Ovarian cancer is one of the most prevalent pathologies in gynaecology. This malignancy can be divided into 2 large groups: epithelial and non-epithelial. Because epithelial ovarian cancers (EOC) are the most commonly diagnosed, this paper focuses on the latest therapies associated with this disease. Due to the difficult diagnosis, EOC is frequently detected in the advanced stage. The treatment is usually complex and requires specialist knowledge. Advances and new ideas, such as identification of various genes and molecules that can serve as prognostic factors, might increase patients’ chances of survival; they may contribute to optimization of patients’ treatment, deciding whether to use aggressive treatment strategies, and predicting chemoresistance. Moreover, new strategies might also improve the quality of life of patients. The study aimed to analyse and discuss the latest reports on new methods of managing EOC.

Key words: ovarian cancer, immunotherapy, chemotherapy, epithelial cancer, treatment.

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What is new about ovarian malignancies?

Kinga Grabska1, Izabela Pilarska1, Marta Magdalena Fudalej2, Andrzej Deptała2, Anna Badowska-Kozakiewicz2

1 Students’ Scientific Organization of Cancer Cell Biology, Department of Cancer Prevention, Medical University of Warsaw, Poland
2 Department of Cancer Prevention, Medical University of Warsaw, Poland

Introduction

Ovarian cancer is one of the most prevalent pathologies in gynaecology. The current histopathological classification distinguishes over 70 types of tumours that may be detected in the ovary. Ovarian tumours can be divided into 2 large groups: epithelial and non-epithelial. Most ovarian cancers are sporadic; only 5–10% of them are presented with a family history [1].

Ovarian cancer incidence is relatively stable in Western countries. However, the percentage of ovarian cancer among gynaecological neoplasms is increasing, which is related to the decline in the incidence of cervical cancer in European countries due to cytological screening programs [2]. Trends in relative survival and population mortality show slight improvement. The changes in mortality can be explained partially by a decrease in the incidence of ovarian cancer (presumably due to the use of oral contraceptives and a reduction in the use of hormones after menopause) [3].

Survival in ovarian cancer is the worst among all gynaecological cancer locations [4]. The main reasons for the poor survival include the lack of early detection methods and the ovaries’ unfavourable anatomical location. Treatment for ovarian cancer is complex and requires specialist knowledge and experience in oncology and surgery [2, 5–7]. Advances in ovarian cancer treatment can significantly improve patients’ chances of survival.

The study aimed to analyse and discuss the latest reports on new methods of managing epithelial ovarian cancer (EOC). Articles published from 1976 to 2020 in the PubMed and Elsevier databases were analysed. The authors focused on analysing risk factors, prognostic and predictive factors, and possible cancer therapies.

Epithelial ovarian cancer – characteristics

The most common type of ovarian cancer is epithelial, occurring in 95% of cases [8]. Every year 220,000 women in the world are diagnosed with this malignancy [9]. The high death rate results from non-specific early symptoms, late diagnosis, and high metastatic rate within the abdomen. Despite modern management and complete response to the treatment, many patients diagnosed with advanced disease develop a recurrence within 2.5 years [10].

Four primary histological types of EOC can be distinguished: serous (high-grade serous carcinomas [HGSC] and low-grade serous carcinomas), endometrioid, mucinous, and clear cell. Epithelial ovarian malignancies are divided into 2 categories – type I and type II tumours. The first type comes originally from the ovary and is caused by continual ovulation cycles, endometriosis, and inflammation. These tumours are more often diagnosed as a disease of a low stage. Therefore, they are less lethal and usually present a better outcome than the second type, which comes usually from a fallopian tube and is related to genetic mutations of the BRCA gene, p53, or other tumour-suppressing genes [8].
Among EOC symptoms, the following might be distinguished: abdominal pain and distention, early satiety, constipation, bloating, nausea, signs from the urinary tract, later fatigue, and loss of weight [11]. The measurement of CA-125 concentration, pelvic ultrasound followed by pelvic computed tomography or magnetic resonance imaging, and diagnostic surgery with pathological examination are the initial diagnostic investigations. The first-line treatment of EOC consists of the primary debulking surgery and platinum-based chemotherapy with or without an anti-endothelial growth factor (VEGF) agent [12].

