A decade of clinical development of PARP inhibitors in perspective

J. Mateo1,2, C. J. Lord3,4, V. Serra1, A. Tutt4,5, J. Balmaña1,2, M. Castroviejo-Bermejo1, C. Cruz1,2, A. Oaknin1,2, S. B. Kaye6,7 & J. S. de Bono6,7*

1Vall d’Hebron Institute of Oncology (VHIO), Barcelona; 2Vall d’Hebron University Hospital, Barcelona, Spain; 3The CRUK Gene Function Laboratory; 4The Breast Cancer Now Toby Robins Research Centre, The Institute of Cancer Research, London; 5The Breast Cancer Now Research Unit, Guy’s Cancer Centre, Kings College, London; 6The Royal Marsden NHS Foundation Trust, London; 7The Institute of Cancer Research, London, UK

*Correspondence to: Prof. Johann S. de Bono, Experimental Cancer Medicine, Division of Clinical Studies, The Institute of Cancer Research, Drug Development Unit, The Royal Marsden NHS Foundation Trust, Downs Rd, Sutton, Surrey SM2 5PT, UK. Tel: +44-2087224028; Fax: +44-2086427979; E-mail: johann.de-bono@icr.ac.uk

Genomic instability is a hallmark of cancer, and often is the result of altered DNA repair capacities in tumour cells. DNA damage repair defects are common in different cancer types; these alterations can also induce tumour-specific vulnerabilities that can be exploited therapeutically. In 2009, a first-in-man clinical trial of the poly(ADP-ribose) polymerase (PARP) inhibitor olaparib clinically validated the synthetic lethal interaction between inhibition of PARP1, a key sensor of DNA damage, and BRCA1/BRCA2 deficiency. In this review, we summarize a decade of PARP inhibitor clinical development, a work that has resulted in the registration of several PARP inhibitors in breast (olaparib and talazoparib) and ovarian cancer (olaparib, niraparib and rucaparib, either alone or following platinum chemotherapy as maintenance therapy). Over the past 10 years, our knowledge on the mechanism of action of PARP inhibitor as well as how tumours become resistant has been extended, and we summarise this work here. We also discuss opportunities for expanding the precision medicine approach with PARP inhibitors, identifying a wider population who could benefit from this drug class. This includes developing and validating better predictive biomarkers for patient stratification, mainly based on homologous recombination defects beyond BRCA1/BRCA2 mutations, identifying DNA repair deficient tumours in other cancer types such as prostate or pancreatic cancer, or by designing combination therapies with PARP inhibitors.

Key words: PARP inhibitors, DNA repair, clinical trials

A deeper understanding of the molecular make up of tumours has brought opportunities to develop more precise treatments, targeting tumour vulnerabilities. Inhibitors of poly(ADP-ribose) polymerase (PARP) are examples of the path towards precision medicine. In this manuscript, we review the clinical development of PARP inhibitors (PARPi) across tumour types, leading to approvals in breast and ovarian cancer, and analyse opportunities ahead to optimize the use of this drug class.

Fast-track for translational research: the launch of PARPi clinical development

In 2005, two seminal studies demonstrated that tumour cells lacking BRCA1 or BRCA2, key tumour suppressor proteins involved in double-strand DNA break (DSB) repair by homologous recombination (HR), are selectively sensitive to small molecule inhibitors of the PARP family of DNA repair enzymes [1, 2]. The model proposed was based on the concept of synthetic lethality: loss of either of two genes is not lethal per se, but concomitant inactivation leads to cell death. PARP1, the major target of PARPi, is primarily involved in the repair of single-strand DNA breaks (SSBs); PARP1 inhibition alone is not lethal as the DNA lesions caused by these drugs can be repaired by other DNA repair pathways, specifically HR. In contrast, in the absence of BRCA1/2 and therefore defective HR, the DNA lesions caused by PARPi are not repaired and cause cytotoxicity.

