Selective inhibitors of PARP1 and PARP2 (PARP1/2) are used to treat cancer patients with deficiencies in the repair of DNA via homologous recombination. Here we provide a perspective on the reported potencies of the most studied of these inhibitors (olaparib, talazoparib, niraparib, rucaparib, and veliparib) in vitro and in vivo and how these numbers relate to the known structures of these inhibitors bound to the active sites of PARP1 and PARP2. We suggest that the phenomenon of PARP trapping is primarily due to the inhibition of the catalytic activity of PARP1 and that the basis for the higher potency of talazoparib compared to the other inhibitors lies in its more extensive network of interactions with conserved residues in the active site. We also consider the potential role of the recently characterized protein “Histone PARylation Factor 1” (HPF1), which interacts with PARP1/2 to form a shared active site, for the design of the next generation of inhibitors of PARP1/2.

DNA is the genetic material found in every cell and contains the instructions for all processes required for life. DNA damage occurs at frequencies of about 1 million DNA changes per cell per day (1) and is caused by external insults such as ultraviolet (UV) radiation, reactive oxygen species, and toxic chemicals or by internal problems such as replication errors, free radicals, and spontaneous mutations. DNA damage can lead to disease, and in particular, accumulated mutations in DNA often underly tumor formation and cancer progression (2). As a reflection of the importance of maintaining genomic integrity, there exist at least 150 different proteins in the human proteome devoted to DNA repair (3). PARP1 is a key protein involved in the DNA damage response (4–7). PARP1 serves as one of the first responders by detecting single- and double-strand DNA breaks (SSBs and DSBs). Upon binding to damaged DNA, PARP1 becomes activated to utilize NAD$^+$ to add chains of poly(ADP ribose) (PAR) onto itself and other nuclear proteins, especially histones. These PAR chains recruit the appropriate DNA repair machinery, in particular proteins containing PAR-binding motifs, to the sites of DNA damage (8–10). PARP1, the most abundant nuclear protein after histones (11), has many other functions in the nucleus (12), including regulation of replication and transcription via its ability to shape chromatin structure (13), regulation of transcription through direct interactions with transcription factors, and control of cell death via depletion of NAD$^+$ (parthanatos) (14). Surprisingly, the parp1 gene is not essential for cell viability, which led to the discovery of the less abundant yet related PARP2 that assists crucially in the DNA damage response (15–18).

PARP1 and PARP2 both belong to the large class of proteins known as diphtheria-toxin–like ADP ribosyltransferases (ARTs) (19, 20). PARP1 and PARP2 (PARP1/2) both share a highly conserved catalytic domain that harbors the binding sites for NAD$^+$ and the attachment sites for the extending PAR chains (Fig. 1A). The N-terminal domains of PARP1/2, although both responsible for binding DNA, are quite different from each other (Fig. 1A). PARP1 has five known DNA-binding domains, namely three Zn fingers, one BRCT domain, and one WGR domain. The Zn1, (Zn2), Zn3, and WGR domains cooperate in the binding of damaged DNA that leads to the conformational changes required for activation of PARylation (21–23). In contrast, the Zn1, Zn2, Zn3, and BRCT domains cooperate in the binding of intact DNA that does not trigger PARylation (24). PARP2 has only two known DNA-binding domains, namely the unstructured highly positively charged N-terminal region and the WGR domain, which for PARP2 has been shown to mediate the alignment of two DNA ends in close proximity (25–27).

PARP1/2 gained much clinical interest upon the discovery that targeting these proteins could serve as the basis for treatment of breast cancer through a mechanism known as synthetic lethality (28, 29). Synthetic lethality was first observed by Bridges (30) in 1922 in the “Fly Room” at Columbia University and the term was coined by Dobzhansky (31) in 1946, both studying hidden recessive homologous phenotypes in the next generation of inhibitors of PARP1/2.