**What is new about epithelial ovarian cancer?**

**Risk factors**

Having a family history of breast and ovarian cancer, mutations of breast cancer type 1 susceptibility protein (BRCA 1), breast cancer type 2 susceptibility protein (BRCA 2), and DNA mismatch repair genes, uninterrupted ovulation cycles (early onset of menses, childlessness, late menopause), endometriosis, ethnicity, smoking, hormonal replacement therapy, and diet are well-known risk factors of the occurrence of EOC [8]. Although these endangering agents are well-known and widely studied, many studies about them and newly discovered factors are reported each year.

The latest studies have revealed that the gender of the offspring might also present an impact on EOC risk. Bearing a male offspring was associated with an 8% lower risk of EOC. Compared to bearing all-female offspring, having all-male offspring was associated with a 14% decrease in EOC risk. The protective effect seems to be enhanced by the increasing number of male children. The biological explanations for this phenomenon might be associated with different maternal hormone concentrations [13]. Despite the fact that the study failed to reach statistical significance, previous studies – in eastern Pennsylvania [14] and Sweden [15] – reported similar findings. Slightly different results were reported in a recent pooled analysis among participants from 12 case-control studies, which included 6872 EOC patients. It was found that each additional offspring was associated with an 8% decrease regardless of whether the child was male or female. The sex of the offspring was not linked with EOC risk for serous, clear cell, and endometrioid histotypes. However, bearing male offspring was associated with lower protection against mucinous histotype, but the male sex appeared to have no significant relation to this kind of EOC [16].

Modugno et al. showed in their research that breastfeeding for at least 3 months provides significant protection from EOC. This protection is increased by an earlier age at first breastfeeding and a larger number and longer duration of breastfeeding episodes. Although this protection decreases over time, it can last for more than 30 years [17]. Another study dealt with the protective value of breastfeeding among patients with BRCA mutations. It showed that breastfeeding history was associated with a decrease of 23% in ovarian cancer risk among BRCA carriers. From 1 to 7 months of breastfeeding increased the protective effect, and after that period the association was relatively stable. The use of oral contraceptives was reported to be a significant independent protective factor, which enhanced the positive effect of breastfeeding [18].

Other research dealt with the link between benign ovarian tumours and the risk of EOC. The cohort group consisted of over 150,000 women with a primary or secondary diagnosis of benign ovarian tumours (e.g. serous cystadenoma, mucinous cystadenoma, clear cell adenofibroma, benign Brenner tumour, thecoma, and fibroma) between 1978 and 2016. The occurrence of benign ovarian tumours doubly increased the risk of mucinous ovarian cancer. The risk was independent of the age and was accelerated up to 20 years after diagnosing benign ovarian tumours. No link was found between other histological types of EOC and previous benign ovarian tumours [19].

In the Ovarian Cancer Cohort Consortium study, a connection between tumour dominance and different reproductive and hormonal risk factors of EOC was observed. Women with a dominant tumour mass were less likely to be parous (among those who have fewer children), have ever smoked, have ever taken oral contraception (OC), have had a hysterectomy, tubal ligation, or unilateral oophorectomy, in comparison to women with a non-dominant tumour mass. Additionally, patients with a right-dominant tumour mass were more likely to be parous or to have ever used OC than those with a left-dominant tumour mass. The body mass index is related to a meaningly increased risk of left- dominant ovarian cancer. The linkage between the histological type of a tumour and dominance was also observed. There were more clear cell subtype masses in left-dominant cases and more serous subtypes of tumour in the right-dominant ones [20] (Table 1).

