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# Targeting homologous recombination addicted tumors: challenges and opportunities

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Abstract: Recent advances in next generation sequencing (NGS) and molecular subtyping of tumors have opened the door to clinically available targeted therapies. Although the treatment of many solid tumors still rely on a steady regimen of non-targeted chemotherapeutic agents, it is becoming increasingly more apparent that certain tumors with defects in DNA damage repair (DDR) genes may be exquisitely sensitive to DNA damaging agents or therapies targeting key elements of this pathway such PARP1, ATR, or ATM. Still, for tumors with DDR defects the challenges are multi-fold including: (I) identifying these tumors in patients in time for a window of opportunity of treatment; (II) ensuring that these tumors are still reliant or addicted to this pathway; and (III) making sure these tumors are matched with the precise treatment option. Herein, we will discuss the opportunities, challenges, and future of targeting a subset of DDR-defective tumors.

Keywords: Precision medicine; pancreatic adenocarcinoma; DNA damaging repair genes

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### Background on the discovery, cloning, and importance of genome maintenance and DNA repair genes

### *Linking BRCA1/2 to cancer predisposition: Let the games begin*

In the early 1990s, *BRCA1* was mapped to chromosome 17 (1), and thus, initiated the international effort to link half the families from a consortium with breast and ovarian cancers (an inherited predisposition) to this loci (2). This investigation, amongst others, pioneered the efforts to discover inherited germline mutations in genes that could explain the reason why some families had a strong predisposition to certain tumors. For instance, it is well established that individuals who carry a germline heterozygous *BRCA1* (or *BRCA2*) mutation can have a lifetime risk of ~85% of developing breast and ovarian

cancers (3), and a higher incidence of developing pancreatic and prostate cancers than the general population. These genetic linkage studies were not only enlightening for genetic testing and identifying cancer predisposition syndromes, which have highlighted the importance of DNA repair for cancer prevention (4), but also opened the field to a new understanding about the intersection between a defective DDR pathway and tumorigenesis. Finally, the culmination of this work plus the sequencing of thousands of cancer genomes have underscored the importance of the DDR pathway. For instance, to date the clinical concept of personalizing therapy for the treatment of DDR defective tumors is very much in vogue.

Although *BRCA1/2* have been implicated in a number of cellular functions, the fact that these genes are mutated with a high frequency in tumors and have a well described genome maintenance and repair mechanism,

has led the field to focus on its role in tumorigenesis (5). Specifically, *BRCA1/2* have been established to play a key role in collapsed replication forks and double strand DNA break (DSBs) via homologous recombination (HR), where the damaged site in DNA sequenced is repaired in a conventional manner. In a deficient BRCA1/2 setting, cells use a faulty, alternative repair mechanism, and this is believed to facilitate genetic instability and tumorigenesis (6).

### *From genes to an homologous recombination deficiency (HRD) signature/scar: are all DDR genes mutated equally?*

The discovery of DDR genes linked to cancer predisposition and tumorigenesis has forced NGS panels to expand personalized approaches to think beyond BRCAness (i.e., *BRCA1/2* genes). However, there are limitations to just trying to expand the panel of DDR genes. First, it is unclear whether low frequency mutated DDR genes (or even variants) are truly driver alterations of tumorigenesis. Unfortunately, in many instances, including *BRCA1/2*  mutated tumors, the frequency of mutations found in a specific tumor type, may be very low compared to more garden variety cancer driving genes (e.g., *Kras* or *TP53*), making it hard to decipher whether these events are frequently selected for in a given patient cohort. Based on the central dogma of conventional cancer genetics, one must have a mutation frequency in a tumor type which is greater than expected in a cohort of healthy controls (7). Other layers of complexity include whether these DDRrelated genes will have the same Achilles Heel (also known as synthetic lethality) that established DDR genes such as *BRCA1/*2 have and whether these genes fit the classical tumor suppressor rules of needing to have loss of the second allele in the tumor (e.g., Loss of Heterozygosity, LOH) (7). Therefore, in many DDR genes, it is unclear whether these DDR defective genes have predictive therapeutic value. Based on these concerns, many investigators have attempted to design assays that detect a molecular signature that could identify tumors with a defective DDR pathway (i.e., HRD, see below section).

