Nanotechnology and Immunotherapy in Ovarian Cancer: Tracing New Landscapes

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ABSTRACT

Ovarian cancer (OC) is the seventh most common cancer in women worldwide. Standard therapeutic treatments involve debulking surgery combined with platinum-based chemotherapy. Of the patients with advanced-stage cancer who initially respond to current treatments, 50%–75% relapse. Immunotherapy-based approaches aimed at boosting antitumor immunity have recently emerged as promising tools to challenge tumor progression. Treatments with inhibitors of immune checkpoint molecules have shown impressive results in other types of tumors. However, only 15% of checkpoint inhibitors evaluated have proven successful in OC due to the immunosuppressive environment of the tumor and the transport barriers. This limits the efficacy of the existing immunotherapies. Nanotechnology-based delivery systems hold the potential to overcome such limitations. Various nanoformulations including polymeric, liposomes, and lipid–polymer hybrid nanoparticles have already been proposed to improve the biodistribution and targeting capabilities of drugs against tumor-associated immune cells, including dendritic cells and macrophages. In this review, we examine the impact of immunotherapeutic approaches that are currently under consideration for the treatment of OC. In this review, we also provide a comprehensive analysis of the existing nanoparticle-based synthetic strategies and their limitations and advantages over standard treatments. Furthermore, we discuss how the strength of the combination of nanotechnology with immunotherapy may help to overcome the current therapeutic limitations associated with their individual application and unravel a new paradigm in the treatment of this malignancy.

Introduction

Ovarian cancer (OC) ranks as the seventh leading cause of death in women worldwide. According to the American Cancer Society, 14,070 deaths and approximately 22,240 new cases were predicted for 2018 in the United States (Siegel et al., 2018). Of the patients with advanced-stage cancer who initially respond to current treatments, 50%–75% relapse. The asymptomatic nature of early-stage ovarian cancer is the main reason for its late diagnosis, which normally occurs at a metastatic stage, drastically reducing the chances of a successful outcome of the treatment (Das and Bast, 2008; Rauh-Hain et al., 2011). Despite the continuous improvement in screening methods, OC-associated mortality rates remain high due to the absence of routine early detection approaches. The lack of specificity of the available tests and the limitations associated with the application of imaging techniques further complicate the diagnostic process (Sarjini et al., 2012; Terry et al., 2016; Russell et al., 2017). OC comprises five histologic subtypes: low-/high-grade serous, mucinous, clear cells, and endometrioid cancer. Serous OC represents the most common carcinoma and accounts for more than 50% of all cases. It is associated with specific genetic mutations (i.e., BRCA1, BRCA2, MMR, TP53, BARD1, CHEK2, RAD51, and PALB2) spanning from single nucleotide polymorphisms to high frequency of somatic gene copies or epigenetic features.
indicative of defects in homologous recombination repair and gene methylations (Kaldawy et al., 2016; Ducie et al., 2017). These subtypes metastasize to the same area within the peritoneal cavity.

Currently, the treatment of OC includes debulking surgeries, which are meant to excise tumor masses, coupled with extensive chemotherapy, radiotherapy, or a combination of the three depending on the stage and type of cancer. Recommended first-line treatments for OC are platinum-based and taxol drugs (www.nccn.org/guidelines). In some cases, after genetic screening, patients may be eligible for monoclonal antibody therapies such as bevacizumab, which blocks tumor angiogenesis by inhibiting the vascular endothelial growth factor signaling. Other approaches include using olaparib, rucaparib, and niraparib, known as inhibitors of the poly(adenosine diphosphate [ADP]-ribose) polymerases and involved in DNA repair. The use of the latter treatments has been specifically recommended for patients with BRCA gene mutations (Coward et al., 2015). Table 1 explains the current therapies available for OC, including standard and targeted chemotherapies. The state-of-the-art nanotherapies currently being used or tested in clinical trials are also mentioned.

The 5-year survival rate for women with advanced-stage OC is approximately 40% (Timmermans et al., 2018; Torre et al., 2018) but increases if the ovarian tumor has more infiltrating T cells (Zhang et al., 2003). The lack of a curative therapeutic regimen, the frequency of relapse, and the mortality levels underlie the effort needed to refine the curative therapeutic regimen, the frequency of relapse, and involved in DNA repair. The use of the latter treatments has been specifically recommended for patients with BRCA gene mutations (Coward et al., 2015). Table 1 explains the current therapies available for OC, including standard and targeted chemotherapies. The state-of-the-art nanotherapies currently being used or tested in clinical trials are also mentioned.

The diversity of physiopathology (Nezhat et al., 2015) between OC types and the heterogeneity of cells infiltrating the peritoneum calls for the identification of effective approaches to maintain the bioactivity of the payload, precisely aim at the target, and preferentially accumulate the drug at the site of interest while reducing cytotoxicity.

