Exploring the genetic space of the DNA damage response for cancer therapy through CRISPR-based screens

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1. Introduction

Synthetic lethality occurs when combined mutations in two genes give rise to cell death while the single mutations do not impact cell survival. Conversely, synthetic viability occurs when phenotypic defects caused by mutations in one gene are alleviated by mutations in another gene. In suitable genetic model systems, like yeast, worms, flies and zebrafish, genetic screens have served as powerful tools to uncover such complex negative and positive genetic interactions, when they are associated with a distinct phenotype. In their formative paper, Lee Hartwell and Stephen Friend proposed that these defined molecular contexts and alterations in cancer could be exploited for new therapies [1]. Within the context of cancer, synthetic lethal interactions can represent precise and personalized approaches to treat cancers that harbor specific mutations while synthetic viable interactions can represent mechanisms of resistance to a specific lethal interaction (Fig. 1).

2. Synthetic interactions

Genomic instability is one of the hallmarks of cancer, meaning that cancer cells often have dysregulated DNA repair pathways [2]. The increased dependency on compensatory DNA repair pathways represents a vulnerability that can be targeted to specifically kill cancer cells. The concept of synthetic lethality within the DNA damage response (DDR) can be illustrated through the interaction between BRCA1/2 and PARP [3,4]. Here, mutations within the BRCA1/2 genes give rise to genomic instability due to defects in homologous recombination (HR) and enhanced replication stress. Members of the PARP family, and specifically PARP1, signal DNA single-strand breaks thus

Abbreviations
- ABE, adenine base editors; BER, base excision repair; CBE, cytosine base editors; CRISPRa, CRISPR activation; CRISPRi, CRISPR interference; Cryo-EM, cryo-electron microscopy (EM); DDR, DNA damage response; FACS, fluorescence-activated cell sorting; FDA, Food and Drug Administration; HR, homologous recombination; HTS, high-throughput screening; KRAB, Krüppel-associated box; NHEJ, non-homologous end-joining; PAM, protospacer adjacent motif; PROTAC, proteolysis-targeting chimeras; SAM, synergistic activation mediator; sgRNA, single-guide RNA; SNV, single nucleotide variant; UGI, uracil DNA glycosylase inhibitor.
activating the repair of these lesions. More recently, PARP1 has also been implicated in the repair of Okazaki fragments that occur during discontinuous DNA replication, since these fragments in essence resemble DNA single-strand breaks [5]. By removing the functions of both BRCA1/2 and PARP, cells accumulate elevated levels of DNA damage, leading to cell death.

Intriguingly, the synthetic lethal interaction between BRCA1/2 and PARP can be overcome through loss of the non-homologous end-joining (NHEJ) factor 53BP1 [6]. Loss of 53BP1 promotes processing of broken DNA ends to make them compatible substrates for HR, even in the absence of BRCA1/2. Thus, 53BP1 and BRCA1/2 are important factors that regulate whether DNA double-strand breaks will be repaired by NHEJ or HR. Similarly, loss of the Shieldin complex renders BRCA1-deficient cells resistant to PARP inhibition due to its function in promoting end-joining by restricting resection of DNA double-strand breaks and subsequent processing by HR [7–9]. These resistance mechanisms give an insight into the competition between HR and NHEJ components at DNA double-stranded break sites. Other resistance mechanisms, such as PTIP deficiency in BRCA2-deficient cells, provide information about replication fork protection. Instead of restoring HR, loss of PTIP inhibits the recruitment of the MRE11 nuclease thus protecting nascent DNA strands from excessive degradation [10].

3. CRISPR-based screens in drug discovery

The identification and use of CRISPR-Cas9 for gene editing coupled with genome-wide libraries for knock-out, knock-down, overexpression and base editing of genes has fuelled our understanding of genetic interactions (Fig. 2). Moreover, the advent of induced pluripotent cells, organoid systems and CRISPR-Cas9 gene editing to recapitulate the precise mutations present in patients have enabled the exploration of genetic space responsible for specific phenotypes.