In these studies, decreasing PARP1 levels by RNA interference also resulted in a significant reduction of cell survival selectively in BRCA1- and BRCA2-deficient cells [1]. Cancer cell lines lacking BRCA1/2 were also sensitive to inhibitors of PARP1, whereas cells with only heterozygous loss of BRCA1/2 genes or those
without BRCA1/2 defects were not. In a back-to-back publication, depletion of BRCA2 using short-interfering RNA (siRNA) sensitized cancer cell lines to PARP inhibition [2]. Later studies demonstrated how loss of other tumour suppressor DNA repair proteins, many of which are involved in HR, also caused sensitization to PARPi [3–5].

PARPi were originally developed for cancer treatment as radio- and chemo-sensitizing drugs, but the aforementioned preclinical observations supported the development of PARPi as single agents for the treatment of BRCA1/2-defective cancers. In 2005, most of the data on BRCA1/2 related to the role of these genes as risk susceptibility factors for familial breast and ovarian cancers. Given this, germline BRCA1/2 (gBRCA1/2) mutation carriers with cancer were the initial target population to test the PARPi-BRCA synthetic lethal hypothesis in the clinic. A first-in-human clinical trial of KuDOS Pharmaceuticals/AstraZeneca, later named AZD-2281/olaparib was conducted to establish a recommended dose and to generate preliminary data in a biomarker-defined population [6, 7]. In this proof-of-concept trial, pharmacokinetics and pharmacodynamics (in peripheral mononuclear blood cells (PBMC), hair follicles, and tumour samples) studies were used to optimize the dose-escalation and expansion phases. Expansion cohorts only included patients with gBRCA1/2 mutations. Doses of 60 mg or more twice daily of olaparib resulted in >90% PARPi inhibition in PBMCs, suggesting biological activity at low doses. Dose-limiting toxicities of fatigue, somnolence and thrombocytopenia led to establishing 400 mg of olaparib capsules twice daily as the maximum tolerated dose. A modified tablet formulation with enhanced bioavailability was later developed; the current olaparib approved dose is 300 mg tablet twice a day [8]. Importantly, gBRCA1/2 mutation carriers did not experience enhanced toxicities, supporting the hypothesis of a cancer-specific vulnerability. Overall, 21 gBRCA1/2 mutation carriers were enrolled and evaluated for response, with radiological responses in eight patients with ovarian cancer and one with breast cancer, and a prostate cancer patient with a sustained PSA response.

This rapid translation of preclinical studies into promising clinical data triggered the development of several PARPi in different tumour types.

**Mechanisms of action of PARPi: beyond synthetic lethality**

PARP1 is a DNA damage sensor and signal transducer that binds to DNA breaks and then synthesizes poly(ADP-ribose) (PAR) chains on target proteins (PARylation) in the vicinity of the DNA break and itself (autOPARylation). These PAR chains lead to the recruitment of additional DNA repair effectors that complete the DNA repair process. In its non-DNA bound state, PARP1 has minimal catalytic activity due to an auto-inhibitory helical domain (HD) interaction with its catalytic domain [9]. When PARP1 binds DNA, via zinc finger domains, a conformational change in the PARP1 protein relieves the autoinhibitory interaction between the HD and the catalytic domain, allowing nicotinamide adenine dinucleotide (β-NAD⁺), the PARP1 co-factor, to bind the active site of the enzyme. PARP1 then uses the hydrolysis of β-NAD⁺ to catalyse the transfer of ADP-ribose moieties on to target proteins. This PARylation of proteins in the vicinity of the DNA breaks then likely mediates DNA repair by modifying chromatin structure (e.g. via histone-PARYlation) and by localizing DNA repair effectors (e.g. XRCC1). PARP1 autoPARylation eventually leads to its own release from the site of DNA damage [9, 10].

