Current Preclinical Models of Ovarian Cancer

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Abstract

Despite improvements in surgical and chemotherapeutic intervention of ovarian cancer over the recent decades, ovarian cancer remains the most lethal cancer in women. Notably, after an initial effective response to chemotherapeutic regimen, therapeutic resistance rises up leading to patient’s death. This scenario highlights the urgent need to develop novel diagnostic and therapeutic strategies. Recently, several efforts to better understand the molecular bases of ovarian cancer using integrated multipletumor molecular profiling have revealed an intrinsic complexity and heterogeneity among ovarian cancers. Concurrently, a growing body of evidences implies fallopian tube epithelium as the likely site of origin of the majority of ovarian cancers. This fallopian tube hypothesis has shifted the attention of ovarian cancer research from the ovarian surface epithelium to the fallopian tube epithelium leading to adjustment of in vitro and in vivo ovarian cancer models. In this review article, we critically summarize recent advances in ovarian cancer preclinical models that have the potential to accelerate and facilitate the discovery of more effective biomarkers and target drugs for personalized cancer therapy.

Keywords: Ovarian cancer; Cell models; Mouse models; Xenografts; Fallopian tube

Abbreviations:
EOC: Epithelial Ovarian Cancer; HGSC: High-Grade Serous Carcinoma

Introduction

With approximately 22000 new cases diagnosed and 14000 deaths each year in United States, epithelial ovarian cancer (EOC) is the most lethal gynecological tumor worldwide [1]. High-grade serous carcinoma (HGSC) represents more than 70% of all EOCs and it is responsible of about 90% of EOC-related deaths. Recent clinical-therapeutic innovations improved the disease free survival of patients with HGSC, without impacting on their overall mortality rate [2,3]. This is largely due to the complete darkness on HGSC carcinogenesis and, in particular, on the initial pathogenetic steps. As a consequence, HGSC is usually diagnosed at advanced stages, while precursor lesions remain elusive. The current therapeutic approach for HGSC contemplates surgery either anticipated or followed by the administration of platinum-taxane combined chemotherapy and is associated with good immediate response, but high rate of recurrence, locally and systemically. Notably, HGSC is insensitive to standard targeted therapy options, such as hormonal therapy and anti-HER2 agents, currently available for breast cancer patients [4,5]. Therefore, the most daunting challenge in ovarian cancer research is to develop new effective therapy to prevent and treat HGSC recurrence.

It is emerging that the five main histotypes of EOC (Table 1) must be considered as distinct diseases based on cell of origin, pathogenesis and molecular alterations [6-15]. In this respect, the effort of the scientific community to unravel the molecular alterations of HGSC (one of the first goal of The Cancer Genome Atlas consortium; TCGA), has allowed the identification of only few common somatic mutations, with the only exceptions of TP53 and BRCA1-2 genes, and none new common “druggable” candidate [16]. In addition, although the ovarian surface epithelium has been considered for long time the primary site of origin of all (both benign and malignant) epithelial ovarian tumors, the origin of EOC is still debated with an increasing consideration of extra-ovarian origin [6,10,11]. In particular, emerging evidences indicate the fallopian tube epithelium (FTE) and the endometrium as the sites of origin of ovarian HGSC and endometrioid/clear cell carcinoma, respectively [6]. Therefore, this new conceptual framework has shifted the attention of ovarian cancer research outside the ovary, from the ovarian surface epithelium to the FTE, renewing the interests in refining in vitro (cell cultures) and in vivo (animals) HGSC models (Figure 1). Experimental models used in ovarian cancer research substantially evolved over the last few years. As a general concept, the ability of the experimental models to accurately recapitulate the complexity of human cancer represents a critical issue in preclinical studies for drug discovery. The high rate of failure of novel cancer therapeutics during clinical trials highlights the inadequate predictability of laboratory cancer models currently available for preclinical studies.