3.1. CRISPR knockout

To initiate editing, the Cas9-single guide RNA (sgRNA) duplex unwinds DNA and searches for complementarity in the 20 base-pair region upstream to the Protospacer Adjacent Motif (PAM). Cas9 then cleaves the DNA generating a blunt-ended DNA double-stranded break. Remarkably, this breakage is an ATP and GTP independent process and instead harnesses the binding interactions between the Cas9 and the PAM [11]. The main pathway of choice following a DNA double-stranded break is the error prone NHEJ, which results in small insertions and deletions at the site of the break, typically producing a frame-shift mutation [12].

Given its scale and high-throughput nature, CRISPR knockout screens in the DDR field and elsewhere have utilized a pooled format for discovery where multiple targeting constructs are used in parallel (Fig. 3). This allows for a genome wide interrogation of gene–gene and gene–drug interactions. Following the perturbations and survival challenges, sgRNAs are amplified and sequenced to determine their relative abundance for negative or positive selection. An alternative readout to cell survival, or proliferation, is to assess functional, or phenotypic, parameters through the use of fluorescence-activated cell sorting (FACS)-based readouts. This is typically done by sorting cells
based on their expression for a particular marker. More recently, microscopy has also been utilized as a readout, for example, by measuring γH2AX foci (a marker of DNA damage) coupled with in situ sequencing [13]. Arrayed screens, which have a physical separation between each perturbation, are typically used for validation and follow-up studies.

Using pooled CRISPR-Cas9 knockout screens, PARP activity has been shown to synergize with more than BRCA1/2 loss, and to be synthetic lethal with 73 genes. These range from other components of the HR machinery, as well as factors of the Fanconi anemia repair pathway, ribonucleotide excision repair, splicing and transcription [14]. Within BRCA1/2 deficiency itself, attractive synthetic interactions have recently been identified via CRISPR screening, including CIP2A. Unlike PARP inhibition which promotes replication-induced DNA lesions that require HR repair, the CIP2A-TOPBP1 complex prevents the potentially lethal mis-segregation of acentric chromosomes [15]. Moreover, the use of DNA damaging agents in CRISPR-Cas9 knockout screens allows for the compilation of rich maps of the DNA damage network [16]. Another approach to systematically identify
gene–gene interactions using the CRISPR-Cas9 system is with a double knockout system [17]. These synergistic and buffering interaction patterns can serve as a ‘phenotypic signature’ to effectively cluster genes into pathways and complexes.

As CRISPR knockout screens become increasingly routine, databases have been curated to collect and store screening data from different institutions. One example of this is The Cancer Dependency Map (depmap.org/portal). An integration of these data sets allows for more cell lines and other models to be compiled. It also provides more statistical power for cancer specific dependencies, revealing additional biomarkers for gene dependency [18].

### 3.2. CRISPR interference

To date, most CRISPR-Cas9 screens in the DDR field have utilized gene knockouts as genetic perturbations. However, the CRISPR system has been further engineered with other protein constructs to allow for other effector functions at gene loci. CRISPR interference (CRISPRi) involves a nuclease inactive Cas9, usually abbreviated dCas9, fused with a Krüppel associated box (KRAB) domain that silences gene expression after being targeted to the promoter region. The human genome encodes for over 350-KRAB domain proteins that differ in their potency with regards to gene repression [19]. Out of the plethora of KRAB domains tested, the ZIM3 KRAB domain has proven to be a reliably strong repressor [20].

An interesting tool to precisely titrate levels of gene expression is the use of guides with mismatches in conjunction with the dCas9-KRAB construct [21]. Typically, sgRNAs with single mismatches close to the PAM region have attenuated activity compared to sgRNA mismatches in the PAM distal region. Other complex bio-physical interactions also dictate the mismatch guide activity which can been modeled using neural network predictions. Taken together, the approach of tittering gene expression could be useful especially for the DNA damage community as many DDR genes are essential, such as ATR [22] and RAD51 [23].