Pharmacological PARPi structurally mimics nicotinamide, and have two general effects: (i) catalytic inhibition of PARP1 (i.e. preventing PARYlation) and (ii) locking or ‘trapping’ PARP1 on damaged DNA. Although the precise mechanisms that explain PARP1 trapping are still unclear, two have been proposed: (i) PARP1 either prevents the release of PARPi from DNA by inhibiting autoPARylation [11] or (ii) PARPi binding to the catalytic site causes allosteric changes in the PARP1 structure enhancing DNA avidity [3, 10, 12]. Either way, trapped PARP1 stalls the progress of replication forks (Figure 1). In normal, non-tumour cells, these stalled replication forks would be repaired by HR. In tumour cells that lack one of the key HR proteins, such as BRCA1, BRCA2, PALB2, or RAD51, cells use alternative DNA repair mechanisms to repair DNA lesions caused by PARP inhibition, primarily through non-homologous or micro-homology mediated end-joining (NHEJ or Alt-NHEJ). Rather than restoring the damaged DNA sequence back to its native form, the use of error-prone DNA repair pathways leads to a fragmentation of the genome that ultimately kills the cell.

Several PARPi in clinical development have different potencies as PARP1 catalytic inhibitors and as PARP-‘trappers’. It has been suggested that PARPi that are weak PARPi trappers (e.g. veliparib), fail to elicit the same scale of synthetic lethality in preclinical models, compared with effective trappers (e.g. rucaparib, olaparib, talazoparib, niraparib) [13]. At present, no clinical trial has compared head-to-head different PARPi. Of note, iniparib (BiPar Sciences/Sanoﬁ) was a compound developed as PARPi, but based on the alternative premise of altering PARP1 zinc-fingers (disturbing its activation by DNA breaks). After an unsuccessful phase III study in triple-negative breast cancer, preclinical work demonstrated that iniparib was not a bona fide PARPi [14].

In addition, PARP1 also binds directly to DNA to act as transcription factor and regulate chromatin structure remodelling. Recently, the role of PARP1 as transcriptional regulator for nuclear factors has raised some interest, due to the role of estrogen/progesterone and androgen receptors in breast and prostate cancer, respectively [15].

**PARPi development across tumour types**

**Development of PARPi in ovarian cancer**

Following the phase I trial, a phase II study assigned 33 patients with high-grade epithelial ovarian cancer (HGOC) and gBRCA mutations in a non-randomized manner to receiving olaparib at doses of either 100 (biologically active dose) or 400 mg (maximum tolerated dose) twice daily [16]. The primary end point was objective response rate (ORR), this being 33% at 400 mg b.i.d. and 13% at 100 mg b.i.d. PFS was also longer in the 400 mg cohort (median 5.8 versus 1.9 months). This dose–response relationship, was also suggested in another randomized trial (400
versus 200 mg), which also included a third arm of pegylated liposomal doxorubicin (PLD). Median PFS was 6.5 months (95% CI 5.5–10.1), 8.8 months (95% CI 5.4–9.2), and 7.1 months (95% CI 3.7–10.7) for the olaparib 200 mg, olaparib 400 mg, and PLD cohorts, respectively [17].

A multicentre phase II study enrolled 65 HGOC patients, regardless of gBRCA status. Confirmed RECIST responses with olaparib were seen in 41% and 24% of gBRCA mutation carriers and non-carriers, respectively [18], suggesting there could be a target population beyond gBRCA mutation carriers. Around 15% of HGOC are associated with gBRCA1/2 mutations (albeit regional differences). Some additional 6%–8% of patients can have tumour, not germline, BRCA1/2 mutations. Mutations in other HR genes such as RAD50, RAD51C/D, CDK12, or PALB2 are less common (<1%–3% each) [19].