Nanomedicines are frequently used as engineered drug delivery systems that support the prolonged circulation of drugs, maintain their bioactivity, reduce their side effects, and selectively target diseased cells (Blanco et al., 2015). Targeted nanomedicines include liposomal nanocarriers [small interfering RNA (siRNA)–EphA2, OSI-211, Myocet (liposome-encapsulated doxorubicin citrate; Ben Venue Laboratories)] (Seiden et al., 2004; Shen et al., 2013; Eitan et al., 2014), polymeric nanoparticles (Abraxane [Protein-bound Paclitaxel, Celgene], CRLX101) (Srinivasan et al., 2014; Pham et al., 2015), and antibody-drug conjugates (Howard et al., 2016). Nanotechnology-based strategies for diagnostic tools have been also developed to detect biomarkers and genetic mutations (Engelberth et al., 2014), as well as to combine nano-enabled therapeutic and diagnostic capabilities, giving rise to “nano-theranostics” (Yaari et al., 2016).

In this review, we discuss the potential of cancer immunotherapy, a recently developed field that aims at treating cancer patients by restimulating their immune system. Particular emphasis is given to its applications and pitfalls in OC. We also review how a nanomedicine approach to immunotherapy may overcome the current therapeutic

### Table 1

<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Drug Class</th>
<th>Formulation</th>
<th>FDA Approved/Clinical Trial Phase</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gold standard chemotherapeutic</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Antibiotics/antineoplastics</td>
<td>1995</td>
<td>Bolis et al., 1978</td>
<td></td>
</tr>
<tr>
<td>Carboplatin</td>
<td>Alkylating agents</td>
<td>1989</td>
<td>Adams et al., 1989</td>
<td></td>
</tr>
<tr>
<td>Paclitaxel</td>
<td>Mitotic inhibitors</td>
<td>1998</td>
<td>Khanna et al., 2015</td>
<td></td>
</tr>
<tr>
<td>Cyclophosphamide</td>
<td>Alkylating agents</td>
<td>1959</td>
<td>Handdias et al., 2016</td>
<td></td>
</tr>
<tr>
<td>Gemcitabine</td>
<td>Antimetabolites</td>
<td>2006</td>
<td>Lorusso et al., 2006</td>
<td></td>
</tr>
<tr>
<td>Melphalan</td>
<td>Alkylating agents</td>
<td>2001</td>
<td>Hasan and Jayson, 2003</td>
<td></td>
</tr>
<tr>
<td>Cisplatin</td>
<td>Alkylating agents</td>
<td>1978</td>
<td>Monneret, 2011</td>
<td></td>
</tr>
<tr>
<td>Topotecan</td>
<td>Miscellaneous antineoplastics</td>
<td>1996</td>
<td>Seiden et al., 2004</td>
<td></td>
</tr>
<tr>
<td>Etoposide</td>
<td>Mitotic inhibitors</td>
<td>1998</td>
<td>Long et al., 2005</td>
<td></td>
</tr>
<tr>
<td>Thiotepa</td>
<td>Alkylating agents</td>
<td>2001</td>
<td>Gordinier et al., 2002</td>
<td></td>
</tr>
<tr>
<td><strong>Targeted therapies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bevacizumab</td>
<td>VEGF/VEGFR inhibitors</td>
<td>2004</td>
<td>Rossi et al., 2017</td>
<td></td>
</tr>
<tr>
<td>Olaparib</td>
<td>PARP inhibitors</td>
<td>2017</td>
<td>Moore et al., 2018</td>
<td></td>
</tr>
<tr>
<td>Niraparib</td>
<td>PARP inhibitors</td>
<td>2017</td>
<td>Essel and Moore, 2018</td>
<td></td>
</tr>
<tr>
<td>Rucaparib</td>
<td>PARP inhibitors</td>
<td>2018</td>
<td>Dal Molin et al., 2018</td>
<td></td>
</tr>
<tr>
<td><strong>Current nanotechnology treatments and ongoing trials</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Doxil</td>
<td>Antibiotics/antineoplastics</td>
<td>Pegylated liposomal doxorubicin</td>
<td>1999</td>
<td>Pisano et al., 2013</td>
</tr>
<tr>
<td>Lipodox</td>
<td>Antibiotics/antineoplastics</td>
<td>Pegylated liposomal doxorubicin</td>
<td>2012</td>
<td>Chou et al., 2006</td>
</tr>
<tr>
<td>Genexol-PM</td>
<td>Mitotic inhibitors</td>
<td>PEG-PLA polymermic</td>
<td>Phase II</td>
<td>Lee et al., 2017</td>
</tr>
<tr>
<td>LEP-ETU</td>
<td>Mitotic inhibitors</td>
<td>Liposomal paclitaxel</td>
<td>Phase I</td>
<td>Damjanov et al., 2005</td>
</tr>
<tr>
<td>Pacidal</td>
<td>Mitotic inhibitors</td>
<td>Paclitaxel micelles</td>
<td>Phase III</td>
<td>NCT00989131</td>
</tr>
<tr>
<td>OSI-211</td>
<td>Antineoplastics</td>
<td>Liposomal lurtotecan</td>
<td>Phase II</td>
<td>Seiden et al., 2004</td>
</tr>
</tbody>
</table>

PARP, poly(adenosine diphosphate [ADP]-ribose) polymerase; PLA, polyactic acid; VEGF, vascular endothelial growth factor.
limitations of the treatment of OC and unravel a new paradigm in the cure of this malignancy.