Unlike Cas9-mediated knockout, since CRISPRi does not induce a DNA double-strand break it does not elicit the DDR thus potentially altering the cell’s state. CRISPRi (as well as other CRISPR-based approaches) have been used in concurrent single cell approaches that determine the sgRNA and a high content transcriptional read-out. Several techniques have been developed using this approach including CROP-seq [24], Perturb-seq [25], CRISP-seq [26] and Mosaic-seq [27]. This adds a further layer of complexity allowing for a greater understanding of the gene expression landscape and signalling pathway activities following the loss of function of a specified gene. Recently, novel regulators in chromosomal instability have been identified using Perturb-seq [28]. This technology has also

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Fig. 3. An example of a pooled CRISPR screen. Cells expressing a Cas9 construct are transduced with a lentivirus sgRNA library in bulk. Following perturbation and selection, the cells are then exposed to a survival challenge (e.g. exposure to a compound). Alternatively, the population can be phenotypically characterized via microscopy or fluorescence-activated cell sorting (FACS) or analyzing the growth advantage in the population. Finally, sgRNA abundance can be determined by next-generation sequencing. Multi-omic profiling such as single-cell RNA-sequencing (scRNA-seq) can also be utilized.
been used to achieve transcriptomic read-outs with mismatch guide mediated perturbations of essential genes [21].

An important limitation to CRISPRi is the requirement for continuous expression of the dCas9-KRAB construct and the sgRNA. One technique that has overcome this obstacle is the CRISPRoff system [29] that consists of a dCas9 construct that can elicit DNA methylation and repressive histone modifications for epigenetic silencing. Transient expression of the CRISPRoff system is sufficient to target a broad range of different promoters for long-term gene repression. However, with this tool it is mostly likely not possible to obtain the different levels of gene inhibition that can be achieved with mismatched sgRNA CRISPRi.

### 3.3. CRISPR activation

CRISPR activation (CRISPRa), similarly to CRISPRi, targets the promoters of loci to induce an epigenetic change. Early CRISPRa constructs consisted of dCas9 fused to the transcriptional activator VP64 which has only very modest effects on transcriptional activation. Currently, one of most used is the synergistic activation mediator (SAM), which recruits the HSF1 and p65 transcriptional activators [30]. Other constructs include the dCas9-SunTag system [31], which recruits many copies of VP64 and the dCas9–VPR system including activator domains of the transcription factors p65 and Rta [32]. Further studies are required to assess whether the use of a mismatch gRNA in combination with CRISPRa could produce varying levels of gene activation, which may be interesting to explore.

CRISPRa libraries have been utilized to identify interactions for both PARP and ATR inhibitors. Using a genome wide CRISPRa library, overexpression of ABCB1, encoding for a multidrug resistance protein, was found to confer resistance to PARP inhibition [33]. Interestingly, acquired resistance to olaparib in ovarian cancer cell lines has been reported to occur via upregulation of MDR-1, the protein encoded by ABCB1 [34]. Using ATR inhibitors and a genome scale CRISPRa library in two human cell lines (MCF10A and HeLa), different genes were found to give rise to resistance in the two cell lines [35]. This would suggest that resistance to ATR inhibitors occurs via different mechanisms in HeLa compared to MCF10A cells.

The overexpression of oncogenes has been shown to increase cellular sensitivity to several DDR inhibitors. For example, high MYC expression, correlating to higher levels of endogenous replication stress, has been shown to sensitize lymphoma cell lines to ATR or WEE1 inhibition [36]. The synthetic lethal relationship between MYC and ATR has been demonstrated in an in vivo setting [37]. Overexpression of the cell cycle regulator CCNE1 is synthetic lethal with the inhibition of PKMYT1 kinase [38]. Thus, we propose that CRISPRa-based genetic screens could potentially be useful in identifying biomarkers of sensitivity and resistance to drugs that inhibit the DDR.

### 3.4. Base editors

Base editors are an innovative addition to the CRISPR-Cas9 toolbox. Unlike CRISPR knockout, inhibition and activation which provide ‘gene-level’ information, base editor technology allows for collecting ‘amino acid level’ information by modifying a nucleobase to incorporate a single nucleotide variant (SNV). Many different base editors have been engineered and at present, they can be divided into two classes: cytosine base editors (CBEs) and adenine base editors (ABEs) [39,40]. Along with a base modification enzyme, CRISPR base editors often employ a Cas9 nickase which produces a single nucleotide cut in the non-edited DNA strand to trigger repair. CBEs additionally typically have a fused uracil DNA glycosylase inhibitor (UGI) domain, which protects the newly formed uracil intermediate from uracil-DNA glycosylase [41]. In total, CBEs and ABEs can efficiently elicit four transition mutations in cytosine (C), thymine (T), adenine (A) and guanine (G) as follows: $C \rightarrow T$, $G \rightarrow A$, $A \rightarrow G$, $T \rightarrow C$. However, this approach cannot currently perform the eight transversion mutations $(C \rightarrow A/G, G \rightarrow C/T, A \rightarrow C/T, T \rightarrow A/G)$ [39].