All of these studies were conducted in populations previously exposed to platinum chemotherapy, the standard of care for ovarian cancer. There is a clear association between the platinum-sensitivity, defined based on the time gap from last platinum exposure to disease progression, and response to PARPi [20]. Platinum salts are DNA damaging agents and cause DNA cross-links that are in part repaired by HR. Hence, DNA repair deficient tumours are expected to be highly sensitive to platinum. At the same time, if resistant clones have restored DNA repair capacities, some cross-resistance with PARPi is expected [21]. Clinical trials in ovarian cancer shifted towards using PARPi as ‘maintenance’ therapy in patients who responded to platinum-based chemotherapy. The key initial study, olaparib’s study 19, was carried out in high-grade serous ovarian cancer patients with platinum-sensitive relapse. This study met its primary end point, with improved PFS against placebo (8.4 versus 4.8 m, HR 0.35), even more pronounced in patients with germline/somatic BRCA1/2 mutations (11.2 versus 4.3 m, HR 0.18) [22, 23]. This led to the approval of olaparib as maintenance treatment of BRCA1/2-mutated patients in 2014; in 2018, the approval was expanded to all platinum-sensitive patients, regardless of BRCA1/2 status. The confirmatory phase III trial, SOLO-2 using the new tablet formulation in gBRCA1/2 mutation carriers confirmed these results (median PFS 19.1 versus 5.5 m for olaparib and placebo, respectively) [24]. In 2018, the phase III SOLO1 trial demonstrated an unprecedented benefit for olaparib versus placebo (HR PFS 0.30) in the maintenance setting after first-line platinum-based chemotherapy in FIGO stage III/IV high-grade ovarian cancer [25]. Based on these studies, olaparib is now also approved by the FDA for first-line maintenance of germline/somatic BRCA1/2-mutated HGOC (including fallopian tube or primary peritoneal cancer). Beyond the maintenance setting, olaparib is also FDA-approved for the treatment of gBRCA1/2-mutated ovarian cancer progressing to three or more prior lines of chemotherapy.

At present, two other PARPi have received approval for the treatment of ovarian cancer patients: niraparib andrucaparib. Niraparib (previously MK4827, Merck/Tesaro) was first tested in a dose-escalation trial that established a recommended dose of 300 mg twice daily based on dose-limiting toxicities of fatigue, pneumonitis and thrombocytopenia [26]. Pharmacodynamic analyses confirmed PARP inhibition ≥50% at doses ≥80 mg/day, with responses observed ≥60 mg/day. Forty percent of 20

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**Figure 1.** Proposed mechanisms for PARPi activity in HRR-deficient cells. PARP inhibition impairs repair of single-strand breaks (SSBs) by disrupting the base excision repair (BER) pathway and also causing PARP1 trapping by inhibiting auto-PARylation and/or PARP release from DNA. These result in unresolved DNA double-strand breaks (DSBs) that in homologous recombination repair (HRR)-deficient cells lead to cell death.
gBRCA1/2 mutation carriers with ovarian cancer achieved RECIST responses, with antitumour activity also documented in non-gBRCA associated platinum-sensitive cases. The development of niraparib in ovarian cancer has focussed on the post-platinum maintenance setting, although preliminary results of a phase II trial evaluating the drug in late-stage disease have been reported [27]. The phase III NOVA trial randomized 555 patients with platinum-sensitive recurrent HGOC to maintenance niraparib or placebo treatment after second or later line of platinum-based chemotherapy [28]. The study included two cohorts: gBRCA mutation carriers (median PFS 21 versus 5.5 months, \(P < 0.001\)) and non-gBRCA carriers (median PFS 9.3 versus 3.9 months, \(P < 0.001\)). In the latter, tumour samples were tested for genomic signatures associated with HR function defects [based on the presence of areas with loss-of-heterozygosity, large-scale state transitions (LST) and telomeric allele imbalance (TAI)]. Median PFS for the non-gBRCA carriers but signature-positive patients favoured niraparib (12.9 versus 3.8 months, \(P < 0.001\)). Patients without the HR-related signature still derived some benefit (median PFS 6.9 versus 3.8, \(P = 0.02\)). These data suggest that overall platinum-sensitivity status associates with PARPi sensitivity, although more benefit is seen in patients with canonical HR defects. This study led to niraparib being granted FDA and EMA approval in the maintenance setting, regardless of BRCA1/2 status.

