Research paper

BET, SRC, and BCL2 family inhibitors are synergistic drug combinations with PARP inhibitors in ovarian cancer

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A B S T R A C T

Background: Homologous recombination deficiencies (HRD) are present in approximately half of epithelial ovarian cancers, for which PARP inhibitors (PARPi) are becoming a preferred treatment option. However, a considerable proportion of these carcinomas acquire resistance or harbour de novo resistance, posing a significant challenge to treatment.

Methods: To identify new combinatorial therapeutics to overcome resistance to PARPi, we employed high-throughput conditional RNAi and drug screening of patient-derived ovarian cancer cells. To prioritise clinically relevant drug combinations, we integrated empirical validation with analysis of The Cancer Genome Atlas (TCGA) and Genomics of Drug Sensitivity in Cancer (GDSC) datasets to nominate candidate targets and drugs, reaching three main findings.

Findings: Firstly, we found that the PARPi rucaparib enhanced the effect of BET inhibitors (CPI-203 & CPI-0610) irrespective of clinical subtype or HRD status. Additional drug combination screens identified that dasatinib, a non-receptor tyrosine kinase inhibitor, augmented the effects of rucaparib and BET inhibitors, proposing a potential broadly applicable triple-drug combination for high-grade serous and clear cell ovarian carcinomas. Secondly, rucaparib synergised with the BCL2 family inhibitor navitoclax, with preferential activity in ovarian carcinomas that harbour alterations in BRCA1/2, BARD1, or MSH2/6. Thirdly, we identified potentially antagonistic drug combinations between the PARPi rucaparib and vinca alkaloids, anthracyclines, and antimetabolites, cautioning their use in the clinic.

Interpretation: These findings propose therapeutic strategies to address PARP inhibitor resistance using agents that are already approved or are in clinical development, with the potential for rapid translation to benefit a broad population of ovarian cancer patients.

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1. Introduction

Ovarian cancer has the highest mortality of all gynaecologic cancers, with more than 22,000 new diagnoses and approximately 14,000 deaths per year in the United States alone [1]. The current standard of care consists of radical surgical resection of visible disease and treatment with platinum and taxane-based chemotherapies [2]. Although initial response rates are high, approximately 70% of patients recur [1]. Progress in identifying novel treatments for ovarian carcinoma has...
Research in context

Evidence before this study

Poly(ADP-ribose) polymerase (PARP) inhibitors (PARPi) demonstrate selective activity in ovarian cancer patients with germline or somatic BRCA1/2 mutations. Despite their success in these patients, many patients harbour de novo resistance or develop acquired resistance to PARP inhibitors. Although PARPi combinations including BRD4 or BCL2 inhibitors have been previously proposed, there have not been studies that systematically identify novel combinations by performing high-throughput conditional screening methods in the presence of PARPi using patient-relevant models of ovarian cancer. Aside from the recent FDA approval of olaparib in combination with bevacizumab for maintenance therapy in May 2020, there are currently no other clinically approved combinations with PARPi.

Added value of this study

Our study utilised a recent set of patient-derived ovarian cancer cells that closely recapitulate the genetic and molecular features of their original tumours. We performed high-throughput RNAi and drug screening in conjunction with genomic dataset analyses to reach three clinically relevant findings. Firstly, rucaparib synergised with BET inhibitors, irrespective of clinical subtype or homologous repair deficiency (HRD) status. Dasatinib, a non-receptor tyrosine kinase inhibitor, augmented the synergy of rucaparib and BET inhibitors, proposing a potential broadly applicable triple-drug combination for high-grade serous and clear cell ovarian carcinomas. Secondly, rucaparib synergised with the BCL2 family inhibitor navitoclax, with preferential activity in ovarian carcinomas that harbour alterations in BRCA1/2, BARD1, or MSH2/6. Thirdly, we identified potential antagonistic PARPi drug combinations with vinca alkaloids, anthracyclines, and antimitabolites, suggesting use of these combinations in the clinic may not be ideal.

Implications of all the available evidence

This study systematically identifies and proposes new combinations with PARPi inhibitors using agents that are clinically available or in active development, with potential to be rapidly translated to the clinic. Currently, only subsets of high-grade serous ovarian cancer patients show clear benefit from PARPi inhibitors. Here, we have identified PARPi inhibitor combinations that could benefit broader populations of ovarian cancer patients, including those with high grade serous or clear cell carcinoma, irrespective of BRCA1/2 status.

been relatively slow due to the complex genetic landscape, the lack of commonly occurring “druggable” targets, and scarcity of accurate model systems. A major breakthrough for precision medicine was provided by the discovery that carcinomas with deficiencies in homologous recombination pathway DNA repair genes such as BRCA1/2 utilise alternate PARP-dependent DNA repair pathways for survival, and are therefore more sensitive to PARP inhibition [3]. The implementation of PARP inhibitor (PARPi) therapy has transformed the standard care of treatment for many ovarian cancer patients.

Since this discovery, defects in other homologous recombination repair genes have also been demonstrated to cause sensitivity to PARPi [3–7]. Several clinical studies indicate that the broader category of homologous repair deficiency (HRD) status and/or genomic loss of heterozygosity predicts a more favourable response to PARPi in the maintenance setting (alone or in combination with bevacizumab) [8–10], which has led changes to clinical management of patients for PARPi therapy.

Unfortunately, resistance to PARPi is a clinical problem and disease progression following treatment is nearly inevitable [11]. BRCA1/2 mutated cancers can develop resistance to PARPi through somatic secondary BRCA1/2 (reversion) mutations that restore the open reading frame and functional homologous recombination repair [11–15]. Similarly, loss of BRCA1 methylation in BRCA1 methylated ovarian carcinoma provides another mechanism of PARPi resistance [16]. Other cancers are proficient in homologous recombination repair and are innately resistant to PARPi. Therefore, there is a pressing clinical need to develop therapeutic strategies that augment PARPi activity to overcome common resistance mechanisms and broaden the potential patient population that could benefit.