There are over 75 000 known pathogenic gene variants [42]. These come in a variety of distinct categories ranging from single point mutations, duplications, copy number changes, insertions and deletions, among others. Base editors allow for the correction of transition point mutations at a target pathogenic site without the requirement of a double-stranded DNA break. A major challenge for base editing is to prevent-off target edits. Most base editors can edit with a small window to where the sgRNA is recruited to. Therefore, the gene sequence neighboring the targets of interest should be considered for gRNA design. Base editors with narrower editing windows [43,44] and more flexible PAMs [45,46] have now been developed. To predict base editing outcomes and ‘bystander’ edits, machine learning models have now been developed [47].

By utilizing base editors in a pooled screen format, more can be learnt about SNVs in DDR. Recently, different nucleotide variants of TP53 have been ranked based on their loss of tumor suppression [48]. Novel variants in ATM kinase that promote genome
instability have been described and mutations in CHK2 that were variants of unknown significance for cancer can be recategorized as loss of function mutations [49]. Treating pooled perturbations with DNA damaging agents allows for a greater understanding of variants that can change drug sensitivity [50]. Overall, base editors will be a useful asset to further our understanding of how SNVs underpin DDR related diseases, cancer predisposition and sensitivity to DDR inhibitors and DNA damaging agents.

4. Advancements in chemical compounds

In addition to exploring genetic space, chemical space can also be explored in cancer therapy. High-throughput screening (HTS) has enabled automation and miniaturized bioassays; thus, a large number of candidate compounds or genetic modulators can be assessed for a specific biomolecular activity. In addition, HTS can also support improved understanding of biochemical processes and as such has thus become a powerful discovery tool that relies on suitable cellular systems, which can reflect the disease mechanism. Coupled with advances in developing chemical compounds, HTS has led to improvements in drug discovery, ranging from synthesis and mode of action, all the way to drug metabolism and side effects.

4.1. Small molecule inhibitors for the DDR

4.1.1. PARP inhibitors

PARP inhibitors are the first and best characterized targeted therapy utilizing synthetic lethality in the DDR. To date four PARP inhibitors have been approved by the Food and Drug Administration (FDA): talazoparib,rucaparib,niraparib and olaparib all of which are NAD+ analogs that compete with NAD+ for binding of the PARP-1 catalytic domain [51]. When a given PARP inhibitor binds to PARP, firstly, PARP can no longer PARylate other proteins involved in the repair process. Second, the completion of base excision repair (BER) and DNA single-strand break repair (SSB) requires the dissociation of PARP from the DNA. Consequently, inhibition of auto-PARylation results in PARP ‘trapping’ on DNA. When a replication fork approaches, DNA SSBs with bound PARP are converted to DSBs which cannot be processed in HR-deficient cells [52]. PARP inhibitors vary in their trapping abilities which correlates with cytotoxicity [53]. Talazoparib has the greatest PARP trapping capability out of the four FDA approved drugs [54]. Each of the PARP inhibitors approved, whilst largely target PARP1, has unique off-target effects with other members of the PARP family and has varying pharmacological profiles across the kinome, the set of protein kinases encoded by the genome [55].

With the implementation of pooled CRISPR screens, BER intermediates, including FEN1 and APEX2, have been identified to enhance sensitivity of PARP inhibition in HR-deficient backgrounds [56,57]. This is proposed to be because abasic sites and DNA SSBs accumulate in BER-deficient backgrounds and provide a source for PARP trapping. Another important determinant of PARP sensitivity identified in screens is the chromatin remodeler ALC1/CHD1L [58]. In the absence of ALC1, PARP is retained on chromatin further enhancing PARP trapping.