Significance
PARP is an important target in the treatment of cancers, particularly in patients with breast, ovarian, or prostate cancer that have compromised homologous recombination repair (i.e., BRCA$^{-/-}$). This review about inhibitors of PARP (PARPi) is for readers interested in the development of next-generation drugs for the treatment of cancer, providing insights into structure–activity relationships, in vitro vs. in vivo potency, PARP trapping, and synthetic lethality.
the fruit fly Drosophila. Whereas various individual mutant genes yielded viable Drosophila on their own, some crossed-bred homozygotes did not produce viable offspring. The concept of synthetic lethality was more recently expanded from describing interactions between genetic perturbations to also include a combination between a genetic perturbation and treatment with a chemical compound (32). BRCA1 and BRCA2 are known as breast cancer susceptibility genes wherein mutations in one of these resulted in an increase in the risk of developing breast cancer and ovarian cancer in women (33). Although less common, men with germline mutations in BRCA1 or BRCA2 have shown higher risks of developing breast cancer and prostate cancer than noncarriers (34). BRCA1 and BRCA2 are known to be important mediators of homologous recombination (HR), which is one of the major pathways for repairing double-strand breaks in DNA (35).

With this brief background, we can now describe synthetic lethality as it applies to BRCA1/2 and inhibition of PARP1/2 (Fig. 1B). Healthy cells carry at least one copy of the wild-type BRCA gene, which can carry out HR without the assistance of PARP1/2. Introducing an inhibitor of PARP1/2 (PARPi) into these cells does not affect their viability. Tumor cells that are lacking functional BRCA1 or BRCA2 become sensitive to a PARPi because the stalled replication forks, which cannot be restarted without the action of PARP1/2, result in single-strand breaks and eventually in the more deleterious DSB (36). Since repair of these DSBs is mediated by HR, and both BRCA1 and BRCA2 are involved in maintaining genomic integrity through HR (37), BRCA1−/− or BRCA2−/− tumor cells treated with PARPi undergo apoptosis (38–41). This mechanism of synthetic lethality provides the basis for targeted therapy for cancers that are associated with BRCA gene mutations.

There are now four different inhibitors of PARP that are approved for clinical use in breast, ovarian, and prostate cancer with underlying BRCA1/2 deficiencies, namely olaparib (Lynparza; KuDOS Pharmaceutical/AstraZeneca), talazoparib (Talzenna; LEAD Therapeutics/Phizer), rucaparib (Rubraca; Agouron Pharmaceuticals/Clovis Oncology), and niraparib (Zejula; Merck/Tesaro) (42). These same PARPi, along with veliparib (Abbott/AbbVie) and others, are part of numerous clinical trials for cancers, either as a stand-alone inhibitor or in conjunction with DNA-damaging agents such as chemo- or radiosensitizers (http://ClinicalTrials.gov) or in combination with immunotherapies (43). There have been many excellent and recent reviews of PARPi, from comprehensive coverage of all things PARP (42) to articles that focus more on the structure and activation of PARP (44), PARP trapping (45), structure–activity relationships (46), combination therapies (47), repurposing of PARPi for nononcological diseases (48), and mechanisms of resistance (49). Here we provide our unique view of the field by reviewing the literature from a perspective of measured potencies in vitro and in vivo (Number Crunching) and analysis of interactions between inhibitors and the active sites of PARP1/2 (Structure Gazing). At the close, we provide some insights on new directions in the field of PARPi.
Number Crunching

In Vitro Inhibition of PARPs by PARPi. Evaluating the potency of small molecule inhibitors is a critical part of any drug discovery effort. The requirements for assays that measure the efficacy of small molecules vary, depending on which step of the discovery process is being undertaken. High-throughput screens of large libraries of compounds that determine relative inhibition values at a fixed concentration of inhibitor need to be robust, cost effective, and amenable to miniaturization and automation. Follow-up screening assays are either similar to the assay used in the high-throughput screen or more often (and more ideally) orthogonal and allow for the determination of half-maximal inhibitory concentrations (IC50s). Mechanistic evaluation of inhibitors to determine binding constants (KDs), inhibition constants (Kis), and modes of inhibition (i.e., competitive vs. noncompetitive vs. uncompetitive) requires additional method development and is an essential step in moving inhibitors from the laboratory bench to clinical trials and ultimately patients.

