

Spotlight on olaparib in the treatment of BRCA-mutated ovarian cancer: design, development and place in therapy

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Abstract: Epithelial ovarian cancer is the sixth most common cancer among women worldwide and the first cause of death among gynecological malignancies. Most of the patients present recurrent disease and unfortunately cannot be cured. The unsatisfactory results obtained with salvage chemotherapy have elicited investigators to search for novel biological agents capable of achieving a better control of the disease. In the setting of homologous recombination deficiency, the DNA errors that occur cannot be accurately repaired, and the treatment with poly(ADP-ribose) polymerase (PARP) inhibition results in definitive cell death in a process called synthetic lethality. As a result of two positive clinical trials, Olaparib was approved in 2014 by U.S. Food and Drug Administration and European Medicines Agency as the first-in-class PARP inhibitor. Olaparib is effective and well tolerated in homologous recombination deficient patients. Several studies with Olaparib have been conducted in the recurrent setting either as maintenance in platinum-responsive patients or as a single agent. Ongoing trials are focused on the use of olaparib as maintenance in the first-line ovarian cancer setting alone or in combination with antiangiogenic agents. Future perspectives will probably investigate the association of olaparib with novel agents as check-point inhibitors and PI3K-AKT inhibitors. The PARP inhibitor era is just at the beginning.

Keywords: olaparib, ovarian cancer, PARP inhibitors, homologous recombination deficiency, BRCA mutation

Introduction

Epithelial ovarian cancer (EOC) is the sixth most common cancer among women worldwide and the first cause of death among gynecological malignancies.¹

The standard management of early stage disease consists of comprehensive staging surgery, followed by adjuvant carboplatin (CBDCA) or CBDCA–paclitaxel (PTX) chemotherapy in high-risk cases.^{2,3} Cytoreductive surgery followed by PTX–CBDCA chemotherapy is the backbone of treatment for advanced EOC, whereas neoadjuvant chemotherapy followed by interval debulking surgery is indicated for women with poor clinical conditions or with a great amount of disease suggesting a low likelihood of obtaining an optimal cytoreduction (residual disease [RD] 0 or <1 cm).⁴

For advanced disease (FIGO stage IIIB–IV), bevacizumab, a humanized anti-VEGF monoclonal antibody, has been licensed by the European Medicines Agency (EMA) in combination with carboplatin and paclitaxel and in maintenance at the dose of 15 mg/kg for 15 months on the basis of two randomized clinical trials (GOG-218 and ICON-7) reporting that the combination of bevacizumab with chemotherapy translates into an increase in progression-free survival (PFS) without any differences in overall survival (OS).^{5,6}

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Approximately 13%–31% of patients with early EOC and 75%–80% of those with advanced disease relapse after a median interval of 11–29 months and 18–24 months, respectively.⁷

Patients with recurrent EOC receive second-line chemotherapy, mainly dependent on platinum-free interval, persistent toxicities, and the type of treatment previously received. Sequential single agents, such as weekly PTX, pegylated liposomal doxorubicin (PLD), and gemcitabine (GEM), are suggested for platinum-resistant patients;⁸ on the contrary, CBDCA doublets (ie, CBDCA in combination with PTX, GEM, or PLD), are used in patients with platinum-sensitive disease^{9–11} and the non-platinum combination of PLD+trabectedin is a therapeutic option for those who have partially platinum-sensitive disease as well as for those who do not fit for platinum rechallenge.¹²

Moreover, in this setting, two randomized trials reported that the combination of carboplatin–gemcitabine–bevacizumab or carboplatin–paclitaxel–bevacizumab administered until progression of disease, significantly increases PFS with an hazard ratio of 0.48¹³ with a nonsignificant trend in OS increase (hazard ratio 0.82).¹⁴

However, the unsatisfactory results obtained with salvage chemotherapy have elicited investigators to detect novel biological agents capable of achieving a better control of the disease.¹⁵ In the last two decades several changes in all fields of ovarian cancer management have occurred, from the diagnosis, to the treatment, to the translational research. Moreover, new drugs have been introduced in the treatment algorithm with the intent to increase the quantity and quality of life of ovarian cancer patients.