Previous PARPi sensitiser screens have identified candidate genes and drugs as possible combinations with PARPi; however, the use of long established cell lines raises concerns to the applicability of these results to larger patient populations [17–22]. To discover and prioritise clinically relevant drug combinations to overcome acquired or innate PARPi resistance, we have combined conditional high-throughput siRNA and drug screening employing newer patient-derived ovarian cancer cells (PDCs) that include both high grade serous ovarian carcinoma (HGSC) and ovarian clear cell carcinoma (OCCC) subtypes [23]. These PDCs were derived using an optimised culture media and have been shown to genetically and phenotypically mirror the original patient’s tumours, for example showing more than 95% identity with respect to genomic loss of heterozygosity pattern [23]. In addition, in vitro cisplatin response of PDCs correlated to the patient’s clinical response [23]. Furthermore, the HGSC PDCs have consistent copy number variation pattern and proteomic profiles when compared to primary HGSC within the TCGA, indicating a potential applicability of our findings to larger patient populations [23,24,25].

In this study, we focused on rucaparib, a potent PARPi 2/3 inhibitor that was approved by the FDA in late 2016 for treatment of patients with deleterious BRCA1/2 mutation associated recurrent ovarian carcinoma and for maintenance treatment of recurrent ovarian carcinoma following response to platinum chemotherapy [8,9,25,26]. By focusing on druggable gene targets and independent cross-validation utilising clinically-focused oncology drug libraries, we generated a functional atlas to investigate novel strategies to overcome rucaparib resistance. The results were integrated with population-based genomics for target prioritisation and validated using orthogonal assays. This multifaceted approach identified the clinically relevant drug combinations of rucaparib and navitoclax in the context of HRD, and the three-drug combination of rucaparib, dasatinib, and BET inhibitors (BETi), irrespective of HRD status (Fig. 1a).

2. Materials and methods

2.1. Cell lines and culture

The patient derived Ovarian Cancer Ince (OCI) cell lines (OCI-P5x, OCI-C5x, FCI-P2p, OCI-P9a1, OCI-P7a, OCI-C1p, OCI-C2p, and OCI-C4p) were cultured in previously defined optimal media conditions [23]. These cells are available from the Ince Laboratory at Weill Cornell Medical College and OCMl medium will be available from US Biologicals. PE01, PE04, UW81.289, UW81.289 + Vector, and UW81.289 + BRCA1 cell lines were provided by Dr. Swisher and grown in previously defined optimal media conditions [27,28]. OVCAR8 and CAOV3 cells were obtained from the American Type Culture Collection (ATCC) and cultured in RPMI 1640 and DMEM medium (Gibco) respectively, both supplemented with 10% fetal calf serum (FCS) and PrimocinTM (Invivogen). KURAMOCHI cells were purchased from Sekisui Xenotech (Kansas City, KS) and cultured in RPMI 1640 medium supplemented with 10% FCS and PrimocinTM. Mycoplasma testing was performed either through IDEXX BioResearch (Columbia, MO) or using
the MycoProbe® Mycoplasma Detection kit (R&D Systems). All cells were cultured at 37 °C in a humidified atmosphere with 5% CO₂.

2.2. DNA sequencing

Genomic DNA was isolated using the DNeasy Blood & Tissue Kit (Qiagen) according to the manufacturer’s protocol. The quantity and quality of DNA was analysed using a NanoDrop spectrophotometer. The BROCA Cancer Risk Panel next generation sequencing was performed by the Department of Laboratory Medicine at the University of Washington. This targeted panel consists of ~60 DNA repair genes, covering 320 to >1,000 sequencing reads per bp [29] (Supp. Table 1).

2.3. Gene expression profiling

Total RNA was isolated using the RNeasy Mini Kit (Qiagen) according to the manufacturer’s protocol, and the quantity and quality was analysed by the NanoDrop spectrophotometer. RNA was analysed by gene expression profiling of 770 genes, using the nCounter® Pan-Cancer Pathways panel (NanoString Technologies). Data was normalised to multiple endogenous control genes, calculating a Z-score for each gene.

2.4. siRNA screen

For the two PDCs chosen for screening, OCI-P5x and OCI-C5x, a preliminary feasibility assessment was conducted to identify optimal transfection conditions, rucaparib concentration, and readout timing. A matrix of 192 different conditions, including cell number, type of transfection reagent, siRNA ratio relative to transfection reagent, and final transfection concentrations were tested in triplicate. Viability was assessed with CellTiter-Glo® after ~1 h of incubation at room temperature. The readout timing was tested at 72 and 96 h. The conditions with the highest Z’ factor were then used for screening:
Dharmafect 1 (Horizon Discovery) as the transfection reagent at a ratio of 1:1 volume with siRNAs; the siRNA amount at 1.25 pmol, followed by a 72 and 96 h incubation for OCI-C5x and OCI-P5x, respectively.

The custom siRNA library collection was designed to target all known human kinases, DNA damage repair genes, epigenetic factors, MYC-synthetic lethal genes, and a collection of genes that are commonly altered in ovarian cancer. siRNAs were purchased from Sigma or Qiagen. In total, the cells were transfected with 2400 different pooled siRNAs targeting 2187 unique genes in three technical replicates in the presence and absence of an IC30 concentration (20 μM) of rucaparib. This concentration of rucaparib was chosen because, while it did not dramatically decrease cellular viability, it has been shown to effectively inhibit PARP activity in other cell lines [30]. Cell viability was assessed after 72 and 96 h for OCI-C5x and OCI-P5x, respectively, using CellTiter-Glo® (Promega) and was quantified using an EnVision plate reader (Perkin-Elmer). We defined sensitiser genes as those that when knocked down in the presence of rucaparib, significantly decreased cell viability by > 1.2 fold change or 20% of the vehicle. Suppressor genes were defined as those, that when knocked down in the presence of rucaparib, significantly increased cell viability by > 1.2 fold change or 20% of the vehicle.