Measuring the activity of PARP1/2 is not a straightforward task because these are very unusual enzymes. Most importantly, PARP1/2 are their own best substrates, which causes two major complications. First, it becomes impossible to independently vary enzyme and substrate concentrations as is typically done to optimize assay conditions for determining enzyme activity and inhibitor potency. Second, every automodification event leads to a changed enzyme. Specifically, in the automodification reaction of PARP1/2, which requires the presence of damaged DNA, ADP-ribose (ADPR) moieties from NAD+ are initially attached onto a variety of amino acid side chains (50) (Fig. 2). These protein-attached ADPRs are then extended to form PAR chains that can make polymers with up to 200 ADPRs (51). In a further complication, branch points in the PAR chains occur approximately every 20 ADPRs (51). Thus, during the course of the reaction with NAD+, every addition of ADPR onto PARP1/2 yields a different form of the enzyme that has lower catalytic activity as the enzyme is modified (52) and reduced binding to DNA once the PAR chains become sufficiently long (53–55). It is likely that automodified PARP1/2 are highly heterogeneous, i.e., that every PARP molecule in a reaction mixture carries a different set of modifications. In an additional twist, hydrolysis (“treadmilling,” Fig. 2) can also occur, and this becomes the dominant NAD+-depleting activity at high concentrations of the accessory protein Histone PARylation Factor 1 (HPF1) (below) (56).

There are many different approaches to measuring the activity, and thereby the inhibition by small molecules, of PARP1/2. One of the most widely used and most sensitive method assaying PARP1/2 relies on the use of [adenylate32P]-NAD+, which incorporates 32P-ADPR onto protein and thereby allows the facile detection of radiolabeled protein as captured in precipitates, in gels, or on filters (56–62). The availability and suitability as a substrate of biotinylated NAD+ (6-biotin-17-NAD+) allows for a colorimetric assay with detection using streptavidin that is labeled with horseradish peroxidase, although its specificity and/or activity may be altered compared to that of NAD+. Methods amenable to high-throughput screening include scintillation proximity assays (63, 64), chemiluminescent immunoassays (54), capture of NAD+ as a fluorescent adduct (65), and fluorescence polarization (55). Based on the existing literature and our own experiences, the lower limit for reliable detection of the autoPARylation activity of PARP1/2 is best attained using α-32P-NAD+ and is in the range of 20 to 50 nM enzyme. The need for such a relatively high concentration of PARP1/2 is in large part due to the limitation that PARP1/2 is its own best substrate. Having the required concentrations of PARP1/2 above the IC50s and Kis reported for clinically relevant PARP1 (10 pM to 10 nM; below) runs into the tight-binding limit problem (66). In the tight-binding limit problem, the measured inhibition constant for an inhibitor does not reflect the true binding constant (IC50s and Kis), but instead reflects the concentration of active enzyme in solution as all the inhibitor is depleted from solution by tight binding to

![Fig. 2. Chemical mechanism of PARylation by PARP1/2. In an initiation reaction, the ADPR moiety of the NAD+ substrate is attached to an amino acid side chain on PARP1/2. Elongation occurs wherein additional ADPRs are added to the existing ADPR, resulting in chain lengths up to 200. Branching of the PAR chains can also occur at the site indicated by an arrow. Additionally, PARP1 can simply hydrolyze NAD+ to yield free ADPR and nicotinamide in a reaction known as treadmilling.](https://doi.org/10.1073/pnas.2121979119)
the protein. Specifically in the case of PARP1, all inhibitors with true binding constants for PARP1 that are significantly below the concentration of PARP1 under the assay conditions will have essentially the same observed affinity, reflecting the concentration of active PARP1. In part to overcome this limitation, we have recently reported a method that allows for the accurate determination of $K_d$ for inhibitors of PARP1 that is sensitive to $\sim 10$ pM (67). Other assays that allow for the determination of subnanomolar binding constants for PARP1 include BiaCore (68, 69), although this method suffers from molecular transport issues of the inhibitor across the surface that is used to immobilize the protein (70) and small signal intensities due to the molecular weight ratio of PARPi to PARP1 ($\sim 0.4\%$).

For the “number crunching” of in vitro data, we have combed the literature for reported IC$_{50}$s and $K_i$s derived from inhibition experiments for various PARPi toward PARP1/2 and compiled these in Fig. 3\textit{A} (SI Appendix, Table S1). For PARP1, we observe relatively tight clustering of most of the experimental observations, with olaparib, talazoparib, and rucaparib being slightly more potent (median = 0.5 to 1 nM) than niraparib and veliparib (median = 4 to 5 nM) (Fig. 3\textit{A}). The significant outlier measurements indicated by arrows come from our own experiments using assay conditions developed to overcome the tight-binding limit problem (56) that is likely hampering the determination of true inhibition constants for these more potent inhibitors by most other assay methods. Supporting this interpretation, the higher potency of both talazoparib and rucaparib compared to the other PARPi that we have measured (0.012 and 0.09 nM, respectively) is also partially reflected in BiaCore experiments (0.17 nM or 0.29 and 0.09 nM, respectively) (68, 69). It should be noted that despite the fact that this assay method is based on DNA release, our results do not require the invocation of “PARP trapping,” i.e., an allosteric interaction wherein binding of compounds to the active site leads to tighter binding of DNA to the DNA domains (\textit{PARP Trapping}). Inhibited PARP1 remains bound to DNA solely because it remains unPARylated. We conclude that it is a common misconception that all PARPi have approximately equal potency. In fact, talazoparib is a significantly more potent PARP1 inhibitor in vitro than all the others, with rucaparib falling between talazoparib and olaparib (67).