Homologous recombination repair defects: role of BRCA genes and PARPs

DNA is continuously subjected to injuries by environmental and endogenous exposures that cause a variety of DNA lesions, including double-strand breaks (DSBs) and single-strand breaks (SSBs).¹⁶ DNA repair systems are critical to maintain genomic integrity by allowing cells to replicate and survive.¹⁷ Homologous recombination repair (HRR) is the most important instrument of reparation of DSBs. The BRCA1/2 genes, together with several other genes, code proteins that are necessary for this process. When either BRCA1 or BRCA2 is defective, homologous recombination is dysfunctional and the reparation of DSBs is performed through alternative repair mechanisms such as nonhomologous end-joining (NHEJ) and single-strand repair.^{18,19} SSBs repair involves a variety of mechanisms such as base excision

repair (BER) and nucleotide excision repair, all of which are supported by poly(ADP-ribose) polymerases (PARPs).²⁰

PARPs constitute a family of 18 proteins.²¹ PARP1 and PARP2 are enzymes involved in SSBs and BER, which are activated by DNA damage and facilitate DNA repair.²² PARP1 becomes activated when an SSB occurs and, after binding to the damaged area, increases its catalytic activity and recruits various other proteins to the site of the DNA damage, initiating a repair complex. If a cell is not able to repair SSB before initiating replication, a single break is transformed into a double-strand during replication process.²³

Several studies proposed the model of synthetic lethality, a process by which cancer cells are contemporarily targeted by the inactivation of two genes when the deficiency of either gene alone is nonlethal.^{23,24} This model can be applied to homologous recombination deficient (HRD) cells; in this case, in fact, PARP inhibitors inhibit the repair of DNA SSBs, thus transforming them into DNA DSBs. When homologous recombination is not functional (HRD), as it is in patients with BRCA mutations, the DNA DSBs cannot be repaired and the PARP inhibition ultimately results in cell death, as shown in Figure 1.²⁵

This mechanism is an important therapeutic target, not just for PARP inhibitors, but for many chemotherapeutic agents and radiotherapy acting by inducing DNA damages. Platinum analogs, in fact, induce intrastrand and interstrand cross-links the reparation of which depends on nucleotide excision repair and by DSB formation.¹⁶ The reported elevated platinum sensitivity of BRCA-mutated EOC to platinum is believed to be related to the HRR defects. BRCA1 and BRCA2 are not the only genes involved in the HRD repair mechanism: other

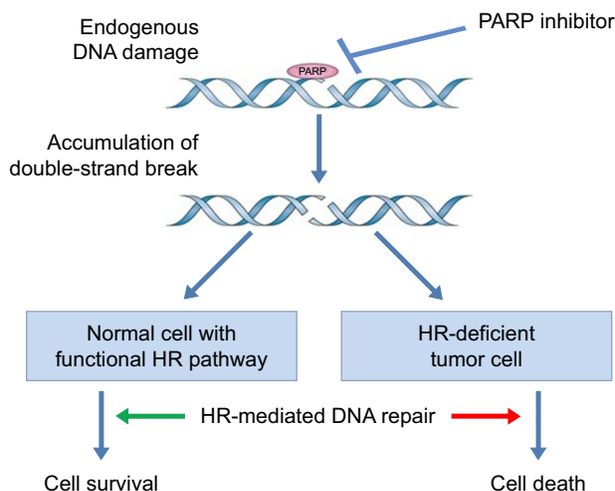


Figure 1 Mechanism of synthetic lethality.

Abbreviations: HR, homologous recombination; PARP, poly(ADP-ribose) polymerases.

members of the Fanconi anemia family, such as RAD51C, RAD51D, and BRIP1^{26–28} as well as ATM, CHEK1, CHEK2, and CDK12 also confer sensitivity to DNA damage and DNA repair inhibition.^{24,29–31} These mutations are responsible for what we actually call the BRCAness phenotype,³² a clinical situation in which, even in the absence of identified BRCA mutations, the disease present repeated platinum sensitivity, long natural history, and the potentiality to respond to PARP inhibitors treatment.