2.5. Pathway analysis

Functional annotation of the suppressor and sensitisir genes was performed using DAVID Bioinformatics Resource 6.8 with the default threshold settings through the R package: gprofilerR_0.6.6. DAVID integrates data from KEGG, REACTOME, BioCarta, among others. To account for potential siRNA library bias, the background genes used in this analysis were the genes probed by the siRNA library.

2.6. Drug screens

The 395-compound library was purchased from Selleck Chemicals and targets a broad range of cancer-related pathways. The drug library contains FDA-approved and tool compounds that target a broad range of oncogenic processes, including PI3K, HDAC, mTOR, CDK, JAK, and RTK. Drugs were diluted to an 8-point dose curve in a broad range of oncogenic processes, including PI3K, HDAC, mTOR, CDK, JAK, and RTK. Drugs were diluted to an 8-point dose curve (20 μM) of rucaparib. This concentration of rucaparib was chosen because, while it did not dramatically decrease cellular viability, it has been shown to effectively inhibit PARP activity in other cell lines [30]. Cell viability was assessed after 72 and 96 h for OCI-C5x and OCI-P5x, respectively, using CellTiter-Glo® (Promega) and was quantified using an EnVision plate reader (Perkin-Elmer). We defined sensitiser genes as those that when knocked down in the presence of rucaparib, significantly decreased cell viability by > 1.2 fold change or 20% of the vehicle. Suppressor genes were defined as those, that when knocked down in the presence of rucaparib, significantly increased cell viability by > 1.2 fold change or 20% of the vehicle.

2.7. Drug synergy assays

OCI-P5x, OCI-C5x, OCI-P5x, FCI-P2P, OVCAR8, KURAMOCHI and CAOV3 cells were seeded in 96-well plates at 1000–2000 cells/well, incubated overnight, and treated with test compounds or DMSO controls the next day. After 6 days, cell viability was determined by incubating with CellTiter-Glo® 2.0 and luminescence was read using a BioTek H4 Synergy plate reader. Drug synergy was calculated using the Bliss and Loewe methods and graphed using Combenefit software [32]. Both methods yielded similar results, and results from the Loewe method are shown in the figures.

2.8. Clonogenic assay

OCI-P5x and OCI-C5x cells were seeded in 12-well plates at 5000 cells/well, incubated overnight, and treated with test compounds or DMSO controls the next day. After 10–14 days, colonies were fixed with 25% methanol and stained with 0.5% crystal violet. Plates were scanned prior to quantitation by dissolving stain with 10% acetic acid for 15 min and then measuring absorbance at 590 nm. Measurements were normalised to the untreated control.

2.9. Western blotting

Protein lysate preparation and SDS-PAGE were performed as previously described [33]. Immunoblot analysis was performed using the following primary antibodies from Cell Signaling Technology at 1:1000 dilution: p-SRC (Tyr416) (#6943), RRID:AB_10013641; SRC (#2123, RRID:AB_2106047), cyclin D1 (#5506), p-H2AX (Ser139) (#9718, RRID:AB_2118009), PARP (#9542, RRID:AB_2160739), cleaved PARP (Asp214) (#5625, RRID:AB_10699459), β-actin (#4970, RRID:AB_2223172). IRDye 680RD or 800CW secondary antibodies (Licor #926-68071, #926-68070, #926-32211, #926-32210; RRID:AB_10956166, AB_10956588, AB_621843, AB_621842) were purchased from Licor and used at 1:5,000 dilution. Images were analysed using the ChemiDoc MP imaging system with Image Lab software (BioRad Laboratories).

2.10. siRNA target validation

Dicer-substrate short interfering RNAs (DsiRNAs) to SRC, CDK9, BRD4, BRCA1, BCL2L1, and BCL2L2 were purchased from Integrated DNA Technologies. OCI-P5x and OCI-C5x cells were seeded in 96-well plates at 1000 cells/well and incubated overnight. The following day, siRNAs (25 nM) were transfected using Dharmafect 1 according to the manufacturer’s recommendations. After 48 h, cells were treated with rucaparib to obtain a 10-point dose curve (0.0045–30 μM). After 6 days treatment with rucaparib, cell viability was determined by incubating with CellTiter-Glo® 2.0 and luminescence was read using a BioTek H4 Synergy plate reader.

2.11. In vivo mouse study

Xenograft studies were performed at Crown Bioscience. OCI-P5x cells were engrafted subcutaneously into female NOG mice. Each mouse was inoculated at the right flank with 5 × 10^6 OCI-P5x tumour cells in 0.1 ml of PBS mixed with Matrigel (1:1) for tumour development. Grouping was performed using StudyDirector™ software (Studylog Systems). Treatments were initiated when the mean tumour size reached ~100 mm³. The drugs were administered to the tumour-bearing mice in 0.5% methylcellulose (Sigma-Aldrich).
according to the predetermined regimen (Supp. Table 2). Tumour volumes were measured twice weekly using a caliper, and volumes were determined using the formula: \( V = 0.5 \times a \times b^2 \), where \( a \) and \( b \) are the length and width of the tumour, respectively. Tumour weight was measured at study termination. Establishment of the OCI-P5x cell line was previously described [23] under approved procedures by the Institutional Review Boards of the University of Miami, Brigham and Women’s and Massachusetts General Hospitals with written consent from the patient. The protocol involving the care and use of animals in this study was approved by the Institutional Animal Care and Use Committee (IACUC) of Crown Bioscience (Pro. No. E3439-U1703). During the study, the care and use of animals was conducted in accordance with the regulations of the Association for Assessment and Accreditation of Laboratory Animal Care (AAALAC).