**Immunotherapy and Cancer**

Cancer immunotherapy aims at stimulating the immune system to provide cancer prevention and treatment. The first discoveries of the crucial role played by the immune regulation in cancer progression have recently led to the 2018 Nobel Prize for Medicine and Physiology to Dr. James P. Allison and Dr. Tasuku Honjo (www.nobelprize.org). Their studies unraveled fundamental mechanisms that govern immune cell (specifically T cells) responses to cancer and provided insights to overcome immune system evasion by cancer. Since then, the use of immune checkpoint blockade has been widely recognized as an effective cancer treatment. In particular, Dr. Allison and his research group have been the first to identify the cytotoxic T-lymphocyte antigen 4 (CTLA-4) protein, an immune checkpoint receptor expressed on the surface of activated T cells that is believed to regulate their proliferation. When the CTLA-4 pathway is activated by costimulatory molecules (CD80, CD86), the result is hindrance of T-cell function, which inhibits the T-cell strong anticancer potential (Leach et al., 1996). Based on these observations, a specific antibody was developed to retain CTLA-4 activation and maintain T cells in an activated status (Chambers et al., 1996). Almost simultaneously, in 1992 Dr. Honjo’s group discovered programmed death-ligand 1, which also acts as a T-cell retainer, finding an alternative way to defeat the tumor-mediated immune evasion. The insights provided by such inspiring scientists have led to many Food and Drug Administration (FDA)–approved drugs for the treatment of various cancers. These drugs span from sipuleucel-T, approved in 2010 to target the immune system for the treatment of prostate cancer (Cheever and Higano, 2011), to ipilimumab, the first monoclonal antibody against CTLA-4 for metastatic melanoma (Lipson and Drake, 2011). By 2018, eight immunotherapies had been FDA approved for the treatment of several cancers (Table 2), including durvalumab (stage 3 lung cancer), blinatumomab (acute lymphoblastic leukemia), and nivolumab (used in combination with ipilimumab for previously untreated kidney cancers) (https://www.cancer.gov/news-events/cancer-currents-blog/fda-approvals).

**TABLE 2**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Mechanism of Action</th>
<th>Targeted Disease</th>
<th>Release Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Durvalumab (IMFINZI, AstraZeneca)</td>
<td>Checkpoint immunotherapy targeting PD-1/PD-L1</td>
<td>Stage III non–small-cell lung cancer</td>
<td>February 16, 2018</td>
</tr>
<tr>
<td>Brentuximab vedotin (Adcetris, Seattle Genetics, Inc.)</td>
<td>Antibody-drug conjugate targeting the CD30 receptor</td>
<td>Untreated classic Hodgkin lymphoma</td>
<td>March 20, 2018</td>
</tr>
<tr>
<td>Blinatumomab (Blincyto, Amgen Inc.)</td>
<td>Bispecific T cell-engaging antibody targeting CD19 receptor</td>
<td>Acute lymphoblastic leukemia</td>
<td>March 29, 2018</td>
</tr>
<tr>
<td>Pembrolizumab (Keytruda, Merck)</td>
<td>Checkpoint immunotherapy targeting PD-1 and PD-L1</td>
<td>Melanoma</td>
<td>April 16, 2018</td>
</tr>
<tr>
<td>Pembrolizumab (Keytruda, Merck)</td>
<td>Checkpoint immunotherapy targeting PD-1 and PD-L1</td>
<td>Relapsed refractory large B-cell lymphoma</td>
<td>May 31, 2018</td>
</tr>
<tr>
<td>Pembrolizumab (Keytruda, Merck)</td>
<td>Checkpoint immunotherapy targeting PD-1 and PD-L1</td>
<td>Relapsed/refractory cervical cancer</td>
<td>June 12, 2018</td>
</tr>
<tr>
<td>Pembrolizumab (Keytruda, Merck)</td>
<td>Checkpoint immunotherapy targeting PD-1 and PD-L1</td>
<td>Relapsed/refractory metastatic melanoma</td>
<td>June 13, 2018</td>
</tr>
<tr>
<td>Pembrolizumab (Keytruda, Merck)</td>
<td>Checkpoint immunotherapy targeting PD-1 and CTLA-4</td>
<td>Relapsed/refractory small-cell lung cancer</td>
<td>July 10, 2018</td>
</tr>
<tr>
<td>Pembrolizumab (Keytruda, Merck)</td>
<td>Checkpoint immunotherapy targeting PD-1 and CTLA-4</td>
<td>Metastatic melanoma</td>
<td>August 17, 2018</td>
</tr>
</tbody>
</table>