Despite good initial response rates with PARP inhibitors, many cancers eventually develop resistance. Numerous mechanisms of resistance have been noted in preclinical studies including the full or partial restoration of BRCA [59,60], restoration of HR in BRCA1 deficient cells [61,62] fork protection in BRCA2-deficient cells [63] and rescues in both PARP trapping [64] and PARylation activity [65]. Inhibitors of poly(ADP-ribose) glycohydrolase (PARG) have generated interest as a mechanism of overcoming PARylation signaling-based resistance [66,67].

The development of a particular resistance mechanism can potentially confer vulnerabilities in other DNA repair pathways and thus provide targets for second-line therapies. For example, BRCA1-deficient cells that lose Shieldin/53BP1 to restore HR become hypersensitive to ionizing radiation and cisplatin [7]. Additionally, BRCA1-deficient cells are hypersensitive to the depletion of nucleotide sanitizers such as DNPH1, resulting in the incorporation of the aberrant nucleotide hydroxyuridine [68]. Taken together, understanding which PARP inhibitors should be used to treat a particular cohort, the development of more selective PARP inhibitors, investigating PARP beyond BRCA1/2, and knowledge on resistance mechanisms in the clinical setting will be important for improving clinical outcomes.

4.1.2. ATM, ATR and DNA PKcs inhibitors

ATM, ATR and DNA PKcs, previously described as the trinity at the heart of the DNA damage response [69], are attractive targets for cancer therapeutics. ATR, despite being an essential kinase, has therapeutic relevance and several inhibitors have been synthesized...
and tested in clinical trials. These include berzosertib (also known as M6620 or VX-970), ceralasertib (AZD6738), BAY-1895344 and more recently M4344. Genome wide interrogations using CRISPR knockout have highlighted potential biomarkers for resistance including the loss of CDC25A \[70\] and loss of cyclin C and CDK8 \[71\] to ATR inhibition.

The kinase ATM, an important protein in signaling DNA DSBs, has three inhibitors that are currently tested in clinical trials. These include AZD0156, AZD1390 and M3541. AZD1390 has particularly good blood–brain barrier permeability profiles \[72\] and is being tested in clinical trials for gliomas (NCT05182905). Both AZD1390 and M3541 are being tested in combination with radiotherapy. Inhibiting ATM kinase activity has been shown to sensitize cells to radiation \[73\] also supported by the sensitivity of Ataxia–telangiectasia patients to radiation \[74\].

DNA-PKcs is a key component in NHEJ and similarly to ATM, inhibition of this kinase sensitizes cells to radiation and synergizes with topoisomerase inhibitors \[75\]. Various DNA-PKcs inhibitors that have entered clinical trials including CC-115, M3814 (neldesertib or peposertib) and AZD7648. An important recent development was the first cryo-electron microscopy (EM) images of DNA-PKcs bound to various inhibitors and ATP analogs \[76\]. Like many kinase targets, the current drug candidates have been developed from high-throughput screening targeting the ATP-binding site. However, molecular details of the modes of action of such candidates have been unclear. This is particularly the case for DNA-PKcs given its large size. From the cryo-EM images generated, precise inhibitor binding can be elucidated as well as the effect this has on DNA-PKcs dimerization and its higher order structures.

4.1.3. Checkpoint and cell cycle inhibitors

Targets of CHK1 and CHK2, downstream of ATR and ATM, are important kinases involved in coordinating the DNA damage response and cell cycle arrest. Several compounds have been tested in clinical trials, many of which have shown low to modest anti-tumor effects alongside toxicity, including AZD7762 \[77\] and GDC-0575 \[78\]. Prexasertib (LY2606368), an ATP competitive inhibitor of CHK1, has shown promising potency in recent trials \[79\]. Further trials with this inhibitor are currently ongoing (NCT04095221, NCT04023669, NCT02649764).

The WEE1 kinase phosphorylates and inhibits CDK1 and CDK2, thereby exerting control over the intra-S and G2-M checkpoints \[80,81\]. Recently, WEE1 has been described to protect against fork degradation at stalled replication forks \[82\]. Cells with higher levels of endogenous replication stress, for instance promoted by high expression of MYC and cyclin E \[36\], were shown to be hypersensitive to WEE1 inhibition. Loss of WEE1 activity is synthetic lethal with diminished histone H3K36me3, and thus cancers with low levels of H3K36me3 benefit from WEE1 inhibition \[83\]. This is due to a reduction of RRM2, a ribonucleotide reductase subunit, leading to a depletion in the nucleotide pool. This potential biomarker is being tested in a phase 2 clinical trial on advanced solid tumors (NCT03284385). AstraZeneca’s WEE1 inhibitor, AZD1775, has been tested on a wide range of tumors and has shown promising results in a uterine carcinosarcoma study \[84\]. Currently, there are 26 clinical trials that are recruiting or are active in order to test AZD1775 in human subjects (clinicaltrials.gov).