Ovarian cancer patients who present a germline or somatic BRCA1 and BRCA2 mutations have a better prognosis compared with BRCA wild-type patients, possibly because of the elevated response rate (RR) to platinum agents, but also to pegylated liposomal doxorubicin and trabectedin. The mutations appear particularly frequent (about 22%–26%) among high-grade serous ovarian cancer (HGSOC),^{33–37} which are also impaired by the presence of mutations of the tumor suppressor gene p53.^{38,39} Moreover, molecular analysis of HGSOC by The Cancer Genome Atlas revealed that around 50% of them present HRD.³⁷ After sequencing 316 HGSOCs germline, BRCA1 and BRCA2 mutations were reported in 9% and 8% of patients, respectively, somatic additional mutations in BRCA1/2 in 3%, EMSY in 8%, PTEN in 7%, RAD51C in 3%, ATM/ATR in 2%, and Fanconi anemia genes in 5% of patients.³⁷ The rates of somatic mutations reported in the literature are variable, and currently, the true prevalence of somatic mutations remains unknown; however it has been estimated between 5% and 8% of cases. This implies that for every five ovarian cancer patients with a germline BRCA mutation, there will be one patient with a somatic mutation.⁴⁰

Sporadic EOCs with HR deficiency not linked to BRCA1-2 mutations have the same biological characteristics and clinical behavior as EOC with either germline or somatic BRCA mutations (“BRCAness” phenotype),^{41,42} and, as such, the potentiality to respond to the same drugs.

Olaparib – the first PARP inhibitor approved

The first PARP enzyme was discovered over 50 years ago and the first drug capable of inhibiting PARP, 20 years later.^{43,44}

In 2005, two outstanding studies, performed by two independent research groups, reported that BRCA1/2-deficient cells were 100–1,000-fold more sensitive to PARP inhibitors than wild-type cell,^{45,46} thus suggesting a particular sensitivity of BRCA-deficient cell lines to PARP inhibitors through a mechanism of “synthetic lethality”. Clinical trials were initiated to explore the clinical activity of PARP inhibitors in HRD-defective tumors.

Olaparib reported 47% RR in a Phase I trial in patients with breast, prostate, and ovarian cancers, harboring BRCA1 or BRCA2 mutations;⁴⁷ moreover, a correlation between RR and platinum sensitivity was reported (RR 69% vs 45% vs 23% in platinum-sensitive, platinum-resistant, and platinum-refractory patients, respectively).⁴⁸

Olaparib was approved in 2014 by the U.S. Food and Drug Administration (FDA) and EMA as the first-in-class PARP inhibitor on the basis of two prospective clinical trials (Study 19 and Study 42).^{49,50}

On December 2014, the FDA approved olaparib capsules (Lynparza; AstraZeneca, Wilmington, DE, USA) for the treatment of patients with germline BRCA-mutated (gBRCAm) ovarian cancer who had received at least three previous chemotherapy lines.⁵¹

On October 2014, the EMA approved Lynparza as maintenance treatment in platinum-sensitive, BRCA-mutated (germline and/or somatic), high-grade serous EOC who were responding to the last platinum-based chemotherapy.⁵²

Study 19 was an international, randomized, Phase II, double-blind, placebo-controlled trial in patients with relapsed platinum-sensitive, high-grade serous, ovarian, fallopian tube, or primary peritoneal cancers.⁴⁹ In the study, 265 patients in complete or partial response to the last platinum-based chemotherapy were randomized to receive maintenance with olaparib (n=136) or placebo (n=129) after completion of at least four cycles of chemotherapy. The primary endpoint was PFS assessed by the investigators per Response Evaluation Criteria in Solid Tumors version 1.0 criteria. The trial reported a significant 3.6 months increase in median PFS in patients treated with olaparib with respect to placebo (median PFS 8.4 vs 4.8 months for patients treated with olaparib and placebo, respectively; HR 0.35; $p<0.001$).⁴⁹ In the preplanned subgroup analysis, the 136 BRCA-mutated patients derived the greatest clinical benefit from olaparib with a significant improvement in PFS of 6.9 months (median PFS 11.2 vs 4.3 months for olaparib and placebo, respectively; HR 0.18; $p<0.0001$). The PFS improvement was confirmed by an independent radiological review.⁵³ Study 19 pressed the fast-track EMA approval of olaparib as maintenance in Europe, but the approval was conditioned by the result of the ongoing randomized Phase III study, which were presented at International SGO Congress in March 2017. The confirmatory randomized Phase III SOLO-2 trial investigated olaparib tablets as maintenance in platinum-sensitive, germline BRCA-mutated ovarian cancer patients, who responded to the last platinum-based treatment. Two hundred ninety five eligible patients were

randomly assigned to receive olaparib (n=196) or placebo (n=99) at the completion of at least four cycles of platinum-based chemotherapy. PFS was significantly longer in the olaparib arm than in the placebo arm (19.1 vs 5.5 months, respectively; HR 0.30; $p < 0.0001$) with an acceptable toxicity profile and without any detrimental effect on patients' quality of life.⁵⁴