2.12. Statistical analysis

For analyses of the siRNA screens and drug screens, a t-test \( p \)-value < 0.05 was considered statistically significant. For validation assays, statistical analyses were performed using Graphpad Prism. Significant differences between treatment groups were determined by one-way analysis of variance followed by Tukey’s post-test. Differences were considered significant if \( p \)-value < 0.05.

2.13. Data deposition

BROCA sequencing data is available at the NCBI BioProject database (Accession No. PRJNA657639). Results from the siRNA and drug screens have been uploaded to PubChem (AID 1508597 & AID 1508598, respectively). Gene expression profiling data has been uploaded to GEO (GSE153440).

3. Results

3.1. Molecular characterisation of PDCs and functional genomic screening

To molecularly stratify seven of the ovarian cancer PDCs by DNA repair capabilities, we used a targeted-deep sequencing panel (BROCA test) (Fig. 1b, Supp. Table 1) [23,34]. Due to differences in mutational status and histology, optimal growth kinetics, and mutational status and PARPi response high-ness, we chose OCI-P5x, derived from a HGSC and OCI-C5x, harbours a deleterious BRCA1/2 mutation (i.e. PE01 or UWB1.289) (Supp. Fig. 1b-c). Despite OCRi-P5x being a top PARPi (Supp. Table 3). There were six genes in common to both PDCs, including the bromodomain-containing gene, BRD4 (Fig. 1c and d). Specific to OCI-P5x, the most represented gene families were members of the Mediator complex, MEDI4/8/10/16/23 (5 out of 27 in the siRNA library) as well as TBP-associated factors (TAFs), TAFII115/55/7/12 (5 out of 11 in the siRNA library), indicating that, in addition to BRD4, other components of transcriptional complexes may modulate the response to rucaparib and potentially other PARPi (Supp. Table 3).

Consistent with the identification of BRD4 as a common rucaparib sensitizer in the siRNA screen, the BETI CPI-203 emerged as a top drug sensitizer in OCI-P5x cells in a rucaparib conditional drug screen (Fig. 2a), where cells were screened with a library of 395 oncology-focused compounds alone or in the presence of the IC30 concentration of rucaparib (Fig. 2a). Analysis of TCGA revealed that BRD4 mRNA expression levels are highest in HGSC when compared to all other cancer types (Fig. 2b), highlighting the potential clinical utility of BETI for ovarian cancer. To further validate rucaparib and BETI as an effective drug combination, we obtained a panel of clinically relevant and commonly used BETI for testing: CPI-0610, Q1, I-BET-762, and OTX015 [39–41]. Confirming the findings of the siRNA and drug combination screen, the effect of each BETI was improved with the addition of rucaparib in OCI-P5x cells (Fig. 2c). Rucaparib and BETI synergy was also observed in 6 additional ovarian carcinoma PDCs and cell lines irrespective of HR status or clinical subtype, including OCI-C5x, FCI-P2p, OCI-C4p, OVCAR8, CAOV3, and KURA-MOCHI (Supp. Fig. 2a). Further, this synergy was not restricted to rucaparib, as two other PARPi, olaparib and niraparib, also demonstrated synergistic activity with the BETI CPI-203 and CPI-0610 (Fig. 2d, Supp. Fig. 2b).

Analysis of over 1000 cell lines within the Genomics of Drug Sensitivity in Cancer (GDSC) dataset revealed that ovarian cancer is the sixth most sensitive cancer type to the BETI, Q1 (Supp. Fig. 3a) [42]. HGSC patients with high BRD4 expression (BRD4 mRNA overexpression (Z-score > 2) or copy number amplification) have a significantly decreased overall survival than those that do not (Supp. Fig. 3b). BRD4-high alterations are also mutually exclusive with BRCA1/2 mutations in the TCGA HGSC cohort (Fig. 2e). BRCA2-mutated colorectal cancer cell lines are significantly more sensitive to BET inhibition than their wild type counterparts (Supp. Fig. 3c) [43]. Additionally, when using a recently published aneuploidy score that reflects the total number of arm-level copy-number alterations in a sample, we found that BRD4-high HGSC have a significantly higher rate of aneuploidy than unaltered BRD4 HGSC (Supp. Fig. 3d–f) [44]. These analyses are consistent with the role BRD4 plays in NHEJ and copy number variation [45,46], and suggest the mechanism of observed synergy between rucaparib and BETI could involve...
perturbing BRD4-mediated adaptation of cancer cells to genomic instability.

For *in vivo* validation, OCI-P5x cells were engrafted into NOG female mice for a four-arm drug study (Supp. Table 2). CPI-0610 was chosen as the modifying agent due to its active clinical development (NCT02158858) and demonstrated synergistic *in vitro* effects with rucaparib (Supp. Fig. 2). The combination of rucaparib and CPI-0610 resulted in a significant 58% decrease in tumour weight when compared to vehicle (Fig. 2f). Single agent CPI-0610 treatment also significantly decreased tumour weight by 42% when compared to vehicle (Fig. 2f), supporting the potential for BETi to be used for ovarian cancer. The body weights of the mice did not decrease more than 10% compared to initial values over the course of the study, suggesting the regimens used were relatively well tolerated (Supp. Fig. 4a).