PLK1 is also a kinase involved in the G2-M checkpoint and overexpressed in many cancers \[85\]. Volasertib is the most advanced PLK1 inhibitor in clinical development with recent phase 2 and 3 clinical trials in acute myeloid leukemia, among others. In combination with chemotherapies, volasertib has had variable response rates \[86,87\]. An orally available ATP competitive inhibitor of PLK1 is currently undergoing clinical trials including in colorectal cancer with KRAS mutations (NCT03829410). Like with many of the DDR inhibitors, biomarkers for sensitivity are in short supply. A recent CRISPR knockout screen has highlighted a potential non-canonical cell cycle gene, ARID1A, that when lost promoted sensitivity to PLK1 inhibition \[88\]. Additionally, high PRC1 expression, which correlated with poor survival, increased sensitivity to PLK1 inhibition in Ewing sarcoma \[89\].

4.1.4. New and emerging small-molecule inhibitors for the DDR

The DNA polymerase POL\(h\) is upregulated in a number of cancers and is associated with a poor prognosis \[90,91\]. The polymerase has several described functions involving POL\(h\)-mediated end joining \[92\] and translesion polymerase synthesis \[93\]. Inhibition of POL\(h\) is synthetically lethal with BRCA1/2 deficiency \[94\]. One hypothesis for this is that the loss of both HR and POL\(h\)-mediated end joining that provide a synthetic lethal relationship. An alternative hypothesis is that POL\(h\) functions in processing the single-stranded DNA gaps that occur at replication forks in a BRCA-deficient background. POL\(h\) inhibition may also provide a therapeutic option to treat BRCA-deficient
cancers that are resistant to PARP inhibition due to mutations in 53BP1/Shieldin [95]. POLh contains a polymerase and a helicase domain, both of which are druggable targets [96,97]. Inhibition of either the polymerase domain [95] or the helicase domain [98] of POLh has been reported to benefit BRCA deficient cancers. Building on these studies, Artios Pharma recently dosed their first patient in a phase 1/2a study (NCT04991480) with a POLh inhibitor targeting the polymerase domain (https://www.artiospharma.com/2021/09/28/artios-doses-first-patient-in-phase-1-2a-study-of-pol%CE%B8-inhibitor-art4215/). Conversely, novobiocin, an antibiotic first described in the 1950s that inhibits the helicase domain of POLh, is currently being tested in clinical trials [98]. Additionally, POLh pre-clinical programs have been disclosed by various pharmaceutical companies including Ideaya, Breakpoint Therapeutics and Repare Therapeutics.

KSQ pharmaceuticals has announced a first dose in a cancer patient in 2021 with a USP1 inhibitor. A phase 1 clinical trial of KSQ-4279 is currently recruiting both as a monootherapy and in combination with PARP inhibition in patients with advanced solid tumors (NCT05240898). In addition, other companies, including Tango, have disclosed a USP1 inhibitor program. USP1 is a deubiquitinase implicated in DNA repair. Along with USP1 associated factor 1 (UAF1), it removes monoubiquitin signals from PCNA, FANCI and FANCD2 [99–101]. USP1 is synthetic lethal with BRCA1-deficient cell lines and is involved in binding and stabilizing replication forks [102]. Like POLh inhibition, it may provide a potential avenue for exploiting HR deficiency.

Other companies are also expected to announce their first patient dosing of new drugs including Cyteir Therapeutics’ RAD51 inhibitor (NCT03997968), Repare Therapeutics’ PKMYT1 inhibitor (NCT05147272) and ATR inhibitor (NCT04497116), Artios’ ATR inhibitor (NCT04657068), WEE1 inhibitors from Debiopharm (NCT03968653) and Zentalis (NCT04158436). Inhibitors for the WRN helicase, which had generated a lot of interest due to being synthetically lethal with microsatellite instability [103–107], are currently in the discovery phase in several companies (https://www.nimbustx.com/pipeline-targets/, https://www.beactica.com/pipeline, https://www.xposetx.com/pipeline, https://ryvu.com/pipeline/).

