B) in ectopic compared to eutopic tissues in women with endometriosis (60). Endometriosis is steroid responsive disease. The increase in ERβ mRNA levels in endometriotic tissues when compared to control tissue is due to promoter hypomethylation (61). Studies have similarly shown hypermethylation of the PR-B promoter region in endometriosis leading to its downregulation (62). Steroidogenic factor-1, SF-1, a nuclear receptor that regulates many genes in the hypothalamic-pituitary-adrenal endocrine axis, is expressed in ectopic endometriotic tissue but not eutopic tissue (63). Its aberrant expression in endometriosis is due to its hypomethylation in the CpG islands. SF-1 is also recruited to the promoter of steroidogenic genes and is a key regulator for estradiol synthesis and essential for growth and inflammation of endometriotic tissue. In endometriosis, E-cadherin, involved in cell-cell adhesion, is down-regulated and hypermethylated in its promoter region (64). Its expression can be induced by treatment with a histone deacetylase. The hypermethylation of Homeobox A10, HOXA10, a highly conserved homeobox gene, results in its reduced expression in the eutopic endometrium in women with endometriosis, resulting in defective uterine receptivity (39,65). Decreased expression of HOXA10 results in the activation of autophagy in endometriosis (66).

In addition to the various enzymes involved in histone modifications (HDAC, HATs, HMTs and MeCP2) (39), studies from our laboratory provided the first evidence of an increased expression of EZH2/H3K27me3 in endometriotic lesions (67) which was later confirmed by other groups (68). EZH2 also enhances epithelial-mesenchymal transition in endometriosis (69). In endometriosis, the levels of H3K9 and H4K16 acetylation were lower, but methylation levels of H3K4 and H3K9 were higher (70).

The expression and function of miRNAs, including their role in epigenetic regulation, are an emerging area of focus in endometriosis (71). Many mRNA transcripts are differentially regulated in endometriotic lesions when compared to those of eutopic tissues. There are several studies that have determined circulating as well as tissue specific miRNA profiles in endometriosis. One such
study identified differential expression of 48 miRNAs in a microarray analysis, with the majority of these miRNAs displaying a reduced expression in eutopic and ectopic tissue when compared with control tissue. Validation by qRT-PCR demonstrated dysregulation of 2 specific miRNAs, miR-21 and miR-26a (72). Our laboratory recently showed that oxidation-sensitive miRNAs such as miR-29a and let-7g may play a role in endometriosis associated pain (11). A recent study identified miR-141 by driving the TGFβ1/SMAD2 signaling activates epithelial-to-mesenchymal transition (EMT) in endometriosis (73).

**Epigenetics and ovarian cancer**

DNA methylation is also an important epigenetic biomarker for EOC (74). *Figure 3B* summarizes some but not all of the epigenetic changes observed in EOC. BRCA1’s promoter methylation is shown to be connected to malignancy in ovarian cancer (75). Higher silencing of this gene is observed in high-grade serous ovarian cancer (HGSOC) and is also connected with lower overall survival of patients. The HOXA9 promoter is hypermethylated in early-stage tumors of EOC (76). Comparing HOXA9 promoter methylation in normal endometrium from women with EOC and healthy controls, it was determined that this methylation is predictive of presence of EOC and the more methylation present, the more progressive was the cancer (77). The RASSF1A gene is a tumor suppressor gene, which is often repressed in ovarian cancer due to DNA methylation (78). RASSF1A is also a direct target of H3K27me3 which results in its downregulation in ovarian cancer cells (79). The most commonly mutated gene in HGSOC according to Cancer Genome Atlas is TP53, which is involved in signaling pathways that halt cell cycle progression or induce apoptosis. It is an important tumor suppressor gene and the methylation of its promoter is seen in EOC (80).