There are also some recent studies claiming that viral infections with Papillomavirus (HPV), Cytomegalovirus (CMV), and Epstein-Barr virus (EBV) [21], as well as an exposition to different types of asbestos fibres in occupational and environmental settings, may contribute to the development of ovarian cancers [22, 23].

**Prognostic and predictive factors**

The Federation of Gynaecology and Obstetrics (French. Fédération internationale de gynécologie et d’obstétrique – FIGO) staging system for ovarian cancer is the primary tool that provides prognostic information and guidance of management for ovarian malignancy. The assessment is based on diagnostic surgery and imaging [24]. Besides the FIGO staging, classifying EOC as types I or II, which depends on the histological type and grading, also implicates a prognosis for the patients. Type I is associated with better outcomes and more prolonged survival [25]. However, there are cases in which the disease progression does not correlate with the initial assessment. What else has an impact on the prognosis? The main reason for treatment failure of ovarian malignancy is resistance to platinum-based chemotherapy [26]. After platinum-based chemotherapy, many patients experience a relapse and become drug-resistant within 2 years. As a result, the survival rate for them is approximately 40% [27]. This phenomenon might be correlated with intrinsic or acquired factors and various ovarian cancer mechanisms [28]. Xing
et al. analysed 6 genes [Fanconi anaemia complementation group A (FANCA), Fanconi anaemia complementation group G (FANCG), DNA Polymerase Delta 1 (POLD1), lysine (K)-specific demethylase 1A (KDM1A), Bloom syndrome protein (BLM), and BRCA 1], potentially affecting the answer to platinum-based treatment. The 6 candidate genes were verified in various validation sets. The insightful examination of these genes proved that the mRNA and protein expression levels possessed a meaningful difference in the analysis of clinical and pathological factors. Moreover, notable relationships between these genes and both overall survival and disease-free survival were detected. The real-time polymerase chain reaction assay confirmed that those candidate genes’ expression levels in platinum-sensitive ovarian cancer cell lines were higher than those in platinum-resistant ones [29].

There are also some specific molecules, expression of which can affect disease progression. Therefore, they can serve as a new prognostic marker and a therapeutic target for the EOC. The study of Zou et al. analysed the expression patterns, prognostic value, genetic variation, and biological functions of 12 members of the ubiquitin-conjugating enzyme E2 (UBE2) gene family. They observed that the mRNA levels of UBE2C, UBE2N, UBE2S, and UBE2T were notably upregulated in ovarian malignancy compared to normal ovarian tissue. Patients with serous ovarian cancer and upregulation of UBE2A, UBE2B, UBE2G, UBE2R2, and UBE2T were characterized by poor overall survival. Moreover, upregulation of UBE2A, UBE2N, and UBE2R2, and downregulation of UBE2T and UBE2G were associated with poor progression-free survival. Being significantly upregulated in EOC compared with that in borderline tumours, benign tumours, and normal ovarian tissues, UBE2T is thought to have a high diagnostic accuracy [30]. The study by Li et al. reported that UBE2C levels are significantly higher in cisplatin-resistant cells than in sensitive ones. Silencing of UBE2C expression using lentiviral-mediated short hairpin RNA (shRNA) in model cells resulted in enlarged tumour size, increased proliferation, and decreased apoptosis [31]. Another study proved that UBE2C and histone-lysine N-methyltransferase (EZH2) genes are potential therapeutic targets and should be investigated for their clinical use [32].

Another prognostic and therapeutic target of EOC may be lysine-rich coiled-coil 1 (KRCC1). It is a nuclear protein overexpressed by tumour cells in patients with high-grade serous ovarian cancer. Higher levels of KRCC1 expression correlate with chemoresistance and poor outcomes. Silencing experimentally, KRCC1 inhibits cellular plasticity and invasive properties and enhances apoptosis, leading to tumour growth reduction [33].