The combined data for inhibition of PARP2 by PARPi yield values for olaparib and rucaparib (median = 0.2 to 0.3 nM) that are noticeably more potent than those for PARP1 (Fig. 3\textit{B}). In contrast to PARP1, talazoparib is not significantly more potent than olaparib toward PARP2 (median = 0.2 nM). As for PARP1, niraparib and veliparib (median = 2 to 4 nM) are less potent than the other three PARPi. We discuss these differences in apparent affinities of the different PARPi for PARP1 vs. PARP2 further in Structure Gazing.

\textbf{In Vivo Inhibition of PARPs by PARPi: BRCA$^{\text{--/--}}$ vs. BRCA$^{+/+}$}.

We next combed the literature for reports of PARPi in cell-based studies. Comparing different cell-based studies to each other is incredibly tricky. First, as noted in the Introduction, cells deficient in HR are expected to respond with much greater sensitivity to PARPi than cells with functioning HR. Second, different laboratories use different cell lines, some of which may harbor unknown HR deficiencies or even unexpected resistance to PARPi. Third, treatment and analyses differ widely from one study to the next, with incubation of cells with PARPi varying from 5 to 13 d and evaluation methods including cytotoxicity, clonogenic assays, or colony-forming assays. We therefore focus our number crunching of in vivo data on selected studies that compare matched cell lines (BRCA$^{\text{--/--}}$ to BRCA$^{+/+}$ or BRCA$^{+/-}$, for simplicity BRCA$^{+/-}$ since loss of HR does not occur in the heterozygous genotype).

The initial descriptions of synthetic lethality with respect to PARPi in 2005 led to the striking observation that BRCA1$^{\text{--/--}}$ or BRCA2$^{\text{--/--}}$ cell lines displayed a 60- to 1,000-fold greater sensitivity to KU0056864 (a precursor to olaparib) (28) and AG14361 (a precursor to talazoparib) (29) than BRCA$^{+/+}$ cell lines. This observation has been replicated numerous times with other cell lines and with other PARPi. For example, the potency ratio for PARPi in BRCA$^{\text{--/--}}$ to BRCA$^{+/-}$ is 250- to 300-fold in two different studies (68, 71). For veliparib and niraparib this potency ratio appears to be smaller (10- to 20-fold) (68, 71), consistent with their weaker potency and therefore presumably lower specificity. For rucaparib, the BRCA$^{\text{--/--}}$ to BRCA$^{+/-}$ ratio is an impressive 1,000-fold. For talazoparib the reported ratio is also large but more variable (560- to 10,000-fold). Interestingly, plotting the potency of PARPi as determined by our assay that overcomes the tight-binding limit problem (67) with the ratio (of the median) of these potencies shows a statistically significant correlation ($R^2 = 0.99$; Fig. 3\textit{C}). This correlation suggests that in vitro inhibition values are more predictive of cell-based efficacies than previously thought (72).

\textbf{In Vivo Inhibition of PARPs by PARPi: Thousands of Cell Lines}.

We also undertook a comparison of different PARPi in the large publicly accessible database Genomics of Drug Sensitivity in Cancer (https://www.cancerrxgene.org; release 8.3) (73).