Study 42 was a single-arm, Phase II, prospective study on olaparib 400 mg BID in the treatment of patients with gBRCA1/2-mutated cancers; 193 out of 298 patients were heavily pretreated with mean number of 4.3 prior chemotherapy regimens.⁵⁰ The overall RR was 26.2% in the general population and 31.1% in ovarian cancer setting. Prolonged (≥ 8 weeks) stable disease was observed in 42% of patients overall and in 40% of ovarian cancer patients.⁵⁰

Recently, a 34% RR with a median response duration of 7.9 months, in a subgroup analysis of EOC patients who had received three or more previous chemotherapy lines, was reported.⁴⁹ Study 42 sustained FDA approval of olaparib as a single agent in BRCA-mutated ovarian cancer patients who had received at least three previous chemotherapy lines.

Phase II/III studies of olaparib in ovarian cancer are summarized in Table 1.^{50,53,55–58}

Safety profile

Olaparib is generally well tolerated; adverse reactions are typically of mild or moderate severity (Common Terminology Criteria for Adverse Events grade 1 or 2) and, in most cases, are short term in nature, self-limiting, and do not require treatment discontinuation or dose reductions.

In the pivotal Phase II trial (Study 19), the most commonly reported adverse events (AEs) were nausea, fatigue, vomiting, and anemia.^{49,53} The tolerability profile of olaparib in patients with BRCAm cancer did not differ from that of the overall population.⁴⁹ Subsequent analyses reported that typically AEs occurred within the first 4–8 weeks of treatment and were mainly grade 1 or 2, generally transient, and managed with supportive care without dose reductions.^{58,59} In the overall population, serious AEs (SAEs) were reported in 18% of patients receiving olaparib vs 9% of those who received placebo while SAEs causally related to olaparib and placebo in the investigator's judgment were 5.9% and 0.8%, respectively.⁵³

Treatment interruptions and dose reductions in the olaparib arm were reported in 28% and 23% of patients, respectively.⁴⁹ After a median follow-up of 5.9 years, no new safety findings appeared.⁵⁹ The SAEs leading to permanent discontinuation of treatment occurred in 6% and 2% of patients receiving olaparib and placebo, respectively.^{49,60,61}

In the SOLO-2 study, the most commonly reported grade ≥ 3 AEs were anemia (19% in the olaparib group vs 2% in the placebo group), fatigue (4% vs 2%), and neutropenia (5% vs 4%). SAEs were experienced by 18% of patients in the olaparib group and 8% of patients in the placebo group, respectively. One patient in the olaparib group had an acute myeloid leukemia with death as outcome.⁵⁴

In Study 42, grade 3 AEs were reported in 54.4% of patients (anemia and fatigue were the most common); in 30.9% of cases, they were considered drug related. SAEs were seen in 30.1% of patients; in about 10.0% they were considered causally related to olaparib. Nine patients died as a result of AEs; in 3.7% of patients the AEs led to treatment discontinuation and in 40.3% caused drug dose modification.⁵⁰ The most frequently reported AEs in the trials involving olaparib are reported in Table 2.

Olaparib as maintenance or in combination with chemotherapy?

A synergism between PARP inhibitors and DNA-damaging agents such as cisplatin, carboplatin, or cyclophosphamide has been described.^{62,63} Thus, the combination of olaparib with other chemotherapy agents appears promising. In a randomized, open-label, Phase II study, patients with platinum-sensitive recurrent HGSOc were treated with carboplatin and paclitaxel in combination and maintenance with olaparib or chemotherapy alone. PFS was significantly increased in the experimental arm (12.2 months vs 9.6 months; $p = 0.0012$), and the improvement was larger in patients with BRCA mutations. The increased toxicity reported in the combination arm and the shape of the curves which diverged only in the maintenance phase seem to suggest no additional benefit of the combination vs the maintenance only.⁵⁸

Olaparib: future perspectives

In an attempt to move toward the first-line treatment in ovarian cancer patients, the recently concluded SOLO-1 trial was developed. SOLO-1 trial is a randomized Phase III study with olaparib as maintenance at the completion of first-line platinum–paclitaxel chemotherapy in FIGO stage III–IV, BRCA-mutated ovarian cancer patients. The results are awaited for Q₂ 2018.