Specifically, a number of additional genes have been implicated beyond *BRCA1/2* genes in the HR pathway. It should be noted that these mutations can be categorized and identified as either germline (i.e., constitutional DNA, identified in the normal blood or presumable inherited) or somatic (i.e., found in the tumor DNA, but not in the germline DNA) in nature. As the field rapidly moves into

the next versions of NGS, many of these early concepts or the central dogma of cancer genetics sometimes gets lost in the commercialization of personalized medicine. The gold standard for identifying an actionable tumor suppressor or genome maintenance gene mutation, in this instance would be a germline mutation (i.e., that causes an amino acid change) combined with a second, somatic hit mutation in the other allele, typically via an LOH event. Additional considerations include identifying whether a somatic change is authentic based on the availability to obtain pure DNA from the tumor (without contaminating normal cells that can confound sequencing results) and matched normal tissue in order to compare with tumor sequencing results. An additional level of complexity on determining whether these HR-gene mutations are authentic, is the fact that many are considered Variants of Unknown Significance (VUSs). Many VUSs have been uncharacterized and even with the confirmation of a potential second allele hit, the functional significance of these VUSs may either be linked to a non-driving Single Nucleotide Polymorphism (SNP) or a more prevalent previously described mutation from a patient cohort.

With that written, still a number of DDR-HR related genes have been well described and established. In a recent publication focused on Homologous Repair-DDR genes looking at over 52,000 tumors of different origins (e.g., biliary tract, GI stromal, bladder, etc.) were analyzed to identify pathogenic mutations in the HR-DDR genes including *BRCA1/2*, *BRIP1*, *RAD50*, *RAD51*, *RAD51B*, *ATM*, *ATRX*, *BARD1*, *CHEK1/2*, *FANCA/C/D2/E/F/G/L*, *MRE11A*, *NBN*, and *PALB2* (8). In this study, they found that the overall frequency of these mutations were over 17% across 21 tumor lineages. Other studies have focused on specific lineages, such as a recent study by Pancreatic Cancer Action Network with the Know Your Tumor program finding that sequencing data from a cohort of over 600 pancreatic cancers with attempted profiling across 44 states showed a frequency of over 8% mutations in DDR genes (9). Although these data do not account for the possibility of the complete loss of the gene, they do support, similar to other large sequencing studies (10,11), that the DDR pathway is critically important in the tumorigenesis process for a number of tumor systems. More recently, Jonsson et al screened over 17,000 tumor profiles across 55 different types of tumors and found the prevalence of 2.7% for *BRCA1/2* pathologic germline variants (12). An additional small percentage had somatic mutations. Combing both germline pathogenic and somatic *BRCA1/2*

mutations accounted for 4.9% and covered at least 38 cancer types of which the majority were pancreatic, breast, ovary, and prostate cancers (12). Interestingly, only 61% of all *BRCA1/2* carriers with various cancers harbored a somatic LOH hit in the wild-type *BRCA1/2* allele, this number was significantly enriched over the background rate of LOH in tumors with non-pathogenic variants (20%) (12). These data support the notion that in some instances cancers arise in a setting of BRCA1/2 germline mutations independent of complete loss of BRCAness. Mouse modeling of these gene defects support and also refute some of these findings for the development of cancer. For instance, Venkitaraman and colleagues (13) generated a murine model of familial pancreatic cancer (driven by a *KRAS* G12D mutation) and found that germline heterozygosity for a functional *BRCA2* truncation induced pancreatic ductal adenocarcinomas (PDACs). Not following the two-hit paradigm, tumor cells from these animals did not lose a second *BRCA2* allele. Complementary to this work, in three out of four PDACs from patients who inherited a classic BRCA2999del5 mutation, did not obtain a LOH hit in the second allele (11). A more recent publication demonstrated two out of three PDX models from three glBRCA PDAC patients, obtained LOH in second allele (14). It may depend when the model systems are developed in relation to the progress of the patient's clinical course, since we have shown that over the course of the disease the patients can develop therapeutic resistance, thus changing these genomic events (15). Taken together, these studies: (I) point towards the need of a more comprehensive molecular signature for BRCAness (i.e., HRD scoring); (II) an understanding of how germline and somatic *BRCA1/2* mutations interact with different tumor systems and lineages; and (III) a caution to the field about completely relying both on the central dogma of tumor suppressors in regards to genome maintenance genes and germline testing.

### *Next generation germline/patient testing: the search for a comprehensive molecular signature*

Initially, identifying patients with a deficiency in the DDR pathway was solely based on Sanger Sequencing. The use of bench techniques to study DNA damage were, and still are, being explored as a simple test to find a defect in a pathway, without depicting the exact genetic lesion. For instance, by running a *FAND2* monoubiquitination assay (Western or immunofluorescence), a molecular lab could determine whether a mutation occurred in the

relevant Fanconi Anemia pathway, without knowing the exact gene that is mutated or loss in a cancer (16,17). The rapid development of molecular and –omic signatures have been established since the advent of sequencing, microarray, and proteomic technologies have gotten more advanced and facile. The repertoire and diversity of mutational signatures span from "one-off" publications (i.e., no follow-up or validating study) depicting the prognostic value of a sampling of gene expression levels for the prognostic value and predictive value of a specific therapy. These studies appeared prevalent in the 1990s and 2000s, as the ability to validate a specific focused gene signature in independent cohorts were rare, since largescale, multi-institutional registry and biobanking strategies were inefficient and rare. More recently, the availability of well-annotated samples, advanced technologies, and rigorously validated publicly available datasets has provided researchers with a reality check on their favorite or novel multi-panel gene signature (18) [for a recent comprehensive review see (19)].