Lastly, rucaparib (Clovis Oncology) was initially developed for IV administration but later evolved into an orally available drug, with a recommended dose of 600 mg twice daily [29, 30]. The ARIEL3 phase III trial evaluated rucaparib as maintenance treatment following platinum-based therapy in recurrent high-grade ovarian cancer, randomizing 564 women to rucaparib versus placebo following platinum-based therapy in recurrent high-grade ovarian cancer, randomizing 564 women to rucaparib versus placebo (2:1) [31]. PFS was the primary end point, assessed in a desirous manner, being the time to disease progression or death by RECIST. Median PFS was 4.9 months for patients in the rucaparib arm versus 2.5 months for those in the placebo arm, a statistically significant difference (HR 0.58; 95% CI 0.43–0.78). The study also demonstrated that rucaparib was well-tolerated, with grade 3–4 adverse events occurring in 26% of rucaparib-treated patients and 34% of placebo-treated patients. The most common adverse events were fatigue and nausea (41% each), and anaemia (11% grades 3–4). Despite these initial encouraging results, several stumbling blocks delayed the clinical development in breast cancer [35]. In 2017, results from a randomized phase III trial (OlympiAD) in HER2-negative, gBRCA1/2-mutation-associated, metastatic breast cancer comparing olaparib with physician’s treatment choice (capecitabine, eribulin, vinorelbine, or gemcitabine) were reported [36]. Olaparib had superior median PFS [7.0 versus 4.2 months, \(HR = 0.58\) (95% CI 0.43–0.80)], with a delay on quality of life deterioration (HR 0.44; 95% CI 0.25–0.77), and a tolerable safety profile with only 4.7% rate of treatment discontinuation due to adverse events. This approach was validated in another phase III trial with talazoparib administered continuously compared with the same chemotherapy regimens [37], with a median PFS of 8.6 versus 5.6 months (HR 0.54, 95% CI 0.41–0.71). Of note, neither of the two trials were powered in their design to detect overall survival differences. Results from a phase III trial with niraparib and a similar design (BRAVO, NCT01905592) are awaited. Both olaparib and talazoparib are now approved by the FDA.

A phase II study with talazoparib (ABRAZO) [38] also demonstrated the association between antitumour activity of talazoparib and longer platinum-free interval.

Several trials are now investigatng these therapies in earlier stages of the disease, including the adjuvant setting (NCT02032823), based on the hypothesis that fewer tumour evolution-related resistance mechanisms will be present in earlier stages of disease. Talazoparib has also been tested as monotherapy in the neoadjuvant setting in patients with gBRCA1/2 mutation in a small phase II trial of 20 patients. Promisingly, the pCR was 53% and 63% when including residual cancer burden 0 and 1, respectively [39]. Conversely, veliparib, which has lower trapping potency, has not demonstrated significantly increased antitumour activity when combined with neoadjuvant platinum-based chemotherapy [40].

**Expanding the indications for PARPi to other tumour types with HR defects**

A basket trial of olaparib in patients with gBRCA1/2 mutations identified responding patients beyond the ovarian or breast cancer population, suggesting that other HR-defective tumours could be suitable for PARPi treatment [41]. Indeed, the phase I trials of olaparib, niraparib, and talazoparib have already documented responses beyond the two now approved indications. A recent study analysed the prevalence of BRCA1/2 alterations across tumour types beyond ovarian and breast cancer, in prostate, skin (non-melanoma), endometrial, pancreatic, and biliary duct cancers the prevalence was \(\geq 5\%\) [42]. Of those, clinical trials have been reported primarily for prostate and pancreatic cancer.

DNA repair gene alterations are common in metastatic prostate cancer (mPrC). Several genomic landscape studies have estimated that 8%–12% of all mPrC have BRCA2 mutations and homozygous deletions, with around half of these cases being linked to a germline mutation [43, 44]. When including other HR-related genes (such as PALB2, ATM, BRCA1, FANCA, etc.), up to 20%–25% of all mPrC harbour gene defects in different DDR pathways, which represents a significant enrichment when compared with localized, often indolent, prostate tumours [45].
The original phase I trial of niraparib included a small prostate cancer expansion cohort, with promising signs of clinical benefit [26]. Next, an investigator-initiated phase II study of olaparib in heavily pretreated mPrC patients documented responses in several patients. Using a combined definition of response including radiological responses, PSA falls, and CTC count conversions, 16/49 patients presented some degree of response. Retrospective genomic assessment of trial biopsies identified a strong association between antitumour activity and alterations in different DNA repair genes [46]. All eight patients with BRCA1/2 alterations responded to olaparib, many with durable responses for over 1 year, but mutations in genes such as ATM, PALB2, or FANCA were also observed in some responding patients. Similar data have been reported in a preliminary analysis of a phase II trial of rucaparib [47]. Based on these studies, registration trials of different PARPi in prostate cancer are now ongoing, although with different strategies when it comes to defining the putative predictive biomarker suite. Beyond their use as single agents in HR-defective tumours, PARPi is an interesting target in prostate cancer based on the cross-talk between the androgen receptor (main oncogenic driver of prostate cancer) and DDR pathways; results from phase II combination trials of PARPi- and AR-targeting agents have now been reported [48] and have led to ongoing registration studies.