In the present review, we attempt to summarize the current status of the art of in vitro and in vivo models of HGSC that better recapitulate various features of this cancer, focusing on their advantages and limitations.
<table>
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<th>Histotype</th>
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<th>Frequency (%</th>
<th>Pathogenesis</th>
<th>Cell of origin</th>
<th>Precursor lesion</th>
<th>Common somatic mutations</th>
<th>Type</th>
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<td>FTE</td>
<td>Benign Brenner tumor</td>
<td>KRAS, PIK3CA</td>
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<td>Stepwise/ De novo</td>
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<td>Endometrium</td>
<td>Endometriosis, Borderline tumor, Adenofibroma</td>
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<td>Endometriosis, Borderline tumor, Adenofibroma</td>
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<td>De novo</td>
<td>FTE</td>
<td>Serous tubal intraepithelial carcinoma</td>
<td>BRCA1, BRCA2, TP53</td>
<td>II</td>
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Table 1: Principal histotypes of ovarian epithelial tumors.

**Cell Models**

**Primary cell cultures**

Short-term cultures derived from freshly isolated cells or tissues, also designed primary cultures, have many important applications since they may recapitulate the pathophysiological system closely. However, primary cell cultures display common limiting characteristics: the slow growth capacity, the limited overall lifespan (i.e. they are not able of indefinite serial ex-vivo propagation) and the occurrence of changes and/or selection over passages, which hamper the system reproducibility. Other intrinsic limitations of primary cell culture system are that the tissue histology is lost along with endocrine, paracrine and neural regulators, gradients of nutrients and other factors. The cell cultures of interest to study HGSC biology are those derived from FTE as well as from ovarian cancers.

**Primary cell cultures generated from fallopian tube epithelium:** Human fallopian tube is gaining progressive attention, given its proposed role as the likely site of origin of HGSC. The FTE consists of a simple columnar epithelium composed of ciliated, secretory and intercalated cells [10]. During the last three decades methods for the in vitro expansion of FTE cells have been established [17-20]. The epithelial cells lining the inner surface of the human fallopian tube can be cultured in vitro and kept in culture for 6-8 weeks [19,20]. New epithelial cells appear after 2-3 days of primary cultures and small clusters after 7-10 days [19,20]. The plating and subculturing efficiency of FTE cells is very low, and the risk of contamination by fibroblasts is high [19,20]. Furthermore, serial cell passages result in loss of the ciliary markers of differentiation and in senescence over time [21]. The observed phenotypical plasticity depends strictly on culture conditions. As alternative experimental approach, cultures of FTE have been recently obtained seeding the isolated cells on the top of transwell filters coated with human placental collagen [22]. This model recapitulates faithfully the histological features of normal human FTE, preserving both morphological and phenotypical cell characteristics, although still characterized by finite proliferative capacity [23]. In order to overcome the proliferation limits, two research groups independently immortalized and transformed ex vivo normal human FTE cells by using human telomerase reverse transcriptase (hTERT) plus SV40 large T antigen and by ectopic expression of either oncogenic H-RasV12 or c-Myc, alone or in conjunction with inhibition of p53 and Rb tumor suppressor pathways [23,24]. The transformed
cells, injected in mice, produced tumor phenotypically and
genotypically resembling human HGSC. In an analogous study,
normal FTE cells were immortalized and transformed by ectopic
expression of oncogenic H-Ras V12 [25]. These cells were tumorigenic
when injected in immunocompromised mice, but at the histological
examination the resulting tumors were mostly poorly differentiated
mucinous adenocarcinoma mixed with undifferentiated carcinoma.

**Primary cell cultures generated from ovarian cancer:** Several
methods for the isolation and culture of primary EOC cells, derived
from either fresh solid tumor or ascites liquid, have been described
[26,27]. Primary EOC cells usually adhere and tend to reach
confluence quickly [26,27]. They can be kept in culture for 2-3 months
before going into senescence. Short-term proliferative cells can be used
for immediate experiments or cryostored, and represent a unique
model suited for a multitude of applications [26,27]. Previous
chemotherapy may affect cellularity and cell viability, therefore HGSC
cells that have received recent chemotherapy have less growing
efficiency *in vitro* [26,27].