As expected, due to the volume of cases and the amount of available resources (both clinical specimens and funding) the breast and ovarian cancer fields have led the way in developing a molecular signature for a HRD-score (20). Previous scores were solely based on: (I) allelic imbalance in regards to telomeres (21), (II) LOH (22), and (III) large scale genomic instability (23). Over the last few years, others have combined a combination of these markers with advanced technologies and insights to generate HRD scores with the ultimate goal to find a reliable assay that can determine whether a tumor genome is HRD, or even how immune-active these tumors may be (24,25), for the use as a predictive biomarker (14,20,26,27).

## *A therapeutic opportunity: an example of translating the science to the clinic*

As mentioned previously there are differing and overlapping definitions of HRD in cancer. Furthermore, there may be subtle but clinically meaningful differences of HRD signatures per cancer subtype (19). A genomic signature that defines HRD in ovarian cancer may not have the same clinical relevance in PDAC. For example, the Myriad HRD score, was not significantly associated with a higher response rate or prolonged survival in patients treated with FOLFIRINOX in a small retrospective study (28) (see *Table 1* list of HRD signatures). The actual prevalence of HRD in PDAC is estimated to be around 10–12% based on



**Table 1** Examples of homologous recombination deficiency (HRD) scores discovered and presented in the literature

whole genome sequencing data (25,30). However, additional efforts to further define subgroups of patients with platinum sensitivity are still evolving, and may expand this subgroup further. Based on the definition of HRD by genomic alterations in DDR pathways (identified from exome sequencing NGS), Pishvaian and colleagues demonstrated a superior overall survival benefit when these patients were exposed to platinum-based chemotherapy. In this study, the percentage of patients demonstrating genomic alterations in DDR pathways was approximately 16% (32).

### *Are all DNA damaging agents equally efficacious? And at what time point should these therapies be administered?*

It is important to differentiate between the different mechanisms of actions of the chemotherapeutic agents versus PARP inhibitors and additional, emerging targeted DDR drugs in development. These considerations may have a profound clinical impact, since a DDR-deficient tumor may show sensitivity to a DDR related chemotherapy, but not to a specific targeted DDR drug in development (e.g., PARP inhibitors and ATR inhibitors).

The platinum salts (carboplatinum, cisplatin and oxaliplatin), generate covalent cross-links between DNA bases from DNA-damage-inducing chemotherapies. The cytotoxic effects are determined by the relative amount and specific structure of DNA adducts (33). Alkylating agents (e.g., temozolomide) modify DNA bases. Electrophilic alkyl groups covalently bind to cellular nucleophilic sites, including bases in DNA, these interactions are responsible for cytotoxicity (34). Topoisomerases are essential for all organisms as they prevent DNA and RNA entanglements and resolve DNA supercoiling during replication and transcription. Inhibitors of topoisomerase 1 (camptothecin, topotecan and irinotecan) and topoisomerase 2 (etoposide and doxorubicin) generate TOP-DNA adducts and DNAstrand breaks. These drugs generate non-productive TOP-DNA cleavage complexes before re-ligation occurs (34). There are clear similarities and differences between the DNA-damage-inducing chemotherapies, irinotecan and platinum agents are standard of care treatments in PDAC and therefore the most explored in this setting.

A more specific approach to targeting the DDR pathway includes biological therapeutics specifically Poly (ADP-Ribose) Polymerase (PARP) inhibitors for tumors with defects in DNA repair. Tumors with compromised ability to repair double-strand breaks (DSB) by HR, are highly sensitive to blockage of the repair of DNA single-strand breaks (SSB), via the specific and targeted inhibition of PARP. PARP-inhibition causes failure of the repair SSB. This SSB encountered by the replication fork will cause stalling of the fork and therefore may result in fork collapse or the formation of DSB. In the absence of HR functional proteins (e.g., *BRCA 1/2*), the replication fork cannot be restarted and collapses, causing chromatid breaks. Additional PARP inhibition mechanisms include the "trapping" of PARP-1 protein on the site of DNA damage. This also may interfere with replication fork progression. This approach has demonstrated wide applicability in BRCA associated ovarian, breast, prostate and pancreatic cancer. Furthermore, initial efficacy has also been seen in the treatment of sporadic cancers with additional HR pathway impairments (35).