4.2. Targeted protein degradation

A significant challenge in ‘occupancy driven’ pharmacology is the observation that many proteins lack a ‘druggable’ pocket in which small molecule inhibitors can bind. An alternative promising strategy is proteolysis-targeting chimeras (PROTACs). These drugs hijack the ubiquitin proteasome system and promote the degradation of a target protein [108]. This effectively removes all possible functions of the protein as opposed to occupying a catalytic or allosteric domain [109]. PROTACs are bifunctional. They contain a small-molecule binder, also commonly called a warhead, which attaches to the target protein, and E3 ubiquitin ligase binding domain which are held together by a linker. This induces proximity between the target protein and the E3 ubiquitin ligase binding domain which are held together by a linker. This induces proximity between the target protein and the E3 ubiquitin ligase, thereby promoting its ubiquitination and degradation (Fig. 4).

![Fig. 4. A target protein degraded using a proteolysis targeting chimera (PROTAC). The induced proximity of the target protein and the E3 ligase promotes the polyubiquitination of the target protein. This post-translational modification signals for its degradation. The PROTAC can be catalytically recycled.](image-url)
Another targeted proteolysis strategy includes molecular glues. These are molecules that stabilize the interaction between two proteins without showing a detectable affinity towards (at least) one of the binding partners. They do not contain a linker and may not directly interact with the protein of interest. Some interact with the E3 ubiquitin ligase, thereby influencing the enzymes interaction interface, resulting in the recruitment of the target protein [110]. Molecular glue discovery is challenging and would require a backwards approach involving screening E3 ligase binding molecules or large protein panels for a desired phenotype.

No DDR PROTACs or molecular glues are currently in clinical trials. PROTACs of PARP have been tested in vitro [111]. Recently, a range of PROTACs for WEE1 have been developed [112]. This was achieved using AstraZeneca’s AZD1775 as the warhead, E3 ligase binders of VHL and CRBN along with linkers of different lengths and compositions. Recently, CDK12-cyclin K PROTACs and molecular glues have shown synergistic effects with DNA damage-inducing drugs [113–115].

An increasing amount of proteolysis therapeutics are expected to enter clinical development. Chemoproteomic approaches and fragment-based ligand discovery will accelerate identifying new relevant ligands [116,117]. Identifying functions for E3 ubiquitin ligases will be important as the majority remain poorly understood. Targeting a given protein for degradation will provide its own unique challenges regarding the emergence of resistance. Modifications in the core component of the E3 ubiquitin ligase is one mechanism by which this can occur [118,119].

5. Future perspectives

The ability to introduce specific mutations using CRISPR-based approaches represents an exciting opportunity to investigate how alterations in DNA contribute to resistance or cancer development. Here, we have described how base editors can be used to gain mechanistic insights at the single nucleotide level, yet these approaches are limited with regards to the spectrum of mutations that can be introduced. The discovery of prime editing, a genome editing technology, that can introduce a wide range of mutations has the potential to be very useful in studying disease-relevant mutations [120] yet its reduced efficiency has precluded it from being used in screens. However, much effort is directed towards making prime editing more efficient and thus future CRISPR-based screens might include prime editors [121,122].

Here, we have largely discussed the implementation of CRISPR-based screens in cancer cell lines. Since they better mimic in vivo conditions, developments in the establishment and utilization of organoids represent an exciting opportunity in which to perform such screens. Indeed, genome-scale CRISPR screens have been reported in human intestinal organoids [123] as well as mouse stomach organoids [124] thus highlighting the feasibility of such an approach. Yet, such 3D systems have their limitations. Consequently, researchers are moving towards performing CRISPR screens in vivo to unravel complex signaling networks within their physiological environment [125,126]. Improvements in the performance of in vivo CRISPR screens will lead to a wider use thus allowing genetic space to be explored in more clinically relevant settings.

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Conflict of interest

The authors declare no conflict of interest.

Author contributions

JIL conceptualized the manuscript with JW. Both JW and JIL contributed equally to writing the manuscript.

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