**Ovarian cancer cell lines**

Cancer cell lines are the model most commonly used in cancer
research and their use has undoubtedly altered our understanding of cancer biology [28,29]. An established cell line refers to
a population of cells which has been serially passaged at least 60
times *in vitro*, and that is easily maintained *in vitro* and cultured for
long periods of time. The Broad-Novartis Cancer Cell Line
Encyclopedia and the TCGA projects together reported the genomic
profiles of more than 1000 cell lines from various cancer lineages,
including ovarian cancers [30,31]. Generally, cancer cell lines possess
the same spectrum of genetic aberrations as primary tumors, although
each cell line presents only a limited number of genetic aberrations,
given that each cell line represents the intertumoral heterogeneity
observed among primary tumors [30,31]. The cell immortalization
leads to selection of a cell population phenotypically homogeneous
and genetically clonal. Thus, a well-known caveat of cell lines is that
they possess distinctive molecular genetics alterations driven by
immortalization. It is therefore of extreme importance to confirm that
these immortalized cell lines represent faithfully primary tumors with
respect to original genomic alterations, since genomic alterations may
result in molecular characteristics predictive of response (sensitivity vs
resistance) to specific therapeutics. So far, numerous EOC cell lines
have been established, and commercialized, but the histopathological
origin of the most commonly used EOC cell lines, namely A2780,
CAOV3, IGROV1, OVCA-3 and SK-OV-3, remains unclear. As
reported in a study performed by TCGA network investigators, a
comparison between the molecular alterations of 47 EOC cell lines and
those of 316 primary HGSCs revealed that the EOC cell lines
commonly used by scientists do not recapitulate the molecular
alterations identified in primary HGSC [28]. This may explain the
reason why *in vitro* studies using EOC cell lines usually fail to correctly
predict clinical response and indicates that several EOC cell lines
cannot be considered reliable *in vitro* HGSC models. In the same study
[28], EOC cell lines were ranked based on the molecular similarity
to primary HGSCs. The top ranking cell lines, characterized by major
genomic similarity to primary HGSC are: KURAMOCHI, OVSAHO,
SNU119, COV362 and OVCA-4 [28]. Therefore, these EOC cell lines
can be considered the most suitable models for preclinical studies of
HGSC.

**Ovarian cancer stem cells**

The cancer stem cell or tumor-initiating cell theory states the
existence of a discrete population of tumor cells able to reconstitute
the tumor upon implantation into recipient mice, by virtue of self-
renewal and multipotency abilities [32,33]. This theory holds relevant
implications for cancer therapy since the cancer stem cells sit in a
quiescence status that protects them against various chemotherapeutics. Hence, cancer stem cells are responsible for
minimal residual disease and cancer recurrence [34].

Many studies have isolated, via fluorescent-activated cell sorting
and characterized alleged human ovarian cancer stem cells from
primary HGSC using surface antigens, such as ALDH1, CD44, CD133
and CD117, that define stem cell-like phenotype in other cancers
[35-38]. The aforementioned isolated cells could fully recapitulate
parental HGSC in xenografts and exhibited enhanced capacity to form
spheroids and chemoresistance to conventional HGSC chemotherapeutics [35-38]. Interesting, Steffensen et al. found that
HGSC patients with higher percentage of CD44 positive cancer stem
cells had shorter progression-free survival [39]. Similarly, two other
studies have reported that HGSC patients with increased ALDH1
expression showed poor prognosis [40,41]. In the same way, CD133
expression, alone or in combination with ALDH1, is a negative
prognostic factor in HGSC patients [36,42]. Recently, Schwede et al.
have identified a stem-cell line gene expression signature in a subset of
HGSCs with higher propensity to form spheroids *in vitro*, to
metastasize *in vivo* and with poorer prognosis [43]. Overall these
findings support the existence of cancer stem cells in HGSC and
emphasize the prognostic impact of cancer stem cells in HGSC
patients.