Fig. 2. Rucaparib synergises with BETi in HGSC. (a) Drug screen utilising a 395 compound library in combination with rucaparib on OCI-P5x cells. AUC fold change vs. AUC Z-score of the combination; the size of the dot corresponds to the absolute AUC difference between single agent and combination. Highlighted in red are the top hits: dasatinib and CPI-203. (b) Boxplot of BRD4 mRNA expression across TCGA published datasets sorted by median expression level. (c) Response of OCI-P5x cells to IC30 rucaparib alone and in combination with the BETi CPI-0610, JQ1, I-BET726, OTX015, CPI-203. (d) Dose-response plots of olaparib and niraparib in combination with the BETi CPI-203 or CPI-0610 demonstrating synergistic drug combination effects in OCI-P5x cells. (e) Mutation enrichment of BRD4 overexpression (Z-score > 2) or copy number amplification in TCGA HGSC, demonstrating that BRCA1 and BRCA2 mutations are mutually exclusive with BRD4 overexpression or amplification. (f) Tumour weights from *in vivo* study examining the rucaparib and CPI-0610 combination using OCI-P5x xenografts. Box-and-whisker plots are presented according to the Tukey method: **p < 0.01, ***p < 0.0001, ns=not significant (one-way ANOVA followed by Tukey’s test).
3.3. Dasatinib augments the synergy of rucaparib and BETi

While single agent CPI-0610 and the combination of CPI-0610 and rucaparib significantly decreased tumour weight in vivo, tumour volume was not significantly different between the single agent and combination treatments (Supp. Fig. 4b). Further immunohistochemical analysis of the tumours showed that the mitotic index was significantly reduced in each treatment group compared to the vehicle control, but not between the single agent or combination groups (Supp. Fig. 4c). The tumours also showed a high number of cells with aberrant mitotic shapes, which are hallmarks of mitotic catastrophe (Supp. Fig. 4d–e). Despite these observations, the discrepancy between tumour weight and volume responses is unclear. It is possible that the tumours were not effectively cleared using this immuno-compromised mouse model. Xue et al have previously reported that an innate immune response in vivo promotes tumour clearance [47]. Alternatively, the rucaparib and CPI-0610 combination was not effective enough to completely suppress the growth of OCI-P5x xenografts, and there are limitations to the subcutaneous xenograft model in modeling ovarian cancer.

To explore the contribution of other tumour growth pathways in the presence of rucaparib and CPI-0610, we screened a clinical oncology drug library with this combination. This clinically focused drug library contains 131 FDA-approved oncology drugs and targeted agents in late-stage clinical development. This clinical drug library was tested on OCI-P5x cells as single agents, with rucaparib alone, CPI-0610 alone, and in combination with both rucaparib and CPI-0610. We compared the results of the drug screen to an internal database of drug responses across a panel of 54 PDCs, generating Z-scores for each compound. This method of analysis and internal PDC database has been shown to nominate drug combinations that are confirmed in patient-derived xenografts and led to tumour regression of recurrent chemoresistant cancers [31]. Confirming the results of both the conditional siRNA and high throughput drug screens, the top drug combination for the rucaparib arm was the BETi, CPI-203 (Fig. 3A). Similarly, a top combination in the CPI-0610 arm was talazoparib, another recently approved PARPi (Fig. 3b, Supp. Fig. 5a) [48]. Dasatinib, a dual BCR/ABL and SRC family tyrosine kinase inhibitor, was a top hit in all three of the combinatorial arms (Fig. 3a–c). AT7519, a multi-CDK inhibitor, was also found to synergise with rucaparib and CPI-0610 (Fig. 3a–c). AT7519 potently inhibits CDK9, a kinase that phosphorylates BRD4 and RNA polymerase II, thereby regulating their activity [49–51].

Using OCI-P5x, OCI-C5x, and OCI-C4p cells, we demonstrated that the double combinations (rucaparib and CPI-0610, rucaparib and dasatinib, CPI-0610 and dasatinib) were significantly more effective at inhibiting colony formation compared to the respective single agent treatments (Fig. 3d, Supp. Fig. 5b). The rucaparib and dasatinib combination also demonstrated synergy in these cells (Supp. Fig. 5c). However, the triple combination of rucaparib, CPI-0610, and dasatinib was significantly more effective at inhibiting colony formation compared to each double combination, except for the rucaparib and CPI-0610 combination in OCI-C5x cells, which was not significantly different to the triple combination (Fig. 3d, Supp. Fig. 5b). Rucaparib treatment alone increased phosphorylation of the proto-oncogene SRC at Tyr416 in both OCI-P5x and OCI-C5x cells (Fig. 3e, Supp. Fig. 5d), suggesting activation of the pro-survival SRC pathway by PARPi which could have cytoprotective effects and/or mediate the development of PARPi resistance. Dasatinib treatment alone or in combination with rucaparib and CPI-0610 resulted in potent reduction in phospho-SRC levels, as well as downstream cyclin D1 levels (Fig. 3e). The double and triple combinations also resulted in increased levels of p-H2AX, indicating increased DNA damage. The triple combination treatment induced significantly higher levels of PARP cleavage, indicating greater induction of apoptosis compared to the single or double agent treatments (Fig. 3e).

To verify the molecular targets of CPI-0610 and dasatinib in sensitising OCI-P5x and OCI-C5x cells to rucaparib, we knocked down expression of BRD4 and SRC using siRNA for 48 h, then treated the cells with rucaparib for 6 days. Knockdown of SRC and BRD4 enhanced sensitivity of these cells to rucaparib (Fig. 3f, Supp. Fig. 5e). Knockdown of CDK9, a target of AT7519 that also regulates BRD4 activity, also sensitised cells to rucaparib. As a positive control, knockdown of BRCA1 also sensitised OCI-P5x cells to rucaparib. Collectively, these results suggest that the three-drug combination of dasatinib, rucaparib, and BETi could be a novel and clinically relevant therapeutic strategy for ovarian cancer, regardless of HRD status and clinical subtype, by collective suppression of the SRC pathway, enhanced DNA damage and induction of apoptosis.