*Figure 3* List of epigenetic marks that have been identified and discussed in the review: (A) endometriosis and (B) ovarian cancer.
aggressiveness of the disease (81). DOT1-like protein, Dot1L, is the only known methyltransferase for H3K79 which is involved in cell cycle regulation and DNA repair. Dot1L expression was lost in ovarian cancer cells resulting in the promotion of cell invasion and induction of stem-cell like properties. This loss also downregulated E-cadherin and Tight Junction Protein 1, promoting EMT but upregulated the expression of Aldehyde Dehydrogenase 1 Family Member A1, by Wnt signaling activation (82). A Disintegrin and Metalloproteinase 19, ADAM19, is part of the ADAM family and has been shown to be involved in cell-cell and cell-matrix interactions as well as being an active metalloproteinase involved in cell migration and signal transduction. ADAM19 is also a target of TGF-β1, whose pathway is often dysregulated in ovarian cancer. In ovarian cancer cells that were refractory to TGF-β1 stimulation, ADAM19 expression was lower. These cells with disrupted TGF-β1 signaling also had repressive histone marks, H3K27me3 and H3K9me2, associated with the promoter region of ADAM19 causing epigenetic dysregulation (83).

miR-214 is a miRNA that is involved in pro-growth and anti-apoptotic pathways and is a target in EOC. Its deregulation, leads to survival and resistance of the cell. This miRNA targets phosphatase and tensin homolog (PTEN), a tumor suppressor, which leads to activation of the Akt pathway, and is associated with cell survival (84). Other miRNAs that were studied in ovarian cancer are miR-484, -642 and -217, which predict chemoresistance. When secreted by ovarian cancer cells, these miRNA targets the VEGF signaling pathway which regulates vascular development and angiogenesis by acting directly on the VEGFB protein or inhibiting the VEGFR2 receptor in the endothelial cells (85).

**Common epigenetic marks in endometriosis and ovarian cancer**

**Epigenetic markers**

**ARID1A**

The AT-Rich Interaction Domain 1A (ARID1A) gene encodes a member of the SWI/SNF family and is required for transcriptional activation of genes that are normally repressed by chromatin (86). Mutations in ARID1A result in the loss of BRG-associated factor 250a (BAF250a), a protein with an important role in cell proliferation and tumor suppression, and a possible link between endometriosis and endometriosis-associated carcinomas. Samartzis et al. observed a complete loss of BAF250a in benign endometriotic lesions and that the expression of BAF250a was lower in ovarian endometriosis when compared to eutopic endometrium, peritoneal endometriosis, and deep-infiltrating endometriosis (87). The expression of this gene was also examined using whole transcriptomics on samples obtained from the Ovarian Cancer Research frozen-tumor bank. Mutations in ARID1A were present in 46% of ovarian clear-cell carcinomas and 30% of endometroid carcinomas, but none were seen in any of the HGSOC samples examined. These mutations also correlated with a loss of BAF250a expression in the tumors. Since these mutations were seen in the preneoplastic lesions, it could indicate an earlier event in the transformation from endometriosis to ovarian cancer (88).

**RASSF2**

Ras Association Domain Family Member 2, RASSF2, is highly expressed in humans and is a known tumor suppressor gene whose down-regulation in cells is due to promoter methylation. One study showed that RASSF2 may be a candidate gene for the malignant transformation of ovarian endometriosis. In this study, researchers used fresh tissue samples from endometriosis-associated ovarian carcinoma patients and found RASSF2’s promoter was hypermethylated and the protein expression was significantly lower in the neoplastic tissue when compared to the ectopic endometrial tissue, suggesting that this epigenetic inactivation is an early event in the transformation (89).

**PTEN**

PTEN is another genetic alteration seen in ovarian cancer and endometriotic lesions suggesting a possible malignant transition. Using a mouse model, Dinulescu et al. induced endometrioid ovarian cancer and saw that when PTEN was deleted in the background of oncogenic K-ras activation, endometriotic precursor lesions were formed. These lesions went on to develop invasive endometrioid ovarian carcinoma (90). In endometriosis, overexpression of PTEN significantly increased apoptosis as well as inhibited the cell cycle (91). Cells expressing low levels of PTEN showed increased angiogenesis and the VEGF expression was high (91). In ovarian cancer, PTEN can suppress its growth by negatively regulating the PI3K/Akt pathway and that an abnormal expression of this pathway is associated with poor prognosis (92). Also, as discussed earlier many of the miR-
214 effects in ovarian cancer were mediated by targeting PTEN. One study using a mouse model examined using PTEN as a gene therapy in ovarian cancer and found that overexpression of PTEN resulted in lower tumor growth and an increased survival time (93). Figure 4 shows string networks centered on PTEN for both endometriosis and ovarian cancer.