Some other genes and molecules may influence the EOC prognosis and contribute to treatment failure. Recent studies have concentrated on the following molecules (examples only): Annexin A8 (ANX8) [34], Ten-Eleven Translocation Protein 3 (TET3) [35], Retinoic Acid-Inducible Gene-1 (RIG-I) [36], Desmoglein-2 (DSG2) [37], Maternal Embryonic Lucine Zipper Kinase (MelK) [38], Eukaryotic Translation Initiation Factor 2B Subunit Epsilon (Eif2BS), B-Arresin 2 [39], and RING Finger Protein 126 [40]. The potential of incorporating new molecules into prognostic and predicting schemes for ovarian cancer patients seems to be high; however, further, more complex research is needed.

### Treatment

The primary treatment for early EOC is surgery. It is used for both staging and debulking (cytoreduction). It can be an exclusive and curative treatment in a disease confined to the ovaries. For later stages of the disease (IC and II), postsurgical chemotherapy is recommended. It usually consists of platinum and paclitaxel [41]. Platinum-based chemotherapy has been the primary management of ovarian cancer since the 1980s [3]. Neoadjuvant chemotherapy is also used in some nonoperative cases. Apart from intravenous chemotherapy, intraperitoneal can be ordered after suboptimal cytoreductive surgery in the FIGO III stage. Bevacizumab — a humanized monoclonal antibody targeting vascular VEGF — can be added to the conventional chemotherapy for patients after suboptimal cytoreductive (FIGO III stage) as well as for patients with disseminated carcinoma [42, 43]. Additionally, Olaparib — an oral poly(ADP-ribose) polymerase (PARP) inhibitor — can be ordered for patients with serous ovarian cancer and the presence of BRCA 1/2 mutations. Olaparib serves as a maintenance treatment for patients after relapse, who responded to platinum-based chemotherapy [44].

Epithelial ovarian cancer therapy has barely changed in the last 40 years, and it still seems to lack effective therapeutic targets. Although EOC was supposed to be scarcely immunogenic, immunotherapy appears to have more therapeutic potential than was previously thought. A lot of emphases is placed on improving the management of ovarian cancer and making it more personalized [43]. So, what is new about the treatment of ovarian cancer?

The primary goal of surgical treatment is to accomplish a complete resection with clear margins. Score R0 or R1 in the residual tumour (R) classification implicates better survival rates [45]. Ceppi et al. proved that targeted molecule-based fluorescence imaging helps achieve complete tumour resection on the microscopic scale. They used a fluorescence imaging system with an orthotopic mouse model.
to assess tumour detectability and to evaluate the effect of fluorescence imaging-guided surgery. The contrast agent used in this study was an intra-peritoneal nanomolecular probe, composed of single-walled carbon nanotubes, coupled to an M13 bacteriophage carrying a modified peptide binding to the secreted protein acidic and rich in cysteine overexpressed in studied mice. High microscopic tumour detection was observed with a pixel-limited resolution of 200 µm. Additionally, the researchers observed an elevated survival in animals treated with fluorescence image-guided surgical resection compared to the typical surgery [46].

The pleiotropy of transforming growth factor-beta (TGF-β) signalling in tumour tissues covers cancer initiation, development, metastasis, and reaction between stroma and cancer cells [47]. TGF-β plays a crucial role in the metastasizing by providing epithelial-mesenchymal transition (EMT). TGF-β signalling may be activated by over-expression of domain-containing ion transport regulator 5 (FXYS5) in cancer cells. The TGF-β activates the SMAD3/SMAD4 complex, which initiates the transcription of further effectors and promotes FXYS transcription. The FXYS5 creates a positive loop with TGF-β to drive EMT and, therefore, metastasis. It makes TGF-β a potential target in the treatment of ovarian cancer [48]. It is claimed that the debulking signature is centred around the hyperactivation of the TGF-β pathway, which drives the overexpression of genes located in the tumour and its microenvironment.