![Fig. 3.](https://doi.org/10.1073/pnas.2121979119)
This database lists the analyses of ∼1,000 different cell lines for their response to over 500 different drugs. Although different cell lines have known and unknown underlying deficiencies in HR or other sensitizers to PARPi, several interesting observations can be made by plotting sensitivity to PARPi for different cell types according to tumors from which they were derived. First, and as noted previously (74), cells from Ewing sarcoma harboring the EWS-FLI1 gene translocation are heavily represented in the list of cell lines sorted by sensitivity to inhibition by PARPi (shown for olaparib in Fig. 4A). Unfortunately, this potential lead for treating patients with Ewing sarcoma with PARPi has not panned out in clinical trials due to lack of response to treatment (NCT01583543, NCT01286987, NCT02116777). Second, as expected, most breast cancers are not BRCA−/− and thus do not show any special sensitivity to PARPi (shown for olaparib in Fig. 4A). In fact, many of these cell lines are defective

Fig. 4. (A) Distribution of potencies for triggering cell death in 1,000 different cell lines by olaparib. The leftmost column includes all 1,000 cell lines studied in the Genomics of Drug Sensitivity in Cancer. Subsequent columns to the right display cancer-specific cell types wherein the breast cancer cell lines are not predominantly BRCA−/−. The line indicates the median value. The data were obtained from https://www.cancerrxgene.org. (B–F) Comparison of the potency of olaparib to other PARPi in a panel of 1,000 different cell types. The data were obtained from https://www.cancerrxgene.org. The IC50 values for each inhibitor were downloaded in an array and compared to the IC50 values for the other inhibitors for each cell line for which the database contained a value. Each data point represents the log IC50 value of a PARPi (B, trametinib; C, talazoparib; D, niraparib; E, rucaparib; F, veliparib) vs. the log IC50 value of olaparib for a different cell line. To facilitate interpretation and comparisons between the different graphs, the axes are “square” (one log unit is the same length on both x and y axes), and the scales are the same for all five graphs. The black line represents a best-fit linear correlation between the two inhibitors and the Pearson correlation coefficient and slope value are indicated for each graph.
in TP53BP1, which can promote partial restoration of HR and resistance to PARPi (75). Third, leukemias as a whole, and most especially B cell leukemias, are more sensitive to PARPi compared to most other cell lines (shown for olaparib in Fig. 4A) and further investigation of this lead in preclinical and clinical studies seems warranted (76). In addition, further information from this database might be gleaned using advanced computational analyses that take advantage of the complete genomic sequencing of all these cells along with these inhibition data.

Because this large dataset from thousands of cell lines contains data from different PARPi, it can also reveal interesting aspects related to the targeting and potency of the studied PARPi. We performed a meta-analysis of these data by graphing the measured potency (log IC50) of one inhibitor vs. the potency of another inhibitor, each point representing a different cell line (Fig. 4B–F). As a negative control, we show a comparison between the PARPi olaparib and trametinib, a 1-nM inhibitor of mitogen-activated protein kinase kinase (MEK). Olaparib and trametinib display no correlation, as expected since they target completely different proteins, pathways, and genetic predispositions (Fig. 4B; Pearson score = −0.014, slope = −0.023). By comparing olaparib and talazoparib (Fig. 4C), we observe a very good correlation of potencies (Pearson score = 0.81; slope = 1.3), suggesting that these two drugs are hitting the same target(s), have the same mechanism of action, and are particularly potent for the same subset of cells. Also, in this comparison one can readily see the 10- to 100-fold greater efficacy of talazoparib compared to olaparib by noting the offset between the x and y axes. This increased potency of talazoparib vs. olaparib is in agreement with our measurement of the Kd in vitro (Fig. 3A) (67). A similar correlation of potencies is observed for olaparib vs. niraparib (Pearson score of 0.72, slope = 0.83; Fig. 4D) with about a twofold lower average potency for niraparib vs. olaparib, in agreement with in vitro measurements (Fig. 3A). Surprisingly, rucaparib shows a much weaker correlation with olaparib (Pearson score of 0.41, slope = 0.31; Fig. 4E) despite its apparent equal or greater potency in vitro (Fig. 3A). We speculate that this weaker correlation may be related to the “antitrapper” effect that has been recently described for rucaparib (77) or may be indicative of off-target inhibition against other PARPi or even completely unrelated proteins that also utilize NAD+. The weakest correlation is seen in the comparison of veliparib with olaparib (Pearson score of 0.33, slope = 0.19; Fig. 4E) wherein all cell lines respond similarly and weakly to veliparib (average IC50 of 58 μM), which may help explain the inability of veliparib to make it to the clinic as a stand-alone treatment.