Since HGSC relapse remains the major cause of treatment failure,
targeting cancer stem cells offers an intriguing option to eradicate HGSC [40,41]. In the attempt to kill HGSC cancer stem cells,
monoclonal antibodies and small molecules directed against cancer
stem cell markers have been tested in HGSC preclinical models and
patients. An inhibitor of CD117, Imatinib, is already used as the first-
line chemotherapeutic drug for gastrointestinal stomal tumors and
chronic myelogenous leukemia. Following a study reporting antiproliferative effects of Imatinib on EOC cell lines, Imatinib was
tested in phase II clinical trials in patients with recurrent HGSC and
demonstrated no efficacy [44-46]. In addition, preclinical studies have
demonstrated the efficacy of both CD44/CD133 antibodies and
ALDH1 inhibitor in EOC cell line xenografts [41,47-50]. The growing
understanding of cancer stem cell biology unveiled alternative
therapeutic approaches to eliminate cancer stem cells [51]. These
latter set out to target features that contribute to cancer stem cell
phenotype: cancer stem cell related pathways, ATP-binding cassette
drug transporters and microenvironment (niche) [52]. Notably, our
knowledge regarding EOC cancer stem cells mainly derives from
either primary tumors or EOC cell lines [41].

**Animal Models**

**Spontaneous and carcinogen-induced models**

It is known that hens, some strains of mice, Wistar and Sprague-
Dawley rats and the primate macaques develop ovarian tumors
spontaneously [53-58]. In particular, the laying hen develops EOC that
recapitulate the 4 major histotypes of women (serous, endometrioid,
mucinous and clear cell) [53]. While these strains of mice and rats,
develop a variety of tumors including tubular adenoma, papillary cystadenoma, adenocarcinoma, mesothelioma, germ cell tumor, granulosa cell tumor and sex-cord stromal tumor [54-57]. Finally, macaques spontaneously develop granulosa cell and sex-cord stromal tumors, but not epithelial tumors similarly to humans, even if they are anatomically more similar to humans [58]. In any case, these models have a relatively late-onset of tumor development and low incidence rate that make them useless for *in vivo* studies.

Since the late sixties, ovarian tumors have been induced in experimental animals by direct application of chemical carcinogens, although no chemical carcinogen has been consistently associated to EOC etiopathogenesis [59]. Ovarian tumorigenesis has been induced using many chemical carcinogens including 9,10-dimethyl-1,2-benzanthrene, 7,12-dimethylbenz(a)anthracene, 20-methylcholanthrene, 1,3-butadiene, formic acid 2- [4-(5-nitro-2-furyl)-2-thiazolyl]hydrazide, a nitrofuran antibiotic, and N-methyl-N'-nitrosourea, a direct-acting alkylating agent [59-65]. Interestingly, the initial lesions induced by these carcinogens, when analyzed, were ovarian surface epithelial proliferations, supporting the cancer origin from the ovarian surface epithelium. However, the induced carcinomas were composed by neoplastic cells resembling either endometrium or oviduct (that corresponds to the human FTE in mice), and were organized either in glandular or papillary structures similar to human endometrioid and ovarian serous carcinomas, respectively. Therefore, it remains controversial whether latter tumors originate from the ovarian surface epithelium that has undergone metaplastic transformation and morphologically resembles the oviduct or derive directly from the oviduct.

Genetically engineered mouse models

Genetically engineered mouse models for HGSC have been difficult to generate. Recently, the direct introduction of oncogenes and/or disruption of tumor suppressor genes into the oviduct allowed the generation of mouse models that recapitulate the new pathogenetic model of HGSC including the presence of lesions similar to serous tubal intraepithelial carcinoma, the putative precursor lesion [66,67]. Kim et al. developed an EOC mice model obtained by *Dicer* double knockout [68]. Clinically similar to human HGSC, this EOC arose from the oviduct, spread to the abdominal cavity and to the ovary, and led to ascites and death of 100% of mice [68]. Phenotypically and morphologically the developed tumors resemble HGSC, although differently they harbor unaltered p53 and altered Dicer [68]. Other investigators have generated *de novo* mouse model through the inactivation of *Beca 1/2, Pten* and *Tpf3* in fallopian tube secretory cells, miming the molecular alterations commonly observed in human HGSC [66]. More recently, Sherman-Baust et al. reported a transgenic mouse model that develops HGSC from serous tubal intraepithelial carcinoma through the inactivation of both p53 and Rb pathways, similarly to HGSC [67]. Notably, both latter mouse models recapitulated human HGSC from a clinical, histopathological, and genetic point of view [66,67]. Overall, all these genetically modified mouse models provide new evidence supporting the “fallopian tube hypothesis”, i.e. the origin of HGSC from the fallopian tube (Figure 1). Therefore, they certainly offer a unique opportunity for the investigation of HGSC early events and carcinogenesis, and the exploration of new strategies of early detection, prevention and therapy. Moreover, this kind of syngeneic mouse models are optimal for the study of tumor microenvironment, angiogenesis, epithelial-stromal interaction and antitumor immune mechanisms because preserve intact immune system. The major limitations of these models are that they are labor-intensive, expensive and time-consuming and do not fully replicate the genetic and epigenetic complexity of a spontaneous HGSC. Finally, being murine these tumors own species-specific characteristics and do not necessary behave as human HGSCs (Figure 2).