Immunotherapeutic approaches include the use of targeted antibodies and vaccines against immune checkpoint inhibitors directed toward a specific immune cell population (Ventola, 2017). For instance, due to their antigen-presenting capabilities, dendritic cells (DCs) have been used to develop immune vaccines (Sabado et al., 2017). Depending on the molecules used to activate them, DCs are able to reprogram or launch a cell-specific cytotoxic response. Conversely, tumor-associated macrophages (TAMs) have been shown to exert different roles in tumor microenvironment development and flourishing (Mills et al., 2016). Approaches that target this macrophage population are currently being evaluated, especially since the discovery that the blockade of TAMs potentiates immune checkpoint inhibitors’ effect (Ries et al., 2014; Zhu et al., 2014). Adoptive T-cell therapy to re-engineer the T-cell populations against tumor initiation is another strategy that has been widely validated (Dzhandzhugazyan et al., 2018). The chimeric antigen receptors re-engineered T cells (CAR-T) system has been recently approved by the FDA for the treatment of patients with leukemia, large B-cell, and non-Hodgkin
lymphomas (Zheng et al., 2018). Other focuses involve the use of a different immune cell population, the natural killer T cells. Natural killer T cells naturally stimulate the innate and adaptive immune system in several ways, such as the release of interferon-γ to activate the CD8+ T-cell population (Mah and Cooper, 2016). They are being investigated as potential immunotherapies both as ex vivo expanded cell vaccines and as combinatorial therapies (Nair and Dhodapkar, 2017).

**Ovarian Cancer: A “Cold” Enemy**

The characterization of the topographic distribution of immune cells within the tumor in a panel of 177 human samples with different cancer types has recently led to their categorization as inflamed (“hot”), noninflamed (“cold”), and “immune excluded” patterns according to where the cells are positioned (Kather et al., 2018). Cold tumors are malignancies that display a very limited response to immunotherapies compared with other cancer types. OC is considered a “cold” tumor (Preston et al., 2011) despite the significant association between tumor immunity and ovarian patient outcomes and the strong correlation between the presence of infiltrating lymphocytes in the primary tumor and patient survival (Zhang et al., 2003). The reasons behind this lack of effectiveness have yet to be clarified. A possible explanation, proposed for pancreatic cancer, suggests that the difference between hot and cold tumors reflects the way tumor-infiltrating immune cells are recognized by cancer cells or engage in the tumor. If so, the properties of the microenvironment make a tumor hot or cold. Hot tumors are more sensitive to treatments that activate the T-cell population, as they are considered to be the main drivers of the adaptive immune response against tumor initiation (Haanen, 2017).

The tumor microenvironment is a complex hub where different cell types interact with each other and with the extracellular matrix, and it is plausible that other cells, including antigen-presenting cells (APCs), play an active role in downregulating the immune system. APCs, including the aforementioned DCs, are highly responsive to external stimuli, and the tumor surroundings can negatively affect their physiologic behavior. Indeed, it has been shown that endoplasmic reticulum stress is also crucial for triggering cancer resistance mechanisms by activating the unfolded protein response, which in turn disrupts the physiologic immune response (Yadav et al., 2014). Specifically, through the constitutive activation of the endoplasmic reticulum stress response factor XBP1, DCs undergo an abnormal lipid accumulation that leads to their ineffective functioning (Cubillos-Ruiz et al., 2015). While low infiltration of immune cells both inside and outside the tumor is found in OC samples, the coexistence of different immune microenvironments within the same patient partly explains the heterogeneity in the response to treatment often observed in patients with recurrent disease (Jiménez-Sánchez et al., 2017).

Currently, there are no FDA-approved immunotherapies for OC, although there are several ongoing clinical trials. Of the 98 total clinical trials, 26 have been completed, 40 are actively recruiting patients, and nine have been terminated before their planned end due to the inefficacy determined by the limitations described in the previous paragraphs (https://clinicaltrials.gov). In 2016, Gaillard et al. reported a comprehensive analysis of all clinical trials on checkpoint inhibitors,

![Fig. 1. Schematic of the current immunotherapies for ovarian cancer. Arrows show the interactions between immune system players (dendritic cells, T cells, macrophages, and monoclonal antibodies) and ovarian cancer cells. Each specific immune cell type can be used to deliver specific therapies that can differently alter the immune system toward a more efficient activity rate. EpCAM, epithelial cell adhesion molecule; VEGF, vascular endothelial growth factor.](image-url)
and discovered that, on average, the efficacy of these treatments is surprisingly poor in OC patients (Gaillard et al., 2016). The positive outcome was found to be around 10%–15%. A schematic representation of the immunotherapy-based approaches used in OC and the interactions between different immune players and tumor cells is provided in Fig. 1.

**Monoclonal Antibodies in OC.** In the attempt to enhance treatments for OC, a number of monoclonal antibodies capable of inhibiting the function of molecules involved in tumor progression and immune suppression. Catumaxomab is a monoclonal antibody directed against the epithelial cell adhesion molecule, a glycoprotein highly expressed in OC (Tayama et al., 2017). This antibody is currently being evaluated in a phase II clinical study on patients resistant to chemotherapy (Berek et al., 2014). Following the identification of cancer antigen 125, which is the most renowned OC marker (Bast et al., 1981), its role as a suppressant of both natural killer cell activity (Patankar et al., 2005; Tyler et al., 2012) and antibody-dependent cellular toxicity (Kline et al., 2017) has been widely investigated. Several anti–cancer antigen 125 monoclonal antibodies have been developed and tested, including oregovomab (Berek et al., 2009) and abagovomab (Sabbatini et al., 2013), although they did not prove to be effective in improving the outcomes in advanced OC when used as a single-agent maintenance immunotherapy. Anti–CD25 (daclizumab) has been clinically tested for its capacity to suppress the T regulatory cell populations, which are responsible for shorter patient survival rates when infiltrated within the tumor (Barnett et al., 2010). Although the trial has been completed, the results have not been released yet. The translational potential of anti-CD25–based platforms is limited by their nonspecific binding, as CD25 is widely expressed on T-cell populations.