**PARP Trapping.** AutoPARylation of PARP1 (and PARP2) leads to the dissociation of these enzymes from DNA due to the charge repulsion between DNA and the PAR chains, which harbor twice the negative charge of DNA or RNA (53–55, 67). PARP trapping in response to treatment of cells with both a DNA-damaging agent (γ-irradiation) and a PARPi (3-amino-benzamide) was first described in 1992 wherein inhibited PARP1 remained bound to DNA due to its inability to perform the PARylation reaction (78). PARP trapping was characterized in much more detail with more potent inhibitors in response to treatment of cells with methyl methanesulfonate. Two different measures of PARP trapping are used, namely increased association of DNA with PARP1 (79) and increased association of PARP1 with chromatin (53). Interestingly, different PARPi have different potencies with respect to their ability to trap PARP1. Talazoparib is 100- to 1,000-fold better at trapping than olaparib and rucaparib, which are both better trappers than veliparib by another factor of at least 5 to 10 (53, 54, 68). PARP trapping has been broadly correlated with in vivo potency and/or toxicity as exemplified in particular for talazoparib. The reason that PARP trapping is more deleterious than persistent SSBs in the absence of PARP1 is that other repair proteins are occluded from the damage sites and the collapse of replication forks that collide with trapped PARP1.

Because it was long assumed that all PARPi have approximately equal potency against PARP1/2 in vitro, much work has gone toward understanding the biochemical basis for why different PARPi have different PARP-trapping potencies. The most likely hypothesis, and the one most pursued by a variety of experimental approaches, is the concept of allosteric coupling between the binding of DNA and PARPi (53, 54). Such an allostery implies that binding of inhibitor leads to tighter binding of DNA, and by thermodynamic necessity, tighter binding of DNA leads to tighter binding of inhibitor. However, despite best efforts using multiple methods, no evidence for such allosteric coupling as a possible cause for the potent trapping of PARPi such as talazoparib has been found (23, 68, 77, 80, 81). Surprisingly, the best evidence for coupling between inhibitor binding in the active site and DNA binding to the Zn fingers exists for rucaparib, niraparib, and veliparib (77), three PARPi that promote (not inhibit) the release of DNA (i.e., they are antitrappers). In contrast, binding of the nonclinical inhibitor EB-47 does lead to tighter DNA binding and suggests that novel PARPi could be designed to better capture this allosteric interaction (77). Based on our measurements of the higher potency of talazoparib compared to the other PARPi in vitro (67), we propose that the origin of PARP trapping is dominated by the inhibition of the activity of PARP1. That is,
PARP1 accumulates at sites of DNA damage and in the presence of PARPi remains tightly bound to DNA as there is no autoPARylation to drive dissociation. The reason talazoparib is a much more potent PARP trapper than the other PARPi is that it is a much more potent inhibitor, both in vitro (Fig. 3A) and in cells (Fig. 4B). Consistent with the idea that trapping is intimately linked to the inhibition of PARP activity, it has been recently shown that the major mechanism for the detrappping of PARP1 is its automodification at specific sites of PARylation (82).

**Structure Gazing**

**Interactions of PARPi with the Active Sites of PARP1/2.** All clinically relevant PARPi are designed to mimic the interactions between the nicotinamide ring of the substrate NAD⁺ and the active site. This approach grew out of the discovery that nicotinamide analogs such as 3-aminobenzamide were surprisingly potent inhibitors of PARP1 (83). Other early nicotinamide analogs validating this approach were characterized structurally by X-ray crystallography (61). As such, PARPi bind to the catalytic domains of PARP1/2 in competition with NAD⁺, inhibit PARylation activity, and thus prevent the subsequent release of PARP1/2 from sites of DNA damage. Detailed knowledge of the interactions between PARPi and the active sites of PARP1/2 are available from a multitude of high-resolution crystal structures (SI Appendix, Table S3). For purposes of the descriptions below, we selected the highest-resolution crystal structures of each inhibitor and compared their conserved and unique interactions, both among different inhibitors and between PARP1 and PARP2. For a point of comparison between PARPi and nicotinamide, we rely on the first structure of PARPi with a full analog of NAD⁺ (benzamide adenine dinucleotide [BAD]) that was determined in 2018 by the Pascal laboratory (Fig. 5) (84). We focus on the three highly conserved amino acids that play the most important roles in positioning the inhibitors (and the nicotinamide) at the active sites, emphasizing the importance of these interactions for the affinity of PARPi and the nonhydrolyzable BAD. Also, we describe some of the unique interactions of each PARPi that in part explain the different affinities of each inhibitor.