**EZH2**

EZH2, as described earlier, is shown to play a role in endometriosis by activating H3K27me3 and shutting down transcription (94). EZH2 is also shown to play a role in ovarian cancer by enhancing angiogenesis (54,86). Higher levels of EZH2 correlated to a worse prognosis for EOC patients (54). Interestingly, when miR-214 is downregulated in ovarian cancer, EZH2 is upregulated, increasing cell proliferation and transcriptional repression through H3K27me3 in the same manner as that seen in endometriosis (95).

**microRNA**

There are several studies where differential expression of miRNAs have been studied in either endometriosis or ovarian cancer, however, Braicu *et al.* compared miRNAs in both these diseases in the same study. It was found that there were more differentially expressed miRNA in endometriosis compared to ovarian cancer and most were linked to EMT (96). Two common miRNAs overexpressed in both diseases were miR-325 and miR-492 (96). While the expression of miR-325 was upregulated in both diseases, it was more upregulated in ovarian cancer suggesting miR-325 as having a role in the transition from endometriosis to ovarian cancer (96). Not many studies have examined the levels of these miRNAs in either disease but upregulation of miR-492 has been shown to decrease PTEN expression which increases the level of AKT activation in cancer cells (97). miR-325 is downregulated in hepatocellular carcinoma but its upregulation inhibited cell invasion and proliferation by targeting HMGB1 (98). miR-21 which was earlier discussed as potentially having a role in endometriosis (72), can cause a cisplatin resistance in ovarian cancer and that its inhibition can enhance the tumor suppressor Programmed Cell Death 4, PDCD4, as well as promote apoptosis (99).

**miR-199a**

In endometriosis, Wang *et al.* found that miR-199a relative expression levels were higher in samples from endometriosis.
patients versus controls and noted that circulating levels of miR-199 increased as the disease progressed. Interestingly, it was also noted that the concentration of miR-199a was significantly higher in ovarian endometriosis than in other types of endometriosis (100). Downregulation of miR-199a correlated with an upregulation of HIF1α in ovarian tumors when compared to normal tissues and to shorter survival in patients. When miR-199a was increased it promoted cisplatin-induced apoptosis by suppressing HIF1α in the cells (101).

miR-135
MiRNA-135 represses HOXA-10 thereby altering endometrial receptivity and implantation (102). In the endometrial lesions, expression of miR-135a/b was decreased when compared to the normal tissue. However, expression was increased in the secretory phase compared to the proliferative phase, by over 10-fold in the lesions and nearly 8-fold in the tissues, suggesting a role in endometrial function and implantation in women with endometriosis (103). In ovarian cancer, it was found that miR-135a was significantly down regulated in EOC tissues which correlated to a high expression of HOXA10 in these tissues when compared to controls. This low expression of miR-135a also correlated with a shorter progression-free survival. The downregulation of HOXA10 also resulted in changes in apoptosis-associated genes, such as caspase-3 and p53, which may control EOC tumor growth (104).

miR-141
miR-141 is a member of the miR-200 family known to modulate the Notch, p53, MAPK, and ErbB signaling pathways. In endometriosis, expression of this miRNA was decreased in the ectopic endometrium when compared to normal and eutopic endometrium as well as being decreased in the ectopic endometrial stromal cells. When cells were transfected with miR-141, a lower growth rate was observed, but it promoted cell proliferation and reduced their migration. Cells transfected with an miR-141 mimic decreased Kruppel like factor 12 (KLF-12), an oncogene seen in several cancers, suggesting that it is a direct target of miR-141 (105). In ovarian cancer cells, Mak et al. showed that miR-141 significantly increased their anchorage-independent growth. This upregulation also correlated to cancer metastasis. When miR-141 was inhibited in cells, a significant reduction in cell viability by over 50% was seen. miR-141 and KLF-12 were shown to have an inverse relationship in this disease as well, it had a tumor-suppressive function in ovarian cancer. Overall, their study showed that miR-141 suppresses KFL-12 which causes cells to escape apoptosis, survive, and metastasize (106). Wu et al. showed that this oncogenic miRNA was higher in ovarian cancer when compared to its concurrent endometriosis and could be a possible marker that distinguishes the two and could be a possible mediator of transformation between the diseases (107).

Long non-coding RNA
Long non-coding RNAs (lncRNA) are newer biomarkers implicated in various diseases including ovarian cancer (108) and endometriosis (109,110). They are longer than 200 nucleotides and not translated into protein. Though long considered to be “junk”, recent evidences support its role in several diseases (111). As with miRNAs, some of the lncRNA discussed below though shown to be playing a role in both these diseases, have not yet been directly linked to each other.