**Figure 1:** Fallopian tube hypothesis on the origin of high-grade serous carcinoma (HGSC). Fallopian tube epithelium (FTE) cells of the fimbriated ends undergo initial neoplastic transformation, becoming serous tubal intraepithelial carcinoma (STIC), STIC cells possess resistance to anoikis that favors settlement and invasion of the ovarian surface. The ovarian microenvironment, rich in hormonal and inflammatory factors, drives the full neoplastic transformation to invasive HGSC (A). Alternatively, the normal FTE cells are entrapped in the ovary favored by their anatomical proximity and physiological ovulation process. Entrained FTE cells undergo progressive neoplastic transformation inside the ovary through the accumulation of molecular alterations (B).

**Figure 2:** Comparison among the most common animal models used in ovarian cancer research. A summary of the major advantages (PRO) and limitations (CONTRA) of the different animal models used in ovarian cancer research is reported.

**Human xenograft models**

Heterotransplantation of cells is possible exclusively in immunounresponsive environment that prevents transplant rejection. However, engraftment rate of human tumor in immunocompromised rodents is low, likely due to residual adaptive and/or innate immunity,
tumor intrinsic characteristics and experimental approaches employed. Nowadays, human xenografts are generated by engrafting human tumor from either primary tumor or cancer cell line into immunodeficient mice, such as: i) athymic nude mice deficient in functional T lymphocytes, ii) severe combined immunodeficient (SCID) mice lacking both B and T lymphocytes, and iii) the nonobese diabetic (NOD)/SCID/IL2Rnull mice which defect in adaptive and innate immunity due to lack of mature lymphocytes and natural killer T cells. The three main routes of implantation used for EOC xenografts are subcutaneous (ectopic), intraperitoneal and intrabursal (orthotopic) [69]. Subcutaneous implantation facilitates manipulation and serial measurements, but it does not recapitulate clinical tumor progression, since rarely malignant ascites and peritoneal carcinomatosis develop [70]. Intraperitoneal implantation consists in the injection of cells into the bursal membrane that envelops the mouse ovary and oviduct. This implantation site reproduces the physiological environment in which HGSC grows, so that tumor and ovary microenvironment can reciprocity interact and be modulated [71,72]. Therefore, intraperitoneal and intrabursal implantations best reproduce the clinical manifestations of human HGSC, recapitulating the early and late stage of the disease, respectively [69,71]. Moreover, the advent of noninvasive imaging instruments designed specifically for small animals allows monitoring of in vivo intraperitoneal tumor growth over time. Noteworthy, intraperitoneal and intrabursal sites require more time and major technical skill for successful implantation as compared to subcutaneous [70].