**Dendritic Cell Vaccines in OC.** Dendritic cells have a pivotal role in launching the immune response due to their capacity for activating CD4+ or CD8+ T cells (Sallusto and Lanzavecchia, 2002). Their role in the tumor microenvironment is the subject of active contemporary research (Pörschke et al., 2017). As plastic APCs, DCs are currently harnessed for their potential to boost the immune system against tumor initiation and progression. Scarlett et al. (2012) applied an inducible p53-dependent model of aggressive ovarian carcinoma to demonstrate that DCs display differential immunostimulatory capacity during tumor initiation and escape. These changes correspond to significantly lower levels of major histocompatibility complex II (MHC-II) and CD40 on their surface. DCs are tunable cells, capable of either inducing an immune surveillance effect or releasing malignant growth by activating or suppressing antitumor T-cell activity, respectively. DC-based vaccines have also been conceived in the context of OC, by ex vivo pulsing DCs with tumor-derived components, as single tumor-associated peptides or peptide combinations (Liao and Disis, 2013). Cancer testis antigens (CTAs) that are typically expressed in multiple types of tumors have also gained interest for their potential applicability in immunotherapy (Gjerstorff et al., 2015; Seifi-Alan et al., 2018). NY-ESO-1, a member of the CTA family, has been used to produce either DC-based vaccines (National Clinical Trial [NCT] number NCT02387125) or adoptive T-cell therapies (NCT number NCT01567891). Similar immunotherapeutic approaches are being developed using melanoma antigens (i.e., MAGE-A1, MAGE-A4, MAGE-A3, and MAGE-A10) that represent another subgroup of the CTA category (Daudi et al., 2014). Zitvogel et al. (1996) are among the first researchers to use tumor antigen-pulsed DCs to treat mice with fibrosarcoma. They also demonstrated that patient-derived DCs pulsed with a cocktail of tumor antigens [whole tumor antigen (WTA)] can trigger a tumor growth suppression through the activation of CD4+ and CD8+ T cells when reintroduced into the patient. In their study, the activation of T cells correlated with a better prognosis in patients with recurrent ovarian, fallopian tube, or primary peritoneal cancer (Kandalaf et al., 2013; Tanyi et al., 2018). Recently, a pilot study demonstrated an autologous WTA-pulsed DC-based vaccine to be safe and effective in combination with cyclophosphamide and bevacizumab (Tanyi et al., 2018). By priming DCs with patient-derived WTA, Tanyi et al. (2018) were able to overcome two of the limitations associated with the use of immunotherapy for the treatment of OC: the lack of an efficient antigen-specific active treatment and the inability of tumor-specific T cells to home to tumors.

**Adoptive Cell Therapy in OC.** Adoptive cell therapies show potential for the treatment of OC. For example, it has been shown that tumor-infiltrating lymphocytes derived from OC biopsy-derived cells can be expanded ex vivo and be reactivated to produce antitumor cytokines (Owens et al., 2018). Similarly, the abundance of tumor-infiltrating lymphocytes in patients’ ascitic fluid has prompted their evaluation as reintegrable immunotherapies after their demonstrated cytotoxic effect on tumor cells (Abe et al., 2018). CAR-T–based therapy produced by combining programmable antigen receptor specificity with T-cell activation also holds an attractive opportunity for the treatment of OC (Dzhandzhugazyan et al., 2018). The lack of a demonstrable efficacy of this approach is mainly due to the poor T-cell trafficking and the immunosuppressive microenvironment (Zhang et al., 2016; Mirzaei et al., 2017; Jindal et al., 2018). Despite the potential pitfalls of this approach, clinical trials evaluating its efficacy in OC are currently active and specifically target mesothelin (NCT02580747, NCT01583686, MUC16 (NCT02498912), HER2 (NCT01935843), and NY-ESO-1 (NCT02366546), among others (Zhu et al., 2017).

**Nanomedicines and Immunotherapy in Ovarian Cancer**

Synthetic and natural nanotechnologies are currently being investigated to deliver immunotherapies, as they have the potential to improve patient treatment outcomes and reduce mortality rates (Shen et al., 2017). This includes the use of nanoparticles for the delivery of immunostimulatory and immunosuppressive molecules in combination with chemoradiotherapy or as adjuvants to other immunotherapies (Sapiezynski et al., 2016). Nanoparticles have also been designed to produce vaccines to stimulate T-cell response against tumor growth (Fan and Moon, 2015), allowing for the codeelivery of antigen and adjuvants (Dunkle et al., 2013), contributing to the inclusion of multiple antigens to activate DC targets (Xia et al., 2015), and guaranteeing the sustained release of antigens for a prolonged immune stimulation (Engelberth et al., 2014).