Using the TGF-β inhibitors with chemotherapy, for patients after suboptimal debulking surgery or in the neoadjuvant treatment, may improve the interval debulking surgery. TGF-β inhibitors decrease migration and invasion of the tumour cells and increase response to the treatment [49]. Zhang et al. reported that inhibiting the TGF-β pathway with LY2157299 monohydrate (TGF-R1 inhibitor) in animal models reduced tumour cell proliferation, migration, and invasion. Additionally, it was found that LY2157299 is involved in slowing down cancer growth and ascites formation. These effects are correlated with reduced expression of vital stroma proteins COL11A1 and VCAN [50].

In 2020, a study assessing the impact of prolonged pre-operative cycles on survival, accounting for surgical outcomes, was published. The study compared the results of treatment of 199 women with newly diagnosed ovarian cancer. Women who received 3 or 4 neoadjuvant cycles were compared with women who received 5 or more cycles. Apart from the number of cycles between the groups of women, there were no other differences in clinical factors. The rates of complete resection were similar, regardless of the number of chemotherapy cycles received. Unfortunately, more cycles of chemotherapy (5 or more) were associated with poorer progression-free survival. The analysis showed that patients taking 3–4 cycles of chemotherapy had a better prognosis than patients receiving 5 or more cycles [51].

Some recent studies reported spontaneous tumour regressions [52, 53], occasional persistent responses to immune checkpoints-inhibitors [54], and longer progression-free time and overall survival in patients with tumours with a high rate of T-cells [55]. These revelations suggest that EOC patients would potentially benefit from immunotherapy.

Zamarin et al. showed that the combination of nivolumab and ipilimumab in platinum-resistant EOC resulted in better response and longer progression-free survival. However, more studies should be conducted [56].

Combining anti-angiogenic agents and immunotherapy might occur beneficially. The single usage of anti-angiogenic agents resulting in a reduced number of blood vessels and increased tumour hypoxia seems inadequate. However, immunotherapy might be supported by additional effects of drugs targeting VEGF, angiopoietin 2, or hepatocyte growth factor pathways. A few clinical trials
evaluating immune-checkpoint inhibitors in combination with anti-angiogenic agents in patients with EOC are currently in progress [57] (Table 2).

PARP inhibitors (PARPi) have been used in EOC treatment since 2014, when the Food and Drug Administration and the European Medicines Agency approved them. Olaparib, Niraparib, Rucaparib, Talazoparib, and Veliparib are the most relevant PARPi. The last 2 of them are currently under investigation. PARPi are used in patients with recurrent ovarian cancer and BRCA mutation [58]. It seems that combined treatment with PARPi and immune checkpoint blockade might be beneficial for patients with BRCA mutation. A phase I study, in which olaparib and tremelimumab [cytotoxic T-cell antigen 4 (CTLA4) immune checkpoint antibody] were used, was conducted to assess this regimen’s tolerability. No dose-limiting toxicities were identified [59]. Two more studies are currently exploring these treatment strategies. The first one is the combined therapy with niraparib and pembrolizumab [programmed death 1 (PD-1) checkpoint inhibitor] in patients with triple-negative breast cancer or recurrent ovarian cancer. The phase 1/2 study results are promising, with an overall response rate (ORR) of 25% in platinum-resistant ovarian cancer and an ORR of 45% in patients with BRCA mutations. The most common reported side effects were anaemia and thrombocytopenia [60]. Another study suggested a synergistic therapeutic effect of the combination of olaparib and durvalumab [anti-programmed cell death ligand-1 (PD-L1)] in patients with relapsed gastric cancer [61].