**Cell line xenograft models:** Despite many shortcomings, EOC cell line xenografts are the most utilized animal model in EOC research, providing a multifaceted tool to explore EOC biology and treatment [29,69-71,73]. Only selected EOC cell lines develop tumors when injected into immunocompromised mice, and the tumor engrafted usually acquires an indistinct undifferentiated morphology and displays complex genetic makeup since usually derived from patients with advanced stage [72,74,75]. The EOC cell line xenograft models commonly used are obtained by intraperitoneal injection of cell lines A2780, OVCAR-3 and SK-OV-3. Among these cell lines, OVCAR-3 genetically correlates to primary HGSC, whereas A2780 and SK-OV-3 do not harbor the typical genetic alterations of HGSC and are probably derived from endometrioid carcinoma [28]. As a consequence, such models do not accurately predict the clinical response of HGSC to novel anticancer agents, nor do they properly anticipate drug resistance and adverse reactions [4,45,76]. Nevertheless, some preclinical studies, using EOC cell line xenografts, correctly predicted anticancer drug response and effectively contributed to guide HGSC monotherapy demonstrated favorable response in phase II/III clinical trials with bevacizumab, a monoclonal antibody to human vascular endothelial growth factor, confirmed its efficacy in HGSC patients, both as a single agent and in combination with paclitaxel [80,81]. Another example of successful use of cell line xenograft model includes the preclinical efficacy of PARP inhibitors in BRCA-deficient cells in vitro and in vivo [82,83]. PARP inhibitor monotherapy demonstrated favorable response in phase II/III clinical trials in patients with HGSC harboring BRCA1 and BRCA2 germline mutations, confirming the preclinical observations [84,85]. Latter studies demonstrated the ability of EOC cell line xenografts in parallel with in vitro studies in predicting drug efficacy, anticipating drug toxicity, identifying biomarkers related to drug response [79-82,86]. Therefore, the ability to successfully translate preclinical findings to HGSC patients largely depends on the selection of appropriated EOC cell line or cell line panel, that must take into account the molecular drivers that better capture the underlying biology of primary HGSC and the likely key features of drug sensitivity.

In conclusion, the advantages of cell line xenografts include the rapidity of tumor formation, easy predictability, reproducibility, synchronization and the need of only few mice in drug response studies. On the other hand the primary limitations result from the use of EOC cell lines (above-mentioned) and immunocompromised mice, that are unable to fully represent the complex interaction between EOC and its microenvironment (immune system, stroma, inflammation, vasculature) (Figure 2).

**Patient-derived murine xenograft models:** The first patient-derived cancer xenograft (PDX) models were established in 1969, but only in the last decade its use became mainstream practice in cancer research [87]. In contrast to cell line xenografts, these emergent models feature direct transfer of tumor/ascites from patients into NOD/SCID mice, thus avoiding possible tissue culture artifacts. Mice bearing patient’s ascites tumor usually do not survive after 8 weeks and necropsy findings show peritoneal effusion and multiple tumor nodules in the peritoneal wall and mesentry, a characteristic feature of advanced stage HGSC in patients. Of note, the morphological and molecular genetic features (TP53 mutation and DNA copy number alterations) of the PDX tumors are virtually indistinguishable from the tumor obtained from the patient, suggesting that the PDX model can faithfully simulate human HGSC [88-91]. Additional key characteristics of the PDX models in contrast with cell line xenografts are: i) the tumors in PDX maintain the human intratumoral stroma and the vascularization recapitulating the physiological microenvironment; ii) PDX xenografts allow the propagation and expansion of human tumors maintaining significant morphologic and genetic concordance with the parental primary HGSC over multiple murine generations. Thanks to their characteristics, PDX models are able to reflect/predict the therapeutic responsiveness observed in clinic, showing concordance with the original patient’s treatment response [90,92], thus representing a unique opportunity toward personalized therapy for HGSC. However, some limitations affect this model. First, it requires a large amount of human fresh tumor and variable period of time to engraftment (2 to 4 months) [93]. Second, the tumor engraftment rate varies significantly in different studies depending on tumor characteristics, mice strain and implantation site [94]. Third, PDXs do not allow to study the role of immune system and, in the long run, also the tumor-stromal interactions since human stroma is eventually replaced by murine stroma [94]. Finally, the development, propagation and maintenance of PDX are time-consuming, labor intensive and expensive.