MALAT1
Metastasis-associated lung adenocarcinoma transcript 1 (MALAT1) is a lncRNA that plays a role in various types of cancers, including ovarian (112). Its function is mediated through PI3K/AKT signaling pathway and its silencing has resulted in a decrease in EMT and an increase in E-cadherin levels in some cancers (108). In endometriosis, there is a time-dependent increase in MALAT1 in the stromal cells treated with hypoxia correlating with an increase in HIF-1α in the cells. These elevations caused an activation of pro-survival autophagy pathway, reducing apoptosis in endometriosis (113). MALAT1 is overexpressed in ovarian cancer, with higher expression correlating to a larger tumor size and metastasis. When MALAT1 was knocked down in an ovarian cancer cell line, SK-OV-3, there was a decrease in cell viability as well as proliferation, migration, and invasion suggesting that MALAT1 plays a role in ovarian cancer growth and development (112).

H19
In normal cells, H19 is an imprinted lncRNA whose function is acting as a molecular sponge for miRNA let-7 and inhibiting its function. Endometrial H19 expression is regulated by estrogen and progesterone positively and negatively, respectively (114). Ghazal et al. showed that the expression of H19 was significantly decreased in women with endometriosis when compared to normal women,
which correlated with an increase in let-7 activity. The increase in this miRNA inhibited the post-transcriptional expression levels of the IGF1 receptor (IGF1R) and resulted in decreased proliferation of endometrial stromal cells (115). In ovarian cancer, H19 expression was shown to be increased in cisplatin-resistant cells and this expression had a strong correlation with cancer recurrence. When H19 was knocked down in cisplatin-resistant cells, resistance was reversed in cells that were more sensitive to hydrogen-peroxide treatments (116).

**BANCR**

BRAF-activated non-coding RNA, BANCR, is increased in cancers such as thyroid, lung, and bladder (117). In an endometriosis rat model, the volume of the grafts in the BANCR inhibitor group was smaller than that of the control group as well as the development of the graft tissues being repressed prominently. Using H&E staining, it was shown that there was fewer interstitial cells and a declined blood supply when compared to the control group. BANCR can suppress production of angiogenic factors in endometriosis which could repress development of ectopic endometrial tissues (118). BANCR expression was examined in EOC patient tissues and shown to be increased when compared to normal tissues. This high expression also correlated with an advanced FIGO stage, high CA125 expression, and with an occurrence of metastasis in the lymph nodes (119).

**Drug therapy**

With the evidence accumulating that epigenetic changes may be the underlying cause for the initiation and progression of both diseases, many physicians and scientists have turned to targeting epigenetic pathways for treatment. In ovarian cancer, one of the most common epigenetic drugs tested is histone deacetyltransferase inhibitors, HDACi which work by increasing the level of acetylated histones, reactivating silenced tumor suppressor genes. Currently, only three HDACi, vorinostat, romidepsin, and panobinostat, have been approved by the FDA (120). These HDACi have very little effect alone, hence many are used in combination therapy. Treatments with DNMT inhibitors and HDACi have been investigated as a means of reversing platinum resistance in ovarian cancer. An HDACi that is currently in the preclinical stage for ovarian cancer is Trichostatin A, TSA. TSA inhibits class I and II HDACs as well as induces expression of P73 and facilitate Bax-dependent apoptosis in cancer cells that are resistant to cisplatin (121). One specific DNMT inhibitor, decitabine, was shown to lead to demethylation of many genes including those that were discussed in this review, such as HOXA10 and BRCA1 (122,123).

Hydralazine is an anti-hypertensive medication that was shown to inhibit non-nucleoside DNA methylation. This drug is in clinical trials along with valproic acid, VPA, which works through oxidative stress and generates reactive oxygen species to inhibit HDACs. This combination was tested in refractory solid tumors, including ovarian cancer. Out of 15 total patients with various solid tumors, four had partial response (3 ovarian cancer) and 8 (4 ovarian) had stable disease 4 weeks after completing one or two cycles of chemotherapy. Results showed that epigenetic drugs could be used to re-sensitize tumors to chemotherapy and should be used in combination (124). Many of the miRNA and lncRNA discussed above were also shown to play a role in chemotherapy resistance and could be future drug targets.