3.4. Rucaparib sensitises cells with alterations in DNA repair to navitoclax

Among the 75 genes that were identified as rucaparib sensitisers in the BRCA1-mutant clear cell carcinoma OCI-C5x cells, BCL2L1, an anti-apoptotic gene, was one of the most statistically significant (Fig. 1d). Consistent with the siRNA screen results, the rucaparib conditional drug screen performed in OCI-C5x cells identified navitoclax, a BCL2, BCL2L1, and BCL2L2 inhibitor [52] as the strongest synergistic drug combination with rucaparib (Fig. 4a). Separate assays confirmed the synergy between rucaparib and navitoclax in the OCCC OCI-C5x and OCI-C4p cells, while little to no synergy was observed in HGS OC1–P5x and FCI-P2p cells (Fig. 4b). To assess the generalisability of the synergy of navitoclax and rucaparib, we tested this combination in a total of 12 ovarian carcinoma PDCs and cell lines. We observed that the synergistic effect of rucaparib and navitoclax was more prominent for the PDCs and cell lines that harbour alterations in BRCA1/2, BARD1 and/or MSH2/6, which are genes involved in homologous recombination and mismatch repair, irrespective of the ovarian carcinoma subtype (Fig. 4b, Supp. Fig. 6a) [53,54]. Treatment of OCI-C5x and OCI-C4p cells with navitoclax resulted in cleavage of PARP that was further enhanced in the presence of rucaparib (Fig. 4c), indicating that enhanced induction of apoptosis contributed to the synergistic effect of this drug combination.

We did not observe a sensitising effect of rucaparib to venetoclax, the BCL2-specific inhibitor (Supp. Fig. 6b), suggesting that the navitoclax sensitisation is due to the inhibition of BCL2L1 and/or BCL2L2 rather than BCL2. To verify this, we silenced expression of BCL2L1 and BCL2L2 in OCI-C5x cells for 48h, then treated with rucaparib for 6 days. BCL2L1 knockdown dramatically sensitised cells to rucaparib, with a less potent effect observed for BCL2L2 knockdown (Fig. 4d). To assess the expression of BCL2 family genes in OCI-C5x and OCI-P5x cells, we analysed mRNA expression of 730 cancer pathway-related genes, which revealed that the pro-survival genes, BCL2L1 and BCL2, were among the genes that were more highly expressed in OCI-C5x when compared to OCI-P5x cells (Fig. 4e). To investigate the application of the experimental findings to larger populations, we performed a pan-cancer TCGA analysis of BCL2L1 and BCL2L2. We observed significant negative correlation of mRNA expression levels between BCL2L1 or BCL2L2 with BRCA1, BRCA2, MSH2 or MSH6 in ovarian cancer and a variety of cancer types (Fig. 4f, Supp. Fig. 6c). These results suggest that carcinomas that harbour defects in DNA repair may have a reliance on the BCL2 family of anti-apoptotic proteins for cellular survival, suggesting a possible biomarker of response for this combination to investigate in future preclinical and clinical studies.

3.5. Suppressor genes and chemotherapies interfering with the effect of rucaparib

The conditional siRNA screen also identified rucaparib suppressor genes, i.e. knockdown of suppressor genes increased survival in the presence of rucaparib. Identification of rucaparib suppressor genes
may inform mechanisms of action of rucaparib as well as mechanisms of resistance. The siRNA screen revealed 109 and 98 genes that suppressed the effect of rucaparib in OCI-P5x and OCI-C5x cells, respectively (Supp. Table 4). Of those, there were 12 genes in common, including SMAD1 and CDK7. SMAD1 and CDK7 are underexpressed (mRNA expression z-score < -2) in 15% and 26% of HGSC, respectively (Supp. Fig. 7a) [55]. Slightly outside the statistical cutoff for common rucaparib suppressors was ARID1A (Fig. 1c and d), a global transcriptional regulator that was recently shown to be mutated in ~67% of OCCC [35,56]. The low expression levels of SMAD1 and CDK7 in HGSC, and high mutational frequency of ARID1A within OCCC could contribute to innate resistance to PARPi.

DAVID pathway analysis identified an enrichment of genes involved with the ribosome, pyrimidine metabolism, nucleotide excision repair, DNA replication, spliceosome, and cell cycle among the rucaparib suppressors (Supp. Fig. 7b) [57]. These results suggest that agents that alter cell growth and division may interfere with the effect of rucaparib. Consistent with this idea, the rucaparib conditional drug screen in OCI-P5x cells (Fig. 2a), also found that addition of rucaparib suppressed the effect of many drugs that target the
mitotic spindle and microtubules, including vinca alkaloids, AURKA inhibitors and PLK inhibitors (Supp. Fig. 7c). Also, DNA intercalators such as anthracyclines suppressed the effects of rucaparib. These findings suggest the use of certain chemotherapies in combination with PARPi could antagonise their activity in the clinic.

4. Discussion

In this study, we employed conditional high-throughput siRNA and drug screening of ovarian cancer PDCs that faithfully recapitulate patient tumour genetics and biology to identify and prioritise effective target and drug combinations with PARPi for HGSC and OCCC. The consistent validation of the siRNA gene hits through chemical functional screens and orthogonal assays demonstrates the accuracy of the automated arrayed siRNA platform to rapidly discover novel targets and drug combinations [58].

We identified the synergistic drug combination of rucaparib with the BETi CPI-203 and CPI0610 in ovarian cancer, irrespective of HRD status and histopathological subtype, potentially broadening the clinical reach of PARPi (Fig. 2). CPI-0610-mediated BET inhibition sensitised OCI-P5x cells to rucaparib-induced DNA damage and apoptosis (Fig. 3), consistent with previous reports using other BET inhibitors [59–62]. While previous reports have focused on BRCA-mutant high grade serous ovarian carcinoma (HGSC), breast invasive carcinoma (BRCA), prostate adenocarcinoma (PRAD), and colon adenocarcinoma (COAD).