Comparison of all the known structures of PARPi bound to PARP1/2 revealed three conserved interactions that are shared by olaparib, talazoparib, rucaparib, niraparib, and veliparib for PARP1 and PARP2 (Fig. 6 and SI Appendix, Fig. S2). First, there are two hydrogen bonds formed by Gly863 in PARP1 (Gly429 in PARP2) with the bi- or tricyclic ring system of each inhibitor, with the amide nitrogen of Gly863 serving as an H-bond donor and the carbonyl oxygen serving as an H-bond acceptor. Second, Ser904 in PARP1 (Ser470 in PARP2) serves as an H-bond donor to a carbonyl in/on the bi- or tricyclic ring system of each inhibitor. The bidentate interaction of Gly863 and the H bond by Ser904 are the basis for the inhibitor’s mimicry of nicotinamide where these same interactions are made with the exocyclic amide of the nicotinamide (Fig. 6A). Third, each of the inhibitors forms a π–π interaction between its aromatic bi-or tricyclic ring and Tyr907 in PARP1 (Tyr462 in PARP2) with a distance of 3.5 to 3.9 Å (Fig. 6). Although this π–π interaction is not seen with the nicotinamide ring, Tyr907 does form part of the hydrophobic pocket that accommodates the nicotinamide and ribose rings of NAD⁺ (Fig. 6A). It appears that this π–π interaction contributes significantly to the much higher affinity of PARPi (nM; Fig. 3) compared to nicotinamide (>)50 μM) (83) and it may play a significant role in the binding of inhibitors with larger aromatic ring structures (e.g., olaparib) than others with smaller, perhaps less ideally placed aromatic rings (e.g., veliparib) (67).

We next examine some of the differences in how the individual PARPi dock in the catalytic pockets of PARP1/2, starting with olaparib (Fig. 6B). In addition to the aforementioned conserved interactions, olaparib forms two more hydrogen bonds with catalytic domain residues. Tyr896 makes a direct hydrogen bond with its backbone amide as well as a π–π interaction with its aromatic ring, thus clamping olaparib more snugly into the catalytic site of PARP1/2. There is also a hydrogen bond between the terminal carbonyl of olaparib and the backbone amide of Arg878, which is a water-mediated interaction in 7KK4 but seen as a direct interaction in 5DS3. This difference in H bonding is accompanied by a slightly different orientation of the “tail” of olaparib in these two structures, which most likely is not due to the absence of the HD helices in 5DS3, since olaparib bound to PARP2 has both the HD helices and a direct hydrogen bond (SI Appendix, Fig. S2A). In the structure of PARP1 with BAD this Arg878 makes a direct H bond to the adenine ring (SI Appendix, Fig. S1). The high similarity in the active sites of PARP1 and PARP2 and their similar binding modes to olaparib are consistent with their essentially identical in vitro potencies (Fig. 3).

Talazoparib is the most potent of the Food and Drug Administration (FDA)-approved PARPi and its more extensive interactions (compared to olaparib) with the active sites of PARPi provide clues for its greater affinity (Fig. 6C and SI Appendix, Fig. S2B). First, talazoparib’s tricyclic structure provides excellent conformational rigidity for the conserved interactions with the nicotinamide-binding residues Gly863, Ser904, and Tyr907. Second, talazoparib interacts with the signature His-Tyr-Glu catalytic triad of the ADP ribosyltransferases, namely His862 (His428 in PARP2), Tyr896 (Tyr462 in PARP2), and Glu988 (Glu558 in PARP2). Specifically, His862, depending on its protonation state, forms either a π–π or a π–cation interaction with the tricyclic ring. Second, the amide nitrogen of Tyr896 forms a water-mediated hydrogen bond with a nitrogen in the triazole ring of talazoparib. Third, the side chain of Glu988, which is required for the polymerization activity of PARP1 (85), forms a water-mediated hydrogen bond with nitrogen in the tricyclic ring of talazoparib. Third, the side chain of Glu988, which is required for the polymerization activity of PARP1 (85), forms a water-mediated hydrogen bond with the helical subdomain residue Glu335. One can speculate that this interaction may hinder the flexibility of the HD subdomain during the initial DNA damage response of PARP1 (21). Overall, the crystal structures of talazoparib bound to PARP1/2 support its tight binding by having a rigid molecular structure, mimicking the bindings of nicotinamide superbly and forming an extensive network of other interactions.