Several new cell cycle checkpoint inhibitors are currently under investigation. A recent study reported the efficacy of prexasertib – a second-generation checkpoint first and second kinase (CHK1/2) inhibitor for patients who are diagnosed with BRCA wild-type high grade serous ovarian cancer and who have undergone intensive treatment. Prexasertib seems to be a useful tool in treating patients with platinum-resistant or platinum-refractory disease [62]. The studies reported that the combination of PARPi with Ataxia Telangiectasia and Rad3-related (ATR) and its downstream Checkpoint Kinase 1 (CHK1) inhibitor is more effective in reducing tumour burden in BRCA mutation models, in comparison to monotherapy [63].

Folate receptor-α (FOLR-1), which is highly expressed in tumour tissue in HGSC patients and usually undetectable in normal tissue, is another potential therapeutic target [64]. FOLR-1 is a glycosylphosphatidylinositol-connected membrane glycoprotein, which is exposed to the extracellular molecules [65]. It is involved in DNA replication and damage repair in cells by mediating cellular responses to folate, including proliferation, cell division, and tissue growth [66]. Some trials focused on monoclonal antibodies binding to the FOLR-1 – farletuzumab [67] and mirvetuximab soravtansine [68]. Moreover, the adoptive cell transfer (ACT) therapy, in which tumour-specific cytotoxic T-cells are expanded in vivo and then infused after lymphodepleting chemotherapy [69], might be used in EOC patients. Westergaard et al. expanded tumour-infiltrating lymphocytes (TILs) from 34 tumour specimens of ovarian cancer and showed the recognition of autologous tumour cells in > 50% of the patients. Moreover, antigen-specific TILs were isolated and further expanded in vivo. These findings supported the hypothesis that patients with OC could benefit from ACT [70] and were followed by the pilot study [71].

Cancer stem cells (CSCs) are a population of tumour cells with self-renewal abilities responsible for both tumour development and resistance to the applied treatment [72]. The drug resistance in CSCs leads to relapses during treatment [73]. Signalling pathways such as Wnt/β-catenin and NOTCH are, among others, in charge of chemoresistance in CSCs [74, 75]. Therefore, inhibition of the Wnt pathway might become an efficient management in EOC treatment. Ipaparotide (OMIP-54F28), a first-in-class recombinant fusion protein with the extracellular part of human Frizzled-8 receptors fused to a human IgG1 Fc fragment that binds Wnt ligands, is undergoing phase I study in pancreatic and ovarian cancers [76]. Additionally, some trials concentrate on Notch pathway silencing in EOC treatment. Delta-like ligand 4 (Dll4), one of the Notch ligands, has been proven to be overexpressed in ovarian cancer. Dll4 is claimed, among others, to be responsible for tumour resistance to the anti-VEGF therapy [77]. The study by Huang et al. showed that combining Dll4 inhibitors (murine REGN1035 and human REGN421) with anti-VEGF treatment (aflibercept) significantly reduces ovarian tumour growth. Dll4 blockade (REGN1035) combined with aflibercept has more significant therapeutic effects. This success may result from the increased apoptosis in tumour cells and increased transcription factor-GATA3 expression under hypoxia conditions [78].

Conclusions

Epithelial ovarian cancer is the most lethal gynaecological malignancy worldwide. It is usually diagnosed in the advanced stage and is associated with poor outcomes. Therefore, searching for and analysing the risk factors of these malignancies is extremely valuable. It can lead to the development of new prevention strategies and a decrease in cancer-related deaths. As the general principle says, prevention is better than cure. Recent years have brought reports about different genes and molecules, which may serve as prognostic factors for EOC patients. They can be used to stratify patients, decide about the usage of aggressive treatment strategies, or predict chemoresistance. Efforts are also being made to increase the clinical effectiveness of the treatment while minimizing toxicities for the patients. Several findings have identified potential new attractive therapeutic approaches, which are still under investigation. Recent studies suggest that EOC patients might benefit from immunotherapy.

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Address for correspondence
Marta Magdalena Fudalej, MD
Department of Cancer Prevention
Medical University of Warsaw
81 Ziürki i Wigury St.
01-091 Warsaw, Poland
e-mail: mmfudalej@gmail.com