The combination of PARP inhibitors with other molecular-targeted agents has been explored in clinical trials.

A synergism between PARP inhibitors and antiangiogenic agents has been reported,⁶⁴ possibly due to the down-regulation of DNA repair mechanism by antiangiogenic agents. A significant increase in PFS (17.7 vs 9.0 months;

Table I Phase II/III studies of olaparib in ovarian cancer

| Study | Patient population and BRCA status | Treatment arms | Total accrual | Primary endpoint | ORR | PFS |
|-------------------------------------|---|--|---------------------------------|------------------|---|---|
| Audeh et al, ⁵⁵ 2010 | Recurrent epithelial ovarian, primary peritoneal, or fallopian tube carcinoma BRCA1/2 positive | Cohort 1: olaparib 400 mg BID Cohort 2: olaparib 100 mg BID | 57 | ORR | Cohort 1: 33% Cohort 2: 13% | Cohort 1: 5.8 months Cohort 2: 1.9 months |
| Kaye et al, ⁵⁶ 2012 | Platinum-resistant, recurrent, epithelial ovarian, primary peritoneal, or fallopian tube carcinoma BRCA1/2 positive | Arm 1: olaparib 200 mg BID Arm 2: olaparib 400 mg BID Arm 3: PLD 50 mg/m ² | 97 | PFS | Arm 1: 25% Arm 2: 31% Arm 3: 18% | Arm 1: 6.5 months Arm 2: 8.8 months Arm 3: 7.1 months |
| Gelmon et al, ⁵⁷ 2011 | Advanced metastatic or recurrent ovarian, primary peritoneal, or fallopian tube cancer (high-grade serous and/or undifferentiated) or breast cancer BRCA1/2 positive AND BRCA1/2 negative | Olaparib 400 mg BID | 91 (65 with gynecologic cancer) | ORR | BRCA1/2 positive: 41% BRCA1/2 negative: 24% BRCA1/2 positive+platinum sensitive: 60% BRCA1/2 negative+platinum sensitive: 50% BRCA1/2 positive+platinum resistant: 33% BRCA1/2 negative+platinum resistant: 4% | BRCA1/2 positive: 221 days BRCA1/2 negative: 192 days |
| Ledermann et al, ⁵³ 2014 | Platinum-sensitive, recurrent, high-grade serous epithelial ovarian, primary peritoneal, or fallopian tube carcinoma BRCA1/2 positive AND BRCA1/2 negative | (Maintenance therapy following platinum-based chemotherapy) Arm 1: olaparib 400 mg BID Arm 2: placebo | 265 | PFS | | Arm 1: 8.4 months Arm 2: 4.8 months Olaparib+BRCA1/2 positive: 11.2 months Olaparib+BRCA1/2 negative: 5.6 months Placebo+BRCA1/2 positive: 4.3 months Placebo+BRCA1/2 negative: 5.5 months |
| Oza et al, ⁵⁸ 2015 | Platinum-sensitive, recurrent, serous ovarian cancer BRCA1/2 positive AND BRCA1/2 negative | Arm 1: Olaparib 200 mg BID+paclitaxel 175 mg/m ² +carboplatin AUC 4×6 cycles followed by olaparib 400 mg BID maintenance Arm 2: Paclitaxel 175 mg/m ² +carboplatin AUC 4×6 cycles | 162 | PFS | Arm 1: 64% Arm 2: 58% | Arm 1: 12.2 months Arm 2: 9.6 months |
| Kaufman et al, ⁵⁰ 2015 | Platinum-resistant, recurrent, ovarian, primary peritoneal, or fallopian tube cancer BRCA1/2 positive | Olaparib 400 mg BID | 193 | ORR | 31% | 225 days |

Abbreviations: AUC, area under the curve; ORR, objective response rate; PFS, progression-free survival; PLD, pegylated liposomal doxorubicin.