![Image](image.jpg)

Fig. 4. Navitoclax and rucaparib synergise in cancers with DNA repair mutations. (a) Drug screen utilising a 395-compound library in combination with rucaparib on OCI-CSx cells. AUC fold change vs. AUC Z-score of the combination; the size of the dot corresponds to the absolute AUC difference between single agent and combination. Highlighted in red is the top hit, navitoclax. (b) Dose-response plots of rucaparib in combination with navitoclax demonstrating synergistic drug combination effects in OCI-CSx, OCI-C4p, OVCA8, and KURAMOCHI cells, but little synergy in OCI-P5x and PCI-P2P cells. (c) Western blot analysis of PARP and cleaved PARP levels in OCI-CSx and OCI-C4p cells in response to rucaparib (10 μM) and navitoclax (1 μM) for 48 h. Blots shown are representative of at least three independent experiments. (d) OCI-CSx cells were treated with siRNA specific to BCL2L1 and BCL2L2 for 48 h. Cells were subsequently treated with rucaparib for a further 6 days before cell viability was measured. Representative of 3 independent experiments, data is presented as mean ± SEM. (e) mRNA expression levels of 770 genes in OCI-CSx and OCI-P5x cells. Highlighted in red are genes involved in apoptosis. OCI-CSx has higher BCL2L1 mRNA expression compared to OCI-P5x. (f) Correlation of BCL2L1 expression with BRCA1/2, MSH2/6, and BARD1 expression in The Cancer Genome Atlas (TCGA): high grade serous ovarian carcinoma (HGSC), breast invasive carcinoma (BRCA), prostate adenocarcinoma (PRAD), and colon adenocarcinoma (COAD).
components of the basic transcriptional machinery that directly or indirectly interact with BRD4 sensitize cells to rucaparib. AT7519, a multi-CDK inhibitor with high selectivity for CDK9, interacts with BRD4 in regulating transcription [49,50]. AT7519 synergizes with both rucaparib and CPI-0610 (Fig. 3a–c), and silencing of CDK9 sensitizes OCI-P5x cells to rucaparib (Fig. 3f). BRD4 may also have important roles in regulating the DNA damage response via NHEJ and the generation of copy number variations [45,46]. Collectively, these results suggest that the mechanism of synergy of rucaparib and BET inhibition may be due to attenuating the transcriptional functions of BRD4 that allow adaptation of cancer cells to genomic instability and/or its role in NHEJ.

We identified that double and triple combinations betweenrucaparib, CPI-0610, and/or dasatinib were effective in both OCI-P5x and OCI-C5x cells (Fig. 3), suggesting broad applicability for both HGSC and OCCC patients regardless of HRD status. The current clinical availability of these drugs points to the possibility of rapid clinical translation. Dasatinib, an FDA-approved inhibitor of SRC/ABL for the treatment of chronic myelogenous leukemia, demonstrated limited activity in a Phase II trial as a single agent in the treatment of recurrent epithelial ovarian cancer [63]. Combinations of dasatinib with carboplatin and paclitaxel have demonstrated some clinical activity, but cited concerns regarding myelosuppression and tolerability and the need for biomarker identification [64]. Future studies will seek to determine whether certain patient populations would benefit the most from rucaparib, CPI-0610 and dasatinib combinations. Assessing the level of SRC pathway activation as a biomarker for response to this combination is a possibility: phospho-SRC is elevated in OCI-P5x, FCI-P2p and OCI-C5x compared to other derived PDCs [23], and SRC amplification or high mRNA expression is observed in approximately 10% of serous ovarian cases in TCGA. A recent study also identified ARID1A-mutant OCCC cells as more sensitive to dasatinib through addiction to the SRC family kinase ABL1 [65]. With regards to mechanism, our study identified that rucaparib treatment alone increased phosphorylation of the SRC proto-oncogene in both OCI-P5x and OCI-C5x cells, pointing to the oncogenic SRC signaling pathway as a potential mediator of acquired resistance to PARPi. Inhibition of SRC signaling with dasatinib treatment alone or in combination with rucaparib and/or CPI-0610 potently reduced p-SRC and downstream cyclin D1 levels that promote cell cycle progression (Fig. 3e). Thus, the potent activity of the triple combination of rucaparib, CPI-0610, and dasatinib involves dual mechanisms that include suppression of SRC pathway-mediated cell growth and proliferation, enhanced DNA damage and induction of apoptosis. SRC activation is also induced by the veliparib and carboplatin combination in triple-negative breast cancer [66], providing a mechanistic rationale to support the use of dasatinib in combination with other PARPi and across other cancers.

This study also identified the synergistic combination of rucaparib and navitoclax in both OCCC and HGSC cells, including OCI-C5x, OCI-C4p, OVCAR8, and KURAMOCHI cells. Navitoclax is a potent BCL2/BCL2L1/BCL2L2 inhibitor currently being investigated in ovarian cancer (NCT02591095) [67]. We observed rucaparib and navitoclax synergy in ovarian PDCs and cell lines with mutations in the HR pathway genes, BRCA1/2, BARD1, and/or MSH2/6, suggesting a patient population that could benefit from this combination. We observe a negative correlation of BRCA1/2 mRNA expression with BCL2L1, BCL2L2 mRNA expression in a variety of cancer types, including ovarian, breast, colon, and prostate, suggesting elevated BCL2L1 or BCL2L2 expression may compensate for BRCA1/2 deficiency. Studies utilising other PARPi have reported synergy between talazoparib and navitoclax [68], or olaparib and navitoclax [89,70], and BCL2L1/2 overexpression mediates ovarian cancer chemotherapy resistance [70], all in the context of HGSC. Our study demonstrates the feasibility of this combination to be extended to OCCC patients, particularly in those harbouring mutations in BRCA1/2, BARD1, and/or MSH2/6. PARP inhibition has been shown to induce DNA replication stress and genomic instability [71]. We hypothesize that BRCA1/2 mutated cancers, which harbour high levels of replication stress, are more reliant upon the anti-apoptotic proteins, BCL2L1 and/or BCL2L2, for cellular survival [72] and indicate a targetable weakness. Indeed, the rucaparib and navitoclax combination results in greater induction of apoptosis compared to navitoclax alone (Fig. 4c). Further, PARPi-induced senescent cells can re-initiate proliferation upon drug withdrawal, potentially explaining the requirement for sustained PARPi therapy in the clinic [73,74]. This renders cells susceptible to secondary synthetic lethality approaches targeting the senescent state using senolytic drugs, such as navitoclax, and providing further rationale for this combination to limit or overcome PARPi resistance.