The literature reports only a few examples of preclinical studies investigating the potential of nanotechnology-based platforms to improve the outcome of immunotherapeutic
regimens in OC. These include polymeric nanoparticles (Cubillos-Ruiz et al., 2009; Hanlon et al., 2011; Ortega et al., 2015; Teo et al., 2015), liposomes (Turk et al., 2004; Rajan et al., 2018), and lipid–polymer hybrids (Anwer et al., 2013).

Nanoplatforms for OC have been synthesized primarily to guide the delivery of RNA oligonucleotides to target cells, thus overcoming the current limitations related to the use of RNA therapeutics. Limitations include low bioavailability, poor cellular uptake, cytotoxicity, and the need to evade the phagocytic cellular components of the immune system (Kole et al., 2012). Polymeric nanostructures have been developed to provide additional control over drug release at tumor sites, as they offer the advantage of being able to respond to specific stimuli provided by the tumor environment, such as pH and enzymatic activity (Uthaman et al., 2018). Among the many polymers available, polyethylenimine (PEI) is one of the most used materials in OC treatment, as it is considered a versatile gene carrier (Teo et al., 2013). PEI displays high efficacy for siRNA encapsulation and delivery for both in vitro and in vivo purposes. Its cationic charge enables the loading of siRNA into nanocomplexes and protects it from enzymatic degradation (Zheng et al., 2011; Höbel and Aigner, 2013). The abundant presence of amine groups allows for the functionalization of the platform and favors further modifications of this polymer to improve the bioactive features, such as its targeting ability and cell specificity. Cubillos-Ruiz et al. (2009) investigated PEI-siRNA nanoparticle uptake by tumor-associated DCs and its effect in reprogrammng their phenotype from immunosuppressive cells to efficient APCs. The authors found that the changes induced in DCs through the use of PEI-siRNA against immunosuppressive determinants consequently activated tumor-reactive human and murine lymphocytes and exerted a direct tumoricidal activity in aggressive ovarian carcinoma-bearing murine models (Cubillos-Ruiz et al., 2009). The induced T cell–mediated tumor regression and prolonged survival were dependent upon the activation of the myeloid differentiation primary response gene 88 (MyD88). PEI alone was sufficient to mediate the upregulation of MHC-II, MHC-I, and costimulatory molecules in tumor DCs in vivo. This suggests that the intrinsic stimulation of Toll-like receptors 5 and 7 by PEI nanoparticles synergizes with the gene-specific silencing activity of the siRNA to transform tumor-infiltrating regulatory DCs into cells capable of promoting therapeutic antitumor immunity. Cubillos-Ruiz et al. (2012) further optimized the platform to achieve the synthetic enhancement of the specific molecular pathway miR-155 signaling in DCs. This pathway is responsible for boosting a potent antitumor immune response that abrogates the progression of established ovarian cancers (Cubillos-Ruiz et al., 2012). Other researchers have taken advantage of polymeric nanoparticles’ capability to be functionalized, thus improving targeting and, consequently, the therapeutic outcome. By applying a different immunotherapy-based approach, Teo et al. (2015) proposed various folic acid (FA)–functionalized PEI polymers to block programmed death-ligand 1/PD-L1 interactions by delivering PD-L1 siRNA to human epithelial ovarian cancer (EOC) cells (SKOV-3 line) and to sensitize them against T cells. With their hypothesis to target PD-L1, the authors responded to the need for a specific targeted delivery of PD-L1 siRNA to epithelial cancer tissues, as PD-L1 is also expressed on healthy tissues (Liang et al., 2003), including placenta and eyes. The polymer/siRNA nanocomplexes knocked down PD-L1 on a luciferase-expressing SKOV-3, enhancing the efficacy of T-cell immunotherapy for the treatment of EOC compared with the respective PEI–FA and PEI–polyethylene glycol (PEG)–FA/scrambled siRNA treated controls. These data highlight the potential use of PEI–FA as specific gene delivery carriers. The modification of PEI with FA or PEG–FA proved to be a valuable tool to reduce cytotoxicity while improving tumor cell targeting toward EOC cells and uptake, with a striking 40%–50% knockdown of PD-L1 expression. Ortega et al. (2015) used click chemistry to produce nanoparticles based on 2-(dimethylamino)ethyl methacrylate polymer further functionalized with the mannose ligand (MnNP). This platform was meant to condense siRNA against the polyoma middle T oncogene and specifically target the mannose receptor (CD206) present on the surface of TAMs (Ortega et al., 2015). MnNP has been demonstrated to be biocompatible in both in vitro and in vivo settings. MnNP is also able to efficiently incorporate and deliver functional siRNA into the cytoplasm of TAMs. This study provides evidence that mannosylation is responsible for TAM selectivity in vivo following intraperitoneal injection with a twofold increase in TAM uptake compared with nontargeted particles and about a 10-fold increase compared with nonmyeloid cells. In this study, the spatial confinement of the MnNP within the peritoneal cavity enhanced the opportunity for the interaction with immune cells associated with OC, and the biodegradability of the system ensured the persistence of the treatment for over 24 hours.