Although the affinity of rucaparib for PARP1 is considered similar (or even tighter) compared to that of olaparib in vitro (Fig. 3A), its weaker potency in vivo is reflected in its structure bound to PARP1 (Fig. 6D; currently, there is no structure available for rucaparib bound to PARP2). In addition to the conserved interactions with Gly863, Ser904, and Tyr907, rucaparib forms only one additional hydrogen bond with the HD-subdomain residue Glu763 at the terminal secondary amine, which was also seen with talazoparib for PARP2 (SI
The seemingly few interactions of rucaparib with PARP1 compared to olaparib (Fig. 6B) and most especially talazoparib (Fig. 6C) may in part be compensated by its rigid molecular structure. The three-ring system in the nicotinamide mimic shares some similarity to talazoparib. The least potent of the FDA-approved PARPi is niraparib (Fig. 3A). The crystal structure of niraparib is available only for PARP1, but its unique interactions in the active site provide clues for the different characteristics of niraparib. In addition to the usual nicotinamide-mimic interactions of Gly863, Ser904, and Tyr907, niraparib makes two unique interactions with HD-subdomain residues, a water-mediated hydrogen bond with Gln759 and a direct hydrogen bond with Asp766. Because the HD subdomain is conserved in PARP1, PARP2, and PARP3 (21), niraparib is considered one of the most selective inhibitors for PARP1/2 over other PARP family members (86). Thus, although niraparib’s relatively flexible molecular structure and limited interactions may limit its potency, its clamping down of the HD domain, which must be disordered for activation of PARP1 (21), allows niraparib to function as a clinical PARP1/2 inhibitor.

Veliparib has the smallest molecular weight among the PARPi and makes the least number of interactions with active site residues. For PARP1, only the conserved Gly863, Ser904, and Tyr907 interactions are observed (Fig. 6E), although in another structure (2RD6) there is an additional water-mediated hydrogen bond Glu988. Interestingly, veliparib makes two additional interactions with PARP2 (SI Appendix, Fig. S2), namely water-mediated hydrogen bonds with Glu558 and Glu335, the latter being the same interaction seen with talazoparib (Fig. 5).

Where Do We Go Next?

Because of the success (and limitations) of PARPi in the clinic, the academic and pharmaceutical research communities are actively pursuing alternative applications for existing PARPi as
The past few years have shown some big surprises in the PARP1/2 field resulting from the discovery of HPF1 by Ahel and coworkers (87). Although the mechanism or specificity of PARP1 had been studied for many decades, it turned out that the PARP research community was missing a fundamental component that dictates how this enzyme functions. First, PARylation, which for years was primarily thought to occur on serine residues, instead of autoPARylation of PARP1, reliable in vitro (and in vivo) transPARylation of histones in nucleosomes (on serine residues), instead of autoPARylation of PARP1/2 (on glutamate), was finally observed in the presence of HPF1 (50, 87). Third, the active site of PARP1/2 turned out to be incomplete without the catalytic acid Glu284 contribution of HPF1 (50, 87). Interestingly, loss of HPF1 also sensitizes cells to PARPi (82, 87, 98), suggesting that perhaps next-generation PARPi should not capture interactions with HPF1 (as suggested above), but instead compete with the binding of both NAD+ and HPF1. Given the proximity of HPF1 to the active site of PARP1/2, it would seem not that difficult to add suitable chemistry to existing PARPi that would interfere with the docking of HPF1. Combining inhibitors of sensitizer proteins with PARPi opens new possibilities for combination therapies. Mining information from screens such as those with genomic sequencing of patient samples may allow for a rapid expansion for the application of PARPi in patients with the appropriate genetic signature.

## Conclusions

With the discovery of the biochemical contribution of HPF1 to the mechanism of the PARP1/2 reaction and the outcome of PARylation, we are entering another era in the drug discovery process for inhibition of PARP1/2. We should be seeking and validating next-generation PARPi that are not mere incremental advances. As the field moves forward, it is important that solid in vitro studies are performed to prevent misguided efforts, an issue not unknown in the PARP field wherein inopportune made it to phase 3 clinical trials before it was discovered not to be a PARPi (102). Additionally, we look forward to further developments in personalized medicine that may allow for a much broader application of these well-tolerated compounds by matching compounds with appropriate cancer patients to maximize therapeutic benefits.

## Data Availability

Previously published data were used for this work (as specified in the text).
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