In the initial phase I study of talozaparib, two patients with pancreatic cancer and mutations in BRCA2 and PALB2, respectively, achieved radiological partial responses [49]. Platinum-based chemotherapy represents one of the main systemic treatments for advanced pancreatic cancer. Two recent prospective trials treated patients with stage III/IV pancreatic adenocarcinoma and germline BRCA1, BRCA2, or PALB2 mutations with veliparib or rucaparib. Response rates were 1/16 for veliparib [50] and 3/19 for rucaparib [51]. Of note, almost all patients in these trials had previously progressed on platinum-based chemotherapy. In pancreatic cancer, positive results of a phase III trial of olaparib for maintenance therapy after response to platinum chemotherapy, following the ovarian cancer model, have recently been announced. Other strategies currently being investigated include testing PARPi in combination with chemotherapy or as radiosensitizing agents.

### Developing predictive biomarkers for PARPi

The analytical validation and clinical qualification of biomarkers able to stratify patients is of critical importance to deliver precision patient care with this drug class. During early phases of PARPi development, most efforts centred on detecting gBRCA1/2 mutations, initially using Sanger sequencing [52]. As interest has evolved beyond gBRCA1/2 mutations, multiplexed NGS assays of both germline and somatic DNA have become the preferred tool for patient identification [53, 54].

Multiplexed NGS panels assess a number of genes of interest (mostly exonic regions), either through amplification or capture-based technologies. Clinical implementation of multiplexed panels is, compared with wider whole-exome (WES) or whole-genome (WGS) sequencing, easier due to their lower cost and, critically, lower burden of bioinformatics requirements for data analysis. Accurate copy number assessment from targeted sequencing remains challenging, which is critical to demonstrating biallelic loss of DNA repair genes. This is, in part, due to intrinsic limitations in the technology but also relates to the need to integrate sample tumour purity, ploidy, and intratumour heterogeneity.

Wider WES or WGS assays add value by covering additional areas of the genome and by improving copy number and rearrangement calling. However, their applicability to clinical decision-making is still limited. Low-coverage WGS appears as a promising alternative for accurate and inexpensive copy number profiling, although it still requires significant bioinformatics analyses.

Incorporation of multiplexed biomarkers, such as DNA repair gene defects, has challenged some of the traditional concepts for biomarker validation in anticancer drug development. New genomic variants in the genes of interest are being identified, some of these being unique to individual patients. Hence, it is challenging to pursue clinical qualification of each of these separately in real time. Orthogonal methods to validate the impact on protein function of these findings is critical. Moreover, emergence of new variants also challenges traditional regulatory frameworks, demanding a continuous reassessment of the data in the post-approval setting [55]. Two main approaches could help circumvent the limitations of multiplexed biomarkers: (i) the identification of common genomic or transcriptional signatures in HR-deficient tumours and (ii) assays capable of determining functional states of the HR pathway.

Genomic signatures or scars represent repetitive patterns of DNA alterations that translate an underlying biological condition [56]. Tumours deficient for HR prioritize NHEJ for repair of DSB, resulting in more errors ligating broken chromosomal ends. These tumours accumulate small insertions-deletions and loss-of-heterozygosity events [57]. Conversely, tumours with defective mismatch repair accumulate point mutations and tumours harbouring biallelic CDK12 loss develop a particular genomic profile characterized by repetitive tandem duplications, which increase neoantigen formation [58–60].