Certain chemotherapies and targeted therapies were found to potentially antagonise the effect of rucaparib in OCI-P5x cells, including vinca alkaloids, anthracyclines, antimetabolites, AURKA inhibitors and PLK inhibitors. In contrast to our findings, there is evidence in the literature to suggest that combinations between PARPi and topoisomerase inhibitors, AURKA inhibitors and PLK inhibitors are promising therapeutic strategies [75–77]. However, these studies used different drugs than those used in the present study, different methods of inhibition, or different cell types. Numerous clinical trials involving PARPi in combination with chemotherapies have been conducted or are underway [78], although successes appear to be limited by toxicity and low overall response rates. Further mechanistic studies will be essential to examine other parameters, such as cell cycle status, to determine the implications of our findings for PARPi and chemotheraphy combinations in ovarian cancer.

The siRNA screen in the present study showed that knockdown of ARID1A in both HGSC and OCCC PDCs could suppress the effect of rucaparib, identifying additional potential mechanisms of PARPi resistance. This holds potential clinical significance as PARPi are currently being investigated in OCCC and uterine endometrioid carcinoma – cancer types that harbour mutations in ARID1A –57% and ~58% of the time, respectively [35,79]. In corroboration with our results, in the ARIEL2 rucaparib clinical trial, patients harboring ARID1A-mutated carcinomas showed significantly decreased progression free survival [80]. The siRNA screen also identified novel PARPi suppressor genes, such as SMAD1 and CDK7, which are often under-expressed in HGSC and when knocked down significantly protect against the effect of rucaparib. CDK7 has been shown to be a negative regulator of BRD4, indicating a potential rationale for the suppressing function [51]. The high mutational frequency of ARID1A across cancer types and low expression levels of SMAD1 and CDK7 in HGSC constitute potential means of innate PARPi resistance.

In summary, high-throughput functional genomic and combinatiorial drug screens in ovarian PDCs revealed mechanisms of innate and acquired resistance to PARPi, highlighting potential antagonistic combinations. In both HGSC and OCCC, combinations with BETi and/or dasatinib present a novel strategy to overcome PARPi resistance through suppression of intrinsic or PARPi-induced SRC pathway signaling, enhanced DNA damage and induction of apoptosis. For patients with HR pathway mutations in BRCA1/2, BARD1 and/or MSH2/6, combinations with navitoclax and PARPi may also be a promising therapeutic approach. The studies herein provide a rationale for future efforts to assess the efficacy, tolerability, and toxicity of the proposed combinations in additional preclinical models and clinical trials. Careful assessment of dose escalation for each drug combination (both double and triple combinations) will be necessary, in addition to identification of optimal dosing regimens, to determine the translational feasibility of these findings to patients.

**Author contributions**

Study design: GYLL, RS, FXS, TAI, TCH, VKG, BAG, CJK, EMS, CG

Data collection: GYLL, RS, IS, KG, CB, RLD, RR, HAS, TCH
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Supplementary materials

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References

[1] Howlader N, Noone A, Krapcho M, Miller D, Brest A, Yu M, et al. SEER cancer statistics review, 1975–2016. Bethesda, MD: National Cancer Institute; 2017.

[2] Orr B, Edwards RP. Diagnosis and treatment of ovarian cancer. Hematol Oncol Clin North Am 2018;32(6):943–64.

[3] Lord CJ, Ashworth A. BRCAness revisited. Nat Rev Cancer 2016;16(2):110–16.

[4] Yap TA, Ashworth A. BRCA genes revisited. Nat Rev Cancer 2016;16(2):110–20.

[5] Stover EH, Konstantinopoulou PA, Matulonis UA, Swisher EM. Biomarkers of response and resistance to DNA repair targeted therapies. Clin Cancer Res 2016;22(23):5651–60.

[6] Yap TA, Plummer R, Azad NS, Hellesdy T. The DNA damage revolution: PARP inhibitors and beyond. Am Soc Clin Oncol Educ Book 2019;39:185–55.

[7] Lui GYL, Grando T, Kemp CJ. CDK12: an emerging therapeutic target for cancer. J Clin Pathol 2018;71(11):557–62.

[8] Swisher EM, Lin KK, Ozar AM, Scott CL, Giordano H, Sun J, et al. Rucaparib in relapsed, platinum-sensitive high-grade ovarian carcinoma (ARIEL2 Part 1): an international, multicentre, open-label, phase 2 trial. Lancet Oncol 2017;18(1):75–85.

[9] Coleman RL, Ozar AM, Lorusso D, Agahajianian C, Okain A, Dean A, et al. Rucaparib maintenance treatment for recurrent ovarian carcinoma after response to platinum therapy (ARIEL3): a randomised, double-blind, placebo-controlled, phase 3 trial. Lancet 2017;390(10106):1949–61.

[10] Ray-Coquard I, Pautier P, Pignata S, Perol D, Gonzalez-Martin A, Berger R, et al. Olaparib plus bevacizumab as first-line maintenance in ovarian cancer. N Engl J Med 2019;381(25):2416–28.

[11] Swisher EM, Sakai W, Karlan BY, Wurz K, Urban N, Taniguchi T. Secondary BRCA1 mutations in BRCA1-mutated ovarian carcinomas with platinum resistance. Cancer Res 2008;68(8):2581–6.

[12] Norquist B, Wurz KA, Pennil CC, Garcia R, Gross J, Sakai W, et al. Secondary somatic mutations restoring BRCA1/2 predict chemotherapy resistance in hereditary ovarian carcinomas. J Clin Oncol 2014;32(18):1890–5.

[13] Sakai W, Swisher EM, Karlan BY, Agarwal MK, Higgins J, Friedman C, et al. Secondary mechanisms as a mechanism of cisplatin resistance in BRCA2-mutated cancers. Nature 2008;451(7182):1116–20.