**Humanized-xenograft models:** A critical shortcoming of xenografts is the lack of lymphocyte-mediated response as a consequence of immunodeficient mice used as recipients. Therefore, xenografts preclude the pathophysiological interaction between cancer cells and immune system that plays a key role in EOC initiation, progression and therapy response [95-98]. To overcome this issue, immunocompromised mice may be engrafted with peripheral CD34 positive blood cells, isolated from human umbilical cord or fetal liver, that are enriched for hematopoietic stem cells [99]. This strategy restores almost completely the human immune system in mice, renamed hematolymphoid humanized [100], and allows to study the immunological response to HGSC as well as the biology of HGSC preserving cancer immunoeediting [101].
Another shortcoming of EOC xenographs is its inability to efficiently mimic epithelial/stromal interactions during tumor initiation due to the absence of normal human ovary. Dates back to 2001 the first study in which normal human ovarian tissue has been implanted into mice to test the morphological and functional preservation following cryostorage used to restore fertility in prepubertal children and women undergone to gonadotoxic treatment for cancer [102-104]. This approach can also have interesting applications in ovarian cancer research since allows to recreate in mice the human ovarian microenvironment where HGSC develops.

Conclusions

HGSC still represents an insurmountable challenge in gynecological oncology mainly because of late diagnosis and chemoresistance onset. Experimental cell and animal models are of pivotal importance to study human disease since allow: i) to understand the natural history and pathogenetic steps that lead to a fully developed disease; ii) to identify potential therapeutic targets; iii) to enable preclinical testing of novel therapies, alone or in combination with standard therapies.

To this day, we lack good inbred laboratory animals that develop HGSC. This is largely due to our limited understanding of the initiating factors that trigger HGSC. Moreover, anatomical, physiological and pathophysiological differences between animals and humans female reproductive system, including short lifespan, seasonal mouse reproduction, estrous cycle instead of menstrual cycle, and lack of menstruation may contribute to explain the difficulty in developing a representative laboratory animal model. It has to be underlined that the emerging “fallopian tube hypothesis” has given a new momentum to ovarian cancer research (Figure 1). As a consequence, new preclinical models, in which FTE cells undergo neoplastic transformation in vitro and in vivo, have been generated [23,24,66,67]. The use and spread of these pioneering models portend new opportunities to investigate the molecular and cellular alterations associated with HGSC initiation and may allow the development of novel prevention, early detection, and screening strategies. Nevertheless, EOC cell line xenographs are still commonly used in preclinical drug development [80,81,84,85]. Recent studies highlighted that, in order to increase the possibility to translate preclinical findings to clinic, the choice of EOC cell line (as biological surrogate of HGSC) should keep into account the molecular characteristics crucial to the biology of primary HGSC and to drug sensitivity [28].

As yet, the animal model that most closely mirrors human HGSC is PDX since it offers the relevant advantage of faithfully maintaining the characteristics of patients’ parental tumor, preserving both stromal microenvironment and molecular alterations. Therefore, PDX holds the promise for personalized antineoplastic treatment and discovery of determinants of drug response [93]. In theory, PDXs allow for a rapid in vivo screening of targeted drugs and for the assessment of the chemosensitivity of patients’ cancer. In particular, since different PDXs can be generated implanting tumor fragments obtained at different time points from the same patient, they potentially allow to capture the timeline of molecular alterations, i.e. tumor evolution, through exposure to therapy and development of chemoresistance, and, furthermore, to pre-test the response to new targeted therapies. Another prospective application of PDX models is represented by the screening of new anticancer compounds in a repertoire of HGSC PDX, that allows to capture HGSC interpatient heterogeneity [105]. Therefore, PDX provides a powerful experimental platform for treatment decision-making and for guiding targeted therapy in the clinic. In addition, it has been reported that PDX preserves cancer stem cells through multiple generations and chemotheraphy, making PDX an excellent model to study the biology of cancer stem cells [38,89]. In this respect, the emergent picture of HGSC as a “cancer stem cell disease” is extremely appealing for its diagnostic and therapeutic implications, although further studies are necessary in order to better characterize stem cells in HGSC. In particular, the generation of stable models of HGSC stem cells seems an essential experimental tool needed to gain deeper knowledge and to allow the discovery of specific drug to target HGSC stem cells.

In the opening era of personalized medicine, the optimal choice of experimental cell and/or animal models remain fundamental to broadly our knowledge on HGSC. For this purpose, experimental platforms must keep abreast of the ever-increasing molecular, pharmacological and clinical information from human disease in order to enhance their translational potential.

References