Several assays quantifying LOH events and/or TAI across the genome have been tested as predictive biomarkers of sensitivity to PARPi. Two of them have been approved by the FDA as companion diagnostics for PARPi in ovarian cancer: the ‘FoundationFocus CDx BRCA LOH’, evaluates the frequency of LOH events throughout the genome [57], and the ‘myChoice HRD’ (Myriad), a composite signature of LOH, TAI, and LST events. Most of the data on the predictive biomarker value of such signatures were generated in the randomized trials of niraparib and of rucaparib in ovarian cancer. The aim of such signatures is to identify patients likely to respond to PARPi by harbouring an intrinsic HR defect even when no obvious BRCA1/2 alterations are detected by NGS. However, these signatures present a critical limitation: the mutational/LOH patterns do not revert when a tumour has recovered HR function, so they may not be accurate to predict PARPi sensitivity in patients who have previously received and progressed on DNA damaging chemotherapy, such as platinum.

An alternative approach is to evaluate key DNA repair protein expression. ATM expression by IHC translates ATM genomic status and is associated with survival in patients with colorectal
Figure 2. Described mechanisms of secondary resistance to PARP inhibitors. The potential mechanisms of PARPi resistance can be classified in three main groups: (i) those that result in restoration of homologous recombination repair (HRR), (ii) those leading to mitigation of replication stress (RS) commonly together with slower cell cycle progression, and (iii) other mechanisms not directly related with an HRR or RS pathway but that still alter the response to PARPi, such as mutations in PARP1 or drug effluxion pumps.
cancer treated with platinum [61]. Functional assays may directly inform on the capacity of the tumour cell to repair damage, translating upstream DNA/RNA alterations. Assays looking at accumulation of γH2AX (a marker of DNA damage) and RAD51 nuclear foci formation (indicating correct HR repair) are a gold standard for evaluating HR function in preclinical models. Their challenge raises from the need to be tested in samples with DNA damage and proliferating cells, since HR is cell cycle-related [62]. Functional assays evaluating γH2AX-RAD51 recently showed promising results in breast cancer biopsies, predicting PARPi sensitivity but also capturing the emergence of secondary resistances [63, 64].

Figure 3. Rational combinations of PARPi with other targeted agents. Hypothesis-driven combinations with PARP inhibitors are summarized; (A) combinations of PARPi with other compounds targeting alternatives DDR nodes aim to maximize accumulation DNA damage during G1 and S phases of the cell cycle, together with preventing its repair during G2 by minimizing the time to repair. This would lead to accumulation of DNA damage during mitosis and cell death. (B) Combinations with drugs targeting other biological pathways which have been shown to be modulated and/or to modulate HRR function, such as the PI3K/AKT pathway, RAS, VEGFR, and AR signalling pathways. (C) Rationale for developing PARPi-immunotherapy combinations; defects in DDR might increase genomic instability, leading to accumulation of mutations and, putatively, increased neoantigen production and T-cell activation. An alternative hypothesis supporting PARPi-immunotherapy combinations is the accumulation of cytosolic DNA induced by DDR defects, which would activate the innate immune system through the cGAS-STING pathway, inducing interferon-mediated response. This pro-inflammatory cascade would result in activation of NK cells and macrophages and the infiltration, proliferation and antitumour response of CD4+ and CD8+ T cells into the tumour. Paradoxically, the STING pathway also activates the expression of PD-L1 in tumour cells, therefore limiting the cytotoxic immune response, but potentially rendering the tumour sensitive to PD-L1 blockade (DC, dendritic cell; M\textsubscript{1}, macrophage; NK, natural killer cell; Treg, regulatory T cell).
**Acquisition of PARPi resistance**

Numerous mechanisms of acquired resistance to PARPi have been described in pre-clinical and clinical studies [65, 66]; these can be grouped into three main classes (Figure 2).