Bevacizumab (Avastin) is an antiangiogenetic drug used to target VEGF and inhibit proliferation and migration of the cells. Trials have shown that bevacizumab, when used as an initial therapy, improved progression free survival in ovarian cancer patients (125).

Another combination approach that is being examined is epigenetic inhibitors and immunotherapy, which has been used in cancers such as non-small cell lung cancer and melanoma with good success, however is in its very early stages of being used in ovarian cancer. Stone et al. found that in a mouse model of EOC, DNMTi and HDACi improved the response to immune checkpoint therapy. Specifically, the DNMTi 5-azacytidine, AZA, increased the number of CD45+ immune cells as well as the percentage of NK cells and active CD8+ cells in the microenvironment. This, in turn, reduced the tumor burden and extended the survival. A triple combination therapy consisting of DNMTi, HDACi, and an immune checkpoint inhibitor provided the greatest effects and longest overall survival (126).

In endometriosis, epigenetic drugs have not been explored to the same extent as that in ovarian cancer, but much of the research has focused on HMT inhibitors, specifically those that work against EZH2. The two common drugs of this type are 3-Deazaneplanocin A hydrochloride, DZNep, and GSK126. DZNep is a lysine methyltransferase EZH2 inhibitor and competitive s-adenosyl homocysteine hydrolase inhibitor, that blocks H3k27me3 and H4k20me3 (127). GSK126 is a selective, S-adenosyl-methionine-competitive small molecule...
inhibitor of EZH2 methyltransferase activity and is more than 1,000-fold selective for EZH2 over other HMTs (128). In endometriosis, DZNep was shown to inhibit the growth of endometriotic lesions as well as attenuated EMT and reduced fibrosis in vivo (69). Although GSK126 is currently being investigated as a treatment for endometriosis, it has not been studied as extensively as DZNep (68). However, it has been studied in endometrial cancer (129). In ovarian cancer, GSK126 caused a regression of ARID1A-mutated ovarian clear cell carcinoma and decreased the number of tumor nodules in xenograft models (86). Recent studies showed that GSK126 suppressed the growth and improved the survival of Coactivator Associated Arginine Methyltransferase, CARML-high expressing HGSOC and EOC as CARML promotes EZH2-mediated silencing of target tumor suppressor genes (130).

TSA is also being examined in endometriosis by looking at its effects on nonsteroidal anti-inflammatory drug activated gene 1 (NAG-1) expression and apoptosis. NAG-1, a member of the TGF-β family and associated with apoptosis and cell growth is decreased in the endometrium of patients with endometriosis. Ectopic endometrial tissues when treated with varying concentrations of TSA resulted in induction of NAG-1 leading to apoptosis in a dose-dependent manner (131). Other studies have also shown that TSA can reduce endometrial implants in mice (132). Wu et al. found VPA to induce cell cycle arrest, as well as p21 expression in endometrial stromal cells (133). Another group used a rat model of endometriosis to test the effects of VPA along with progesterone. They examined the serum levels of TNF-α, lesion size, and body weight. Rats receiving VPA and/or progesterone showed a reduced lesion size and increased weight gain compared to the untreated rats as well as an improved response to thermal stimuli, suggesting that this combination may be an efficient treatment for endometriosis (134).

Conclusions

Throughout this review, epigenetic markers that have been identified and studied extensively in endometriosis, ovarian cancer, or both have been discussed. While neither disease has a definitive cause, there are many avenues that could be responsible to play a role in their etiology. And while endometriosis is not a disease that affects all women, and even in those who are affected, its phenotype is different, it is plausible to theorize that epigenetics plays a major role in its initiation and progression. Similarly, not much is known about how endometriosis transforms into carcinoma? Some of the common epigenetic markers found in these two diseases might play a plausible role in this linkage.

Many scientists around the world are focusing their efforts not only in identifying the cause of these diseases, but also on ways for its early detection and treatment. Many of the current treatments available are only of temporary relief, however with the gain of newer knowledge, better treatment options are growing for both these diseases. It is however getting clear from these studies that there is no single gene or marker that is truly responsible for either of these diseases, but however, a combination of multiple biomarker changes all working together is involved in the disease initiation and progression. What remains a fact is that more research is warranted in these two diseases, as more and more women are being diagnosed each year from them and hence need well designed diagnostics and therapeutics to curb them.

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