$p=0.005$) and RR (79.6% vs 47.8%; $p=0.002$) has been reported with the combination of olaparib plus cediranib vs olaparib alone in recurrent, platinum-sensitive, high-grade ovarian tumors.⁶⁵

A Phase I study was performed combining olaparib and bevacizumab; patients with advanced cancers received increasing doses of olaparib (100, 200, and 400 mg BID, capsule formulation) in combination with bevacizumab 10 mg/kg IV every 2 weeks.⁶⁶ A total of 12 patients were enrolled and

the authors concluded that the combination of olaparib 400 mg BID and bevacizumab 10 mg/kg was well tolerated and it represented the maximum tolerated dose for future trials.⁶⁶

PAOLA-1 is an ongoing ENGOT/GCIG Phase III trial evaluating olaparib (tablet formulation) vs placebo in combination with bevacizumab as maintenance treatment in patients with stage IIIB–IV high-grade serous or endometrioid ovarian cancers treated with standard first-line platinum-based chemotherapy plus bevacizumab.

Table 2 Patients (%) in olaparib arm: any-grade AEs reported in >15% or grade ≥ 3 AEs reported in >5% of patients overall

| Adverse event | Study 42 ⁵⁰ | | Study 19 ⁴⁹ | | SOLO-2 ⁵³ | |
|--------------------|------------------------|------------|------------------------|------------|----------------------|------------|
| | Any grade | G ≥ 3 | Any grade | G ≥ 3 | Any grade | G ≥ 3 |
| Fatigue | 60.1 | 6.2 | 48.5 | 6.6 | 66 | – |
| Nausea | 61.7 | 0.5 | 68.4 | – | 76 | – |
| Vomiting | 38.9 | 2.6 | 31.6 | – | 38 | – |
| Anemia | 32.1 | 18.7 | 19.8 | 5.1 | 43 | 19 |
| Diarrhea | 29.0 | 1.6 | 22.8 | – | 33 | – |
| Abdominal pain | 30.1 | 7.3 | 17.6 | – | 25 | – |
| Decreased appetite | 18.7 | 0.5 | 18.4 | – | 22 | – |
| Dyspepsia | 19.7 | 0 | 16.2 | – | – | – |
| Headache | 16.6 | 0 | 18.4 | – | 26 | – |
| Dysgeusia | 20.2 | 0 | – | – | 27 | – |
| Constipation | – | – | – | – | 21 | – |
| Cough | – | – | – | – | 17 | – |
| Arthralgia | – | – | – | – | 15 | – |
| Neutropenia | – | – | – | – | 19 | 5 |

Abbreviation: AEs, adverse events.

Another interesting combination with a strong preclinical rationale is the association between PARP inhibitors and PI3K inhibitors. Juvekar et al reported an in vivo synergism between PI3K inhibitor BKM120 and olaparib in *BRCA1*-mutated breast tumors, thus suggesting an important role of PI3K α in the DNA damage response.⁶⁷

In the Phase I study of olaparib and BKM120, patients with either breast or ovarian cancer were enrolled; clinical benefit was observed in both gBRCAm and gBRCAwt patients but the combination required attenuation of the BKM120 dose. Randomized Phase II studies are needed to further define the efficacy of PI3K/PARP inhibitor combinations as compared with a PARP inhibitor alone.⁶⁸

Michalarea et al recently presented data on the combination of olaparib with AZ5363, an AKT inhibitor.⁶⁹ Common (>15%) G1-2 toxicities were nausea, vomiting, fatigue, diarrhea, and anemia. Based on tolerability, recommended Phase II dose for the combination was established at 640 mg BID 2/7 AZD+300 mg BID olaparib.⁶⁹

Recently, the causes of resistance to PARP inhibitors have been investigated: unfortunately, in most of the cases, the resistance is due to unknown mechanisms, in 15% to a BRCA mutation reversion, and in 10% to TP53BP1 gene mutations, which have opposing activity to BRCA1 in preventing DNA resection and promoting NHEJ.⁷⁰

The future goal of the research will be the identification of biomarkers that can predict the response to PARP inhibitors.

In 2014, Lee et al proposed that tumor FOXO3a expression may represent a predictor of response to the combination

therapy of carboplatin plus olaparib in mBRCA patients with ovarian or breast cancer.⁷¹

Other preclinical data report that the abundance of PARP may reflect cellular DNA repair deficiencies, thus constituting a universal predictive biomarker for the response to PARP inhibitors.⁷² PARP binding protein overexpression is reported in pancreatic cancers, where it induces genetic instability and PARP hyperactivation, thus suggesting that it might predict both HRD and sensitivity to PARP inhibition.⁷³

The characterization and real application of such biomarkers in clinical practice need to be verified in prospective trials.