First, a number of different mechanisms result in restoration of HR function. The most common examples are secondary mutations restoring the open reading frame of HR repair genes (BRCA1/2, PALB2, RAD51C/D) in tumours with frameshift or nonsense mutations [67–70]. Expression of functional hypomorphic variants of BRCA1 has been associated with resistance to PARPi in patient-derived models [63, 71, 72]. Similarly, while epigenetic silencing of BRCA1 and RAD51C by hypermethylation of promoter regions results in PARPi sensitivity, demethylation is associated with mRNA re-expression and development of resistance [64, 73, 74]. There is also evidence in preclinical models of the co-existence of other mechanisms of resistance involving, for example, BRCA1 restoration with the removal of barriers to DNA end resection via loss of 53BP1 or proteins from the Shieldin complex, among others [75, 76].

An alternative mechanism for PARPi resistance is the protection of the replication fork, often combined with slowing cell cycle progression, as described in several preclinical studies. BRCA1, BRCA2 and PARPi, among others, play an important role in the protection of stalled replication forks, a critical step to enable replication fork repair after DNA damage [77–79]. In the absence of the aforementioned proteins, protected forks are extensively degraded, which leads to cell death. For example, BRCA2-mutant cells with loss of the MLL3/4 complex protein, were reported to become PARPi-resistant by fork protection through reduction of MRE11 recruitment to stalled forks [80]. Yazinski and colleagues further demonstrated that PARP-inhibitor resistant, BRCA1-deficient cells become dependent on ATR for survival [81].

Mutations in the DNA-binding domains of PARPi represent other likely relevant mechanisms of resistance, although clinical data are sparse [82]. Similarly, mechanisms that increase PARylation of PARPi, such as loss of PARG, could lead to PARPi-resistance by decreasing PARP trapping [83].

Beyond mechanisms rewiring the DNA damage response, increased expression of ATP-binding cassette transporters, such as the P-glycoprotein efflux pump have been shown to reduce the efficacy of PARPi [84].

**Closing remarks and future directions**

The successful development of PARPi over the last 10 years has resulted in an effective therapeutic option being available for a subset of ovarian and breast cancers, with expansion to other biomarker-driven indications expected in the near future. These treatments exploit a tumour vulnerability, that in normal conditions makes these tumours more aggressive, and constitutes one of the prime examples of success in precision medicine to date.

We envision that in the years to come the utility of PARPi will expand further. First, by better understanding what alterations beyond BRCA1/2 sensitize different tumour types to PARP inhibition; this will require combining preclinical studies and clinical trial data. Secondly, by developing more precise assays to stratify sensitive patients, capturing different predictive biomarkers and, potentially, combinations of genomic events that when co-occurring together may be relevant. Facilitating the implementation of genomics in routine clinical practice will also lead to a wider population of patients being tested. To accomplish that aim, we need to provide better resources to physicians to access these technologies and to support genomic data analyses and interpretation.

Lastly, rational drug combinations including PARPi may extend the patient population who may benefit from this drug class (Figure 3). Combinations with DNA damaging chemotherapy aim to maximize the effect of DNA damage, but they have been proved challenging due to overlapping toxicities [85]. Combination with DNA damaging radiation therapy should nevertheless be explored. Promising preliminary data have been reported combining PARPi with other targeted agents. This strategy could either aim to create synthetic lethal interactions by targeting several levels of the DNA repair machinery (i.e. combinations with ATR or HDAC inhibitors) [86, 87], take advantage of the relation between angiogenesis, hypoxia and DNA damage [88], or exploit the cross-talk between DDR and hormone receptor-driven pathways, such as ER and AR in breast and prostate cancer, respectively [89–91].

The advent of immune checkpoint inhibitors has transformed the management of several tumour types. There is a preclinical rational suggesting that PARP inhibition may trigger neoantigen and non-neoantigen-based mechanisms of tumour cell recognition by the immune system, making PARPi a potential partner for combination with immune checkpoint inhibitors [92–94]. A range of clinical trials exploring these combinations are now underway and may provide evidence of this clinical effect [95]. Optimizing dosage and scheduling of these combinations requires considering the time necessary for PARP inhibition to force tumour adaptation.

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Disclosure

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References