As mentioned, the most well described HRD biomarker in PDAC is germline *BRCA1/2* mutations. The global prevalence of germline *BRCA1/2* is around 7% (36). This subgroup of patients have shown a superior overall survival (OS) when treated with platinum based chemotherapy in retrospective studies (37). However, the toxicity profile of platinum treatment including the accumulating neuropathy and hematological toxicity is well described and needs to be considered here (38). PDAC associated with a germline *BRCA1/2* mutation demonstrate efficacy to platinum treatment. However, the side effects are debilitating and dose reductions or cessations are usually mandatory, thus limiting the profound therapeutic usefulness in BRCAassociated cancers. Therefore, additional maintenance strategies have been explored. The aim of a maintenance treatment is to provide an alternative treatment approach without compromising the patient's quality of life. The clinical trial design in maintenance studies, include comparison of drugs in the maintenance setting that have a potentially superior therapeutic window. For instance, the aim of the POLO study: Olaparib as Switch Maintenance Therapy after Response to platinum-based treatment of metastatic germline BRCA-mutant (gBRCAm) pancreatic cancer (36). Patients had to have received a minimum of 16 weeks platinum-based first line chemotherapy, and they had to demonstrate SD or PR or CR in order to be eligible for the clinical trial. Identified patients were randomized in a 2:1 ratio olaparib 300 mg twice daily or placebo. The primary endpoint PFS was 7.4 months on olaparib versus 3.8 months in the placebo arm, HR 0.53 (95% CI: 0.35–0.82; P=0.0038). Interim OS data (at 46% maturity) showed no difference between arms. Final OS results will be evaluated at 69% data maturity. No statistical differences



**Table 2** Opportunities and challenges of targeting DDR in cancer

DDR, DNA damage repair; LOH, loss of heterozygosity; HRD, homologous recombination deficiency; PDAC, pancreatic ductal adenocarcinomas; VUS, variants of unknown significance.

were noted in quality of life measurements between the olaparib versus placebo arm. Olaparib-arm patients were more likely to achieve a response to treatment or maintain disease control; responses were durable lasting a median of over 2 years. Of note, this strategic approach of first-line platinum-based chemotherapy followed by maintenance PARP inhibitor together has an extended PFS benefit to patients with a germline *BRCA1/2* mutations and metastatic disease. This study is the first Phase III trial to validate a targeted treatment in a biomarker-selected population of pancreatic cancer patients, highlighting the importance of gBRCAm testing in this setting.

### *Concluding thoughts and future directions*

Clearly both the medical oncology community and our patients diagnosed with cancer are eager to launch into the arena of personalized/precision oncology. In this era of facile genomic sequencing of tumor genomes, identifying defects in DDR genes or finding a relevant HRD score has provided a shiny glimmer of hope. The concept and the success of synthetic lethality for these tumors are real, as clinical trial data keeps emerging that supports this therapeutic strategy in numerous oncologic settings, including maintenance therapy. Simply put, this work is very promising and exciting for a disease like PDAC, where only 9% of patients live 5 years; we may offer a significant subset of patients an enhanced quality of life and overall survival.

Still there are numerous questions and challenges to

be worked out as we launch into this next generation of personalized medicine (*Table 2*). As outlined above, optimizing a rapid screening platform against pure tumor cells will be critical as we attempt to take advantage of a therapeutic window. However, based on recent data, the central dogma concept that two hits in a tumor suppressor or genome maintenance gene may have to be reconsidered in future and ongoing clinical trials in regards to selection criteria and retrospective analyses. Additional ramifications for these findings may not only affect the predictive value of identifying these mutations, but also have diagnostic implications for family members of patients who harbor the same germline mutations. Finally, additional genetic and beyond genomic alterations (e.g., the immune system and post-transcriptional gene regulation) in these tumor related to the DNA repair pathway are still being uncovered (25,39). These molecular lesions and gene regulatory mechanisms within tumor cells may provide novel targets, biomarkers, and important insights into innate and acquired resistance mechanisms for the above described therapies.

Certainly, a key factor in moving this field forward will be the prospective clinical and molecular analyses of ongoing clinical trials (*Figure 1*). The recent advances in the development and use of patient derived models of cancer (14,40,41) will provide an invaluable resource to study drug-target (gene) relationships, understanding resistance mechanisms, and may also complement personalized medicine approaches (*Figure 1***,** *Table 1*). Taken together, we are optimistic that validated multiplexed next generation platforms (that include patient derived models) (*Figure 1*) that can reliably



**Figure 1** A vision for a precision therapy strategy for the treatment of HR deficient tumors. HRD, homologous recombination deficiency.

identify DDR-deficient tumors along with optimized, targeted therapeutic strategies will be a game changer for many lethal cancers (e.g., PDAC) with a DDR-defect.

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