Poly(lactic-co-glycolic acid) nanoparticles (PLGA-NPs) are biodegradable, and their composition can be tuned to temporally control the release of the payload (Corradetti et al., 2012; Minardi et al., 2016). PLGA-NPs have been used as an alternative route to deliver whole WTA to DCs since the injection of soluble antigens presents inherent limitations due to instability and poor internalization rates. These factors result in the transient and inefficient activation of T cells (Hanlon et al., 2011). At the same time, PLGA-NPs protect antigens from enzymatic degradation and maintain their bioactivity, leading to a more efficient presentation of MHC-peptide complexes by recipient cells following uptake and processing. In vitro studies have confirmed the effectiveness of PLGA-NPs in the activation of a CD8+ cell response characterized by a significant increase in the production of inflammatory cytokines and a greater expression of costimulatory molecules, providing encouraging evidence for their potential clinical translation. Interestingly, the delivery of WTAs through PLGA-NPs appeared to facilitate the antigens’ access to the MHC class I compartment in the cytoplasm, providing a reservoir for a prolonged and enhanced Ag presentation.

Liposomes are small, artificial, spherical vesicles synthesized primarily from natural nontoxic phospholipids (Akbarzadeh et al., 2013). Their wide application as drug delivery systems in biomedical settings is due to their biocompatibility, biodegradability, low toxicity, and capability to load both hydrophobic and hydrophilic drugs (Johnston et al., 2007). Moreover, liposomal encapsulation offers the advantage of effectively enhancing the solubility of lipophilic and amphiphilic drugs and improving site-specific drug delivery to tumor tissues through surface functionalization (Hofheinz et al., 2005; Corradetti et al., 2012). The latter aspect is crucial to increase the retention time, which can be modulated by drug-lipid interactions, and permit the accumulation of
liposome-encapsulated chemotherapeutic agents at the tumor site (Deshpande et al., 2013).

Doxil (Ben Venue Laboratories) was the first pegylated liposome-based drug to enter the market in 1995. The nanoparticle formulation includes doxorubicin, a DNA intercalating agent used against a variety of cancers, including gynecologic cancers (Howard et al., 2016). While no significant differences were observed in terms of efficacy compared with the free drug, the liposomal formulation allowed the reduction of cardiotoxicities related to the use of doxorubicin and the preferential accumulation of the drug at the tumor site (Green and Rose, 2006). More recently, the FDA approved the use of an RNA interference therapeutic delivered by lipid nanoparticles: patisiran (Adams et al., 2018). Although developed for the treatment of degenerative diseases, patisiran shows promise as a new breakthrough in patient care, as it heralds the arrival of an entirely new class of medicines to treat human diseases. However, despite the wide interest in the use of liposomal formulations for OC treatment, only one group has tested liposomes as nanocarriers for immunotherapy. Turk et al. (2004) developed folate-conjugated liposomes to target intraperitoneal ovarian carcinoma cells, as they over-express the folate receptor. Data revealed that this formulation was also absorbed by TAM through folate receptor-mediated internalization, with a 10-fold increase in the engulfment of macrophages compared with ascitic tumor cells in vivo, corroborating the need to develop combinatorial strategies aimed at modulating TAM and inhibiting cancer cell growth. Lipid–polymer hybrid nanoparticles (LPNs) are core–shell nanoparticle structures constituted by a polymeric core and a lipid shell. LPNs have been considered by other researchers to confer a high degree of physical stability to the platform, resulting in a superior in vivo cellular delivery efficacy (Hadinoto et al., 2013) compared with polymeric and liposomal nanoparticles. The combination of the two LPN platforms formulated with a lipopolymer PEG–PEI–cholesterol was used as an effective tool to deliver an interleukin 12 plasmid at the tumor site. Interleukin 12 was chosen for the therapeutic action it plays in OC, which relies on its potential to activate the antitumor immunity (Whitworth and Alvarez, 2011). This approach proved to be safe and effective in platinum-sensitive OC patients treated with intravenous carboplatin and docetaxel (Anwer et al., 2013).

Physical and Biologic Barriers Challenging the Treatment of OC

Innovative immunotherapeutic-targeted strategies mediated by nanotechnology offer the promise of enhancing host antitumor responses which may improve clinical outcomes in women with OC. Although preclinical studies have demonstrated the induction of an antitumor response, there is no clinically effective nanomedicine-based immunotherapy available for OC patients. The biologic barriers that physically and mechanically influence the processes involved in tumor spread and immune cell infiltration must be considered when developing new strategies for the treatment of OC. As mentioned earlier, one of the main mechanisms by which OC cells spread is through transcoelomic metastasis, which involves dissemination throughout the peritoneal cavity (Tan et al., 2006). Ascite formation is determined by the accumulation of cancer cells, growth factors, and immunosuppressive ligands (vascular endothelial growth factor and fibroblastic growth factor β), which increase peritoneal capillary permeability (Ahmed and Stenvers, 2013) and thus the leakage of plasma proteins (i.e., albumin, fibrin, and fibrinogen) from newly developed vessels (Stanoevic et al., 2004). The obstruction of lymphatic vessels by cancer cells also occurs, which leads to an impaired reabsorption of the physiologic peritoneal fluid (Kripps et al., 2013). As a consequence of the compromised lymphatic drainage of the peritoneal cavity, fluid confinement in the peritoneum occurs, contributing to the pathogenesis of malignant ascites. The environment that these biologic and physical processes create impedes immune cell migration and infiltration within the metastatic tumors (Cai and Jin, 2017) and induces a peripheral tolerance that attenuates their function (Kulshrestha et al., 2017). For instance, ascites proved to recruit and immunologically suppress a population of neutrophils through cell contact in a cohort of newly diagnosed OC patients (Singel et al., 2017). The release of macrophage migration inhibitory factor from ascite-derived cancer cells has also been proposed to halt the tumor-killing ability of NK cells by transcriptionally downregulating the expression of the surface receptor NKG2D (Krokenberger et al., 2008). These findings confirm the proactive role of malignant ascitic fluid in physically supplying cells and chemical stimuli to favor an immune-suppressed environment. Additionally, another physical barrier to immune cell penetration is represented by the tumor vascular endothelium (Motz and Coukos, 2013). In a physiologic environment, the presence of adhesion molecules, such as intercellular cell adhesion molecule or vascular cell adhesion molecule, allows T cells to adhere to and travel through the endothelium. In the tumor milieu, the release of angiogenic growth factors prevents T cells from passing through by inhibiting the adhesion molecules’ expression (Bouzin et al., 2007).

Exosomes: An Alternative Tunable and Nanoscopic Strategy

Recently, biologic nanoparticles (called exosomes) have also emerged as a powerful translational platform to be harnessed in the development of naturally inspired delivery systems. Exosomes are nanoscopic lipid vesicles with a size range spanning from 30 to 150 nm that are released by cells and thus retain their bioactive moieties. Due to their small size and architecture, exosomes can penetrate across the lymphatic vessels and tumor interstitium and reach target organs (Srinivasan et al., 2016). Their composition and cargo can be further modified by conditioning parental cells or by improving their natural potential with the addition of functional drugs, thus giving them additional functions (Conlan et al., 2017). Exosomes play a crucial role in cell-to-cell communication and are characterized by a precise targeting potential that allows for the activation or repression of specific molecular
cascades in targeted cells (Syn et al., 2017). Currently, their role in the exchange of information between the tumor and the surrounding microenvironment is being explored (Maia et al., 2018), as is their potential as delivery vessels for both therapeutic and imaging purposes (Luan et al., 2017; Shen et al., 2018).

Recent advances in the field of immunotherapy unveiled the role of appropriately stimulated exosomes released from cancer cells as potent endogenous nanocarriers responsible for the suppression of T cells and the facilitation of tumor growth (Chen et al., 2018). Once injected for therapeutic purposes, exosomes are not susceptible to further modifications determined by the microenvironment, offering a great advantage over the use of CAR-T cells or DCs, which are amenable to acquiring a different phenotype (Yamashita et al., 2018).

Interestingly, they have also been proposed as useful tools to predict the patient response to immunotherapy. On the other hand, exosomes derived from immune cells, APCs, or TAMs are now at the forefront for the development of innovative vaccine strategies for cancer immunotherapy against tumor initiation and are the subject of current clinical trials for the treatment of other tumor types (Hong et al., 2017; Liu et al., 2017).

Conclusions and Perspectives

In this review we discussed the widely recognized impact of immunotherapy in the treatment of cancers, highlighting the challenges researchers face in the effort to overcome the limitations provided by OC. These include its cold nature, determined by the immunosuppressive environment and the transport oncophysics, which urgently calls for the conception of alternative approaches to deliver immunotherapies. Ideally, these approaches are meant to preferentially accumulate the drug at the tumor site; sustain the temporal and spatial release of the payload, thereby reducing cytotoxicity; and selectively target specific cell types to stimulate antitumor immunity (Fig. 2). Nanotechnology offers advantageous drug delivery systems with demonstrated therapeutic efficacy, with a direct or indirect effect on cancer cells. However, the potential of nanomedicines for the treatment of OC has been harnessed to a limited extent. Although capable of identifying and targeting the cell population of interest, none of the nano-enabled strategies proposed have yet shown significant clinical benefits. Furthermore, the literature lacks a comprehensive discussion about the in vivo biodistribution of the proposed nanoplatforms, reinforcing the concept that the drastic changes within the peritoneal cavity in terms of transport oncophysics and metastases heterogeneity largely limit their capability to reach tumor masses. A deep understanding of the role exosomes play in traveling and mediating cell interaction within the OC environment will successfully lead to the development of cutting-edge approaches to prime the body’s immune system against tumor initiation. The continuous advancements in the field of nanotechnology will provide the tools needed to synthesize exosome-resembling particles to be used as alternative immunotherapy treatment of OC. Another approach may include the coupling of naturally derived exosomes with established multistage vectors, demonstrated to achieve efficient delivery of chemotherapeutics to metastatic breast cancer (Xu et al., 2016) and ovarian tumor tissues (Shen et al., 2013). The possibility to exploit the physical properties of the ascitic fluid and the geometry of the peritoneal cavity during metastatic OC to tailor the architecture of multistage vectors paves the way for the fabrication of nanotechnology-based immunotherapies to accomplish the challenge of boosting...
the anticancer immune system and minimizing tumor relapse.

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