Other PARP inhibitors

Other PARP inhibitors have been developed. Veliparib, a PARP1 and PARP2 inhibitor, reported 20% and 35% RR in platinum-resistant and platinum-sensitive, BRCA-mutated ovarian cancer patients, respectively.⁶² A Phase III study on veliparib as maintenance therapy in first-line treatment of ovarian cancer is ongoing (NCT02470585).

The ARIEL2 Phase II study tested rucaparib as a single agent in the treatment of BRCA-mutated, BRCA-wild type, and BRCA-like patients selected according to the genome-wide loss of heterozygosity (LOH) next generation sequencing test. RR was 69% vs 39% vs 11% in BRCA-mutated, BRCA-like, and BRCA-wild type ovarian cancer patients.^{74,75}

Based on this study, rucaparib received FDA approval for the treatment of BRCA-mutated ovarian cancer patients (either germline or somatic) who had previously received two or more chemotherapy lines.

ARIEL3 (NCT 01968213) is a Phase III trial designed to evaluate the efficacy of Rucaparib vs placebo as maintenance treatment after platinum-based chemotherapy in women with relapsed, platinum-sensitive, high-grade serous or endometrioid ovarian cancer. Responses to treatment will be analyzed based on homologous recombination (HR) status of tumor samples evaluated by LOH test. The study has been recently completed and the results were presented at ESMO 2017. The most robust clinical outcomes were observed among patients with a germline or somatic BRCA mutation. The median PFS for patients treated with rucaparib was 16.6 months (95% CI 13.4–22.9) vs 5.4 months (95% CI 3.4–6.7) in the placebo arm. The median PFS for the HRD patients treated with rucaparib was 13.6 months (95% CI 10.9–16.2) vs 5.4 months (95% CI 5.1–5.6) among those who received placebo. The median PFS of the intention to treat population was 10.8 months (95% CI 8.3–11.4) vs 5.4 months (95% CI 5.3–5.5) for rucaparib- and placebo-treated patients, respectively. The most common grade ≥ 3 AEs in

the rucaparib group were anemia (18.8%) and liver enzyme increase (10.5%).⁷⁶

Niraparib maintenance treatment showed a significant improvement in PFS with respect to placebo in recurrent, platinum-sensitive, BRCA-mutated, ovarian cancer patients responsive to the last platinum-based treatment (21.0 vs 5.5 months; $p < 0.001$).^{73,74} Of note, PFS was significantly increased also in patients without germline BRCA mutation (9.3 vs 3.9 months; HR 0.45), thus suggesting that platinum sensitivity, and not BRCA mutation, represents the most performing predictive biomarker for HRD.⁶⁷ Niraparib received FDA approval (March 2017) for maintenance treatment of platinum-sensitive, recurrent ovarian cancer regardless of BRCA mutations.^{77–79}

Conclusion

Recurrent ovarian cancer represents a challenging clinical situation – cure is almost impossible and prolongation of survival at the prize of an acceptable toxicity and no detrimental effect on quality of life of the patients are the goals of treatments. In this scenario, the availability of new drugs will further contribute to the process of chronicization of the disease. PARP inhibitors represent very interesting drugs interfering with mechanism of DNA repair in patients harboring HRD deficiency (a process called synthetic lethality). The drugs reported outstanding clinical activity with manageable toxicity profile in all the setting of disease where they have been employed. Olaparib is the first-in-class PARP inhibitor approved in Europe as maintenance in BRCA-mutated, platinum-responsive patients and in the USA as a single agent in BRCA-mutated patients who have received at least three previous chemotherapy lines. The future of olaparib is to upgrade into the management of newly diagnosed disease either alone (SOLO-1 trial ongoing) or in combination with bevacizumab (PAOLA-1 trial ongoing) and in non-gBRCA patients in the light of the continuous discovery of several genes involved in BRCA-independent HRD impairment. Other interesting combinations are with immunotherapy and PI3K-AKT inhibitors. The exciting sensation is that these new compounds are here to stay and to change the natural history of disease at least in a subset of patients – actually 15% of patients initially enrolled in Study 19 are still receiving olaparib after >5 years. The PARP inhibitor era is just at the beginning.

Disclosure

Francesco Raspagliesi was the principal investigator of the SOLO-1 trial. Domenica Lorusso was the principal investigator of NOVA and ARIEL3 trials. The other authors report no conflicts of interest in this work.

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