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Targeting Ataxia telangiectasia mutated (ATM) and DNA dependent protein kinase catalytic subunit (DNA-PKcs) for synthetic lethality application in breast cancer

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Abstract

BRCA1 germ-line mutations predispose to hereditary breast and ovarian cancers. Cells lacking functional BRCA1 protein are deficient in the homologous recombination (HR) DNA repair pathway. Base excision repair (BER) is essential for processing base damage induced by endogenous and exogenous sources. Recently, BRCA1 was shown to transcriptionally regulate expression of genes involved in BER such as OGG1, NTH1, APE1 and XRCC1. The primary aim of the work described in this thesis was to investigate whether targeting the double-strand break (DSB) pathway in BRCA1-BER deficient cells using ATM or DNA-PKcs inhibitors would be synthetically lethal.

Firstly, DNA repair gene and protein expression in BRCA1 deficient and BRCA1 proficient cells were investigated. Initially the RT² Profiler™ DNA Repair PCR Array was used to investigate the expression of 84 DNA repair genes. Data demonstrated down-regulation of several DNA repair mRNAs in BRCA1 mutant/knockdown cell lines as compared to BRCA1 proficient cell lines. Quantitative real time PCR (RT-qPCR) was performed for selected DNA repair genes (APE1, XRCC1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs) and confirmed statistically significant down-regulation of these genes in BRCA1 mutant/knockdown cell lines as compared to BRCA1 wild type/control cell lines. Protein expression of APE1, XRCC1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs was assessed by western blot analysis and showed down-regulation consistent with the results of mRNA expression. These results suggest that BRCA1 deficiency may be associated with a global defect in the BER pathway.
Secondly, as BRCA1 deficient cells demonstrated evidence of BER deficiency, these cells may be amenable to a synthetic lethality approach by targeting additional DSB repair targets. To examine this, BRCA1 deficient cells were targeted by ATM inhibitors (KU55933 and KU60019). Data demonstrated that BRCA1 mutant/knockdown cell lines were significantly more sensitive to ATM inhibition than BRCA1 proficient cell lines. Functional studies were conducted to understand the mechanism behind this selective cytotoxicity of ATM inhibitors in BRCA1 deficient cells. BRCA1 deficient cells treated with ATM inhibitors showed significant increase in DSB formation compared to untreated BRCA1 deficient cells. Cell cycle progression was arrested significantly at G2/M phase in treated BRCA1 HeLa cells as compared to untreated BRCA1 HeLa cells. On the other hand, MDA-MB-436 cells showed a modest increase in G2/M arrest after 48 hours treatment which suggests disturbed cell cycle progression. BRCA1 deficient cell lines also demonstrated increased levels for apoptosis in response to ATM inhibition. This in vitro study suggests that a potential synthetic lethality relationship exists between BRCA1 deficiency and ATM inhibition.

In light of the observed BRCA1/ATM synthetic lethality relationship, BRCA1 deficient cells were targeted with DNA-PKcs inhibitors (NU7441 and NU7026) to investigate whether these drugs would also induce synthetic lethality. Data demonstrated that BRCA1 mutant/knockdown cell lines were significantly more sensitive to DNA-PKcs inhibition than BRCA1 proficient cell lines. Functional studies showed that BRCA1
deficient cells treated with DNA-PKcs inhibitors have a significant increase in DSB formation as compared to untreated BRCA1 deficient cells. The formation of DSBs was followed by a modest arrest at G1 phase and induction of apoptosis. This *in vitro* study suggests that a potential synthetic lethality relationship exists between BRCA1 deficiency and DNA-PKcs inhibition.

Finally, BRCA1 deficient cells were targeted by ATM or DNA-PKcs inhibitor in combination with cisplatin to investigate whether it would enhance the observed synthetic lethality. The results of both combination treatments induced hypersensitivity in BRCA1 deficient cells compared to BRCA1 deficient cells treated with cisplatin alone. Significant increase in DSB formation was observed in BRCA1 mutant/knockdown cells treated with the combination of ATM inhibitor/cisplatin or DNA-PKcs inhibitor/cisplatin compared to each inhibitor alone. Cell cycle analysis showed significant G2/M arrest and induction of apoptosis in BRCA1 deficient cells treated with the combination treatments compared to each inhibitor alone. These results support the hypothesis that cisplatin increases the efficacy of ATM and DNA-PKcs inhibition in BRCA1 deficient cells. Taken together, this study provides the pre-clinical evidence that ATM and DNA-PKcs could be alternative synthetic lethality targets in BRCA1 deficient breast cancer.
Publication arising from this thesis


Poster presentations


Other related publications


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<tbody>
<tr>
<td>µg</td>
<td>Microgram</td>
</tr>
<tr>
<td>µl</td>
<td>Microliter</td>
</tr>
<tr>
<td>µM</td>
<td>Micromolar</td>
</tr>
<tr>
<td>AP</td>
<td>Apurinic/Apyrimidic</td>
</tr>
<tr>
<td>APE1</td>
<td>Human apurinic /apyrimidin endonuclease 1</td>
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<tr>
<td>ATM</td>
<td>Ataxia telangiectasia mutated</td>
</tr>
<tr>
<td>ATR</td>
<td>Ataxia-telangiectasia and Rad3-related</td>
</tr>
<tr>
<td>BER</td>
<td>Base excision repair</td>
</tr>
<tr>
<td>bp</td>
<td>Base pair</td>
</tr>
<tr>
<td>BRCA1</td>
<td>Breast cancer susceptibility 1</td>
</tr>
<tr>
<td>BRCA2</td>
<td>Breast cancer susceptibility 2</td>
</tr>
<tr>
<td>Ct</td>
<td>Threshold cycle</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl sulfoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>DNA-PK</td>
<td>DNA-protein kinase</td>
</tr>
<tr>
<td>DNA-PKcs</td>
<td>DNA dependent protein kinase catalytic subunit</td>
</tr>
<tr>
<td>DSB</td>
<td>Double strand break</td>
</tr>
<tr>
<td>FEN1</td>
<td>Flap structure specific endonuclease 1</td>
</tr>
<tr>
<td>H2AX</td>
<td>H2A histone family, member X</td>
</tr>
<tr>
<td>HR</td>
<td>Homologous recombination</td>
</tr>
<tr>
<td>IC₅₀</td>
<td>Inhibitory concentration of an inhibitor that results in death of 50% of cell population</td>
</tr>
<tr>
<td>kDa</td>
<td>Kilodalton</td>
</tr>
<tr>
<td>L</td>
<td>Litre</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>M</td>
<td>Molar</td>
</tr>
<tr>
<td>MMR</td>
<td>Mismatch Repair</td>
</tr>
<tr>
<td>MMS</td>
<td>Methyl methane sulphonate</td>
</tr>
<tr>
<td>MRN</td>
<td>MRE11/RAD50/NBS1 complex</td>
</tr>
<tr>
<td>NER</td>
<td>Nucleotide excision repair</td>
</tr>
<tr>
<td>NHEJ</td>
<td>Non homologous end joining</td>
</tr>
<tr>
<td>PARP1</td>
<td>Poly (ADP-Ribose) Polymerase 1</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffered saline</td>
</tr>
<tr>
<td>PI</td>
<td>Propidium iodide</td>
</tr>
<tr>
<td>PI3K</td>
<td>Phosphatidylinositol 3-kinase</td>
</tr>
<tr>
<td>Pol β</td>
<td>DNA polymerase beta</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>RNAse</td>
<td>Ribonuclease</td>
</tr>
<tr>
<td>rpm</td>
<td>rotation per minute</td>
</tr>
<tr>
<td>SD</td>
<td>Standard deviation</td>
</tr>
<tr>
<td>SEM</td>
<td>Standard error of the mean</td>
</tr>
<tr>
<td>SL</td>
<td>Synthetic lethality</td>
</tr>
<tr>
<td>SSB</td>
<td>Single strand DNA break</td>
</tr>
<tr>
<td>XRCC1</td>
<td>X-ray repair cross complementing gene 1</td>
</tr>
<tr>
<td>γH2AX</td>
<td>Phosphorylated H2AX</td>
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Chapter 1

Background
1. Introduction

Cancer is a class of complex diseases caused by genetic, epigenetic or translational changes that force the transformation of normal cells into malignant cells (Baylin and Jones, 2011). The major limitations of existing anticancer agents are that their efficacy is restricted by specificity, toxicity and the development of drug resistance. The search for new targeted, less toxic treatments and methods to improve the specificity and effectiveness of existing therapies is a major focus of cancer research.

In the early 1900s, Paul Ehrlich (German chemist) defined “Chemotherapy” as the use of chemicals to treat disease; he was also the first to use animal models successfully in screening a series of chemicals for their potential activity against diseases. Ehrlich's accomplishment remains the cornerstone in cancer drug development (DeVita and Chu, 2008). Since then cancer research has produced a complex and rich foundation of information which has revealed that cancer is a disease involving dynamic changes in the genome. To provide an overview of the complex biological data about cancer, researchers identify six essential features to transform normal cells into cancer cells. These features were referred to as “Hallmarks of cancer” by Hanahan and Weinberg (shown in Figure 1.1), and include; sustaining growth signals, insensitivity to antigrowth signals, resisting cell death, limitless replicative potential, inducing angiogenesis, activating invasion and metastasis (Hanahan and Weinberg, 2000).
Almost ten years after that and with an increasing body of research, Hanahan and Weinberg suggested two additional hallmarks and two enabling characteristics. The new emerging hallmarks are; deregulating cellular energetics and avoiding immune destruction, while tumor-promoting inflammation and genome instability and mutation are enabling characteristics (Hanahan and Weinberg, 2011).

![Therapeutic Targeting of the Hallmarks of Cancer](image)

**Figure 1.1. Therapeutic Targeting of the Hallmarks of Cancer.**

Each ‘characteristic’ can be targeted with a class of drug which is either in clinical trials or in some cases approved for clinical use in treating certain forms of human cancer. Adapted from (Hanahan and Weinberg, 2011). Copyright© 2011 by Elsevier with permission conveyed through Copyright Clearance Center Inc.

Genomic instability and mutation are described as “Enabling Characteristics” as cells develop new ways to use different DNA repair pathways and damage signalling to avoid cell death. One novel approach in the treatment of cancer is
to potentiate the effect of the chemotherapy drug in damaging the DNA by targeting DNA repair mechanisms responsible for repairing this damage. In other words, this field is interested in combining a DNA repair inhibitor with the right DNA damaging agent.

One such novel target is Poly (ADP-ribose) polymerase 1 (PARP1). PARP1 is an enzyme that plays an essential role in single strand break repair (SSBR), a pathway that is related to the base excision repair (BER) pathway. Inhibition of SSBR pathway is associated with accumulation of SSBs; without repair the SSB would be converted to a double strand break (DSB) lesion during DNA replication (Dantzer et al., 2000, Hoeijmakers, 2001, Schreiber et al., 2006).

In 2005, two Nature papers reported that cells deficient in the Breast Cancer Susceptibility genes BRCA1 and BRCA2 were 100 to 1000 fold more sensitive to PARP inhibition than BRCA heterozygote or wild type control cell lines (Bryant et al., 2005, Farmer et al., 2005). Both of these studies were reported independently by two groups using deficient models of BRCA1 and BRCA2 with different PARP inhibitors, which provide evidence that this sensitivity in BRCA mutant cells was due to PARP inhibition. The hypothesis was that with the mutation in BRCA genes cells are defective in the Homologous Recombination (HR) pathway and targeting PARP will inactivate the SSBR pathway inducing ‘Synthetic Lethality’.

‘Synthetic Lethality’ is the process by which cancer cells are selectively targeted by the inactivation of two or more genes or pathways that causes cell death, whereas inactivation in only one of them is not lethal (Kaelin, 2005, Ashworth, 2008). Since then several studies have been conducted to investigate this hypothesis. Now there is evidence that cells with defects in the
HR pathway, or where the \textit{BRCA} gene is silenced by epigenetic changes, are sensitive to PARP inhibitors (Ashworth, 2008), widening the therapeutic potential for these agents.

Chapter one of this thesis sets out to provide the background information on DNA damage and repair pathways in mammalian cells, where mechanism of both base excision repair (BER) and double strand break (DSB) repair will be studied in some detail. Following that the progress to date in the development of inhibitors for DSB repair, and the biological role of ATM and DNA-PKcs will be presented. Understanding the concept of “synthetic lethality” between BRCA and PARP in the treatment of cancers defective in HR DNA repair will follow. Finally, alternative synthetic lethality targets will be investigated.

\textbf{1.1.DNA damage}

DNA is under constant attack from various destructive agents including endogenous, as well as exogenous sources summarised in Figure 1.2. Endogenous damage may result from spontaneous base loss or various types of base modification (for instance; cytosine deamination, converting it to uracil) caused by exposure to metabolic products such as reactive oxygen species, or mispairing errors introduced during replication (Lindahl, 1993, Gates, 2009). Exogenous damage has many sources including; UV light, X-rays or gamma radiation, thermal disruption or chemical exposure. DNA damage types can be broadly subdivided into base damage and backbone damage (Hoeijmakers, 2001).
Cytosine residues in DNA can undergo hydrolytic deamination converting them to uracil residues. Up to 500 deamination of cytosine events take place in human cells per day. Other deamination reactions include conversion of adenine to hypoxanthine, 5-methylcytosine to thymine and guanine to xanthine. 

1.1.1. Base damage
Hydrolysis of the glycosidic bonds holding a base to the DNA backbone is also common (Gates, 2009).

**Oxidation**

One of the causes of DNA base damage is exposure to reactive oxygen species generated during either normal cellular oxygen metabolism, from exposure to UV light or a wide range of other exogenous sources. A frequent oxidative lesion is 8-hydroxyguanine (8-oxoG); this mutagenic lesion shows preference to pair with adenine rather than cytosine during replication. This lesion is estimated to occur at a rate of up to 500 events per day, the same rate as cytosine deamination (Tudek et al., 2003, Gates, 2009).

**Alkylation**

Another source of DNA base damage is where an alkyl group attaches to the DNA base giving alkylation products such as O²-alkylthymine, O⁴-alkylthymine, O⁶-methylguanine and O⁶-ethylguanine. This binding may prevent DNA replication causing mutation or cell death. Alkylation can be generated by both endogenous sources (for instance; oxidative by-product or cellular methyl donors such as S-adenosylmethionine) and exogenous sources (fuel combustion, tobacco exposure or anticancer therapies e.g. cisplatin) (Engelbergs et al., 2000, Fu et al., 2012).

**1.1.2. Backbone damage**

The DNA backbone is under constant exposure to environmental and endogenous agents that create thousands of lesions per cell each day (Lindahl,
1993). While some of these lesions like abasic sites or SSB are considered to be toxic, DSB are considered to be the most harmful. There are three major classes of DSB structures that can be toxic if not repaired.

(1) Two-ended DNA double-strand break, created by direct fracture of a DNA duplex. (2) One-ended DNA double-strand break, created when a replication fork encounters a DNA single-strand break. (3) Daughter strand gap, created when lagging or leading strand progression is inhibited by a DNA lesion (Helleday et al., 2007).

1.2. DNA repair

The accurate and efficient repair of DNA damage is essential for normal cellular function and maintenance of genomic stability (Figure 1.2) (Hoeijmakers, 2001). Genomic instability in cancer cells is often related to deficiency in one or more of the DNA repair pathways, which can account for the development of a more malignant phenotype (Hoeijmakers, 2009). Cells have two strategies to use for damage recognition and to initiate DNA damage responses. Direct recognition is the simplest form of repair (one step) where particular DNA damage can be repaired by cognate proteins (usually an enzyme). The second DNA damage response is multi-step recognition, where several proteins are involved and each one has a particular job to do, starting from recognition of the damage and ending with sealing the new DNA strand. DNA repair pathways can be divided into five categories; direct repair, base excision repair (BER), nucleotide excision repair (NER), mismatch repair (MMR) and double strand break repair (DSB) which includes, homologous
recombination (HR) and non homologous end joining (NHEJ) pathways (Sancar et al., 2004).

### 1.2.1. DNA repair pathways

#### 1.2.1.1. Direct repair

Several mechanisms exist to directly reverse certain DNA damaging lesions, for instance direct reversal of the O$^6$-methyl group from O$^6$-methylguanine (O$^6$-MeGua) by Methylguanine DNA Methyltransferase (MGMT). MGMT is a small enzyme of 20 kDa which forms a low stable complex with the DNA backbone at the damage site. It was believed that the enzyme flips-out the O$^6$-MeGua base and the methyl group is transferred to an active site cysteine. However, O$^6$-MeGua if not repaired can cause errors by mispairing with thymine during replication, causing G:C to A:T transitions or a strand break (Lindahl et al., 1988, Sancar et al., 2004, Luo et al., 2010).

#### 1.2.1.2. Base excision repair

BER is a complex pathway responsible for accurate removal of DNA bases that have been damaged by alkylation or oxidation, along with a variety of other lesions including base deamination and single strand breaks that could otherwise cause mutations during replication. BER in mammalian cells consists of two major sub-pathways that differ from each other in the length of the region they repair and in the subsets of enzymes involved (Figure 1.3). The short patch repair (SP-BER) pathway replaces single nucleotides, whereas the
long patch repair (LP-BER) pathway replaces 2-10 nucleotides. Initiation of BER is thought to be by the DNA glycosylase family proteins, which remove the damaged base creating an abasic site (apurinic/apyrimidinic, AP site). Apurinic/apyrimidinic endonuclease 1 (APE1) then cleaves the phosphodiester bond 5’ to the AP site leaving a nick with 5’deoxyribose-5-phosphate (5’dRP) and 3’-hydroxyl group (3’OH). The second step of BER is adding the first nucleotide to the 3’-end of the AP site and removing the 5’ terminal sugar residue (by the process of β-elimination) by DNA polymerase β (Pol β). Normally, this reaction continues through the SP-BER sub-pathway where DNA Ligase 3 and X-ray repair cross complementing gene 1 (XRCC1) complex then complete the repair.

On the other hand, when the 5’-end is resistant to β-elimination the repair process proceeds through the LP-BER sub-pathway. In this sub-pathway, replication factor C (RF-C) recruits proliferating cell nuclear antigen (PCNA) onto the AP site. PCNA acts as a DNA sliding clamp for the DNA polymerase δ and ε (Poly δ/ε) which performs DNA synthesis to remove the 5’ terminal sugar residue as part of a flap. As an alternative, Pol β and Rad9-Rad1-Hus1 complex (9-1-1) was found to have a similar structure as PCNA. The flap is then removed by flap endonuclease (FEN1) and DNA Ligase 1 completes LP-BER sub-pathway by ligating the DNA ends (Fortini and Dogliotti, 2007, Hegde et al., 2008, Abbotts and Madhusudan, 2010, Balakrishnan et al., 2009).

Poly(ADP-ribose) polymerases (PARPs) have been suggested to be involved in the regulation of the BER process, by binding to the AP site until the appropriate enzyme becomes available to process the repair (Khodyreva et
al., 2010, Yelamos et al., 2011). Once PARP1 is activated it catalyses the successive transfer of ADP-ribose units from the substrate nicotinamide adenine dinucleotide (NAD+) to a variety of acceptor nuclear proteins. This automodification attracts other DNA damage repair enzymes to the site such as, Pol β, Ligase 3 and XRCC1 (Schreiber et al., 2006, Megnin-Chanet et al., 2010). However, PARP1 does not play a major role in catalysis of DNA damage processing via either base excision repair pathway (Allinson et al., 2003, Strom et al., 2011).
Figure 1.3. Base Excision Repair (BER) pathway.

A damaged base is removed by a DNA glycosylase to generate an AP site. The repair patch may be a single nucleotide repair short patch (SP-BER) or a 2–10 nucleotide long patch (LP-BER). When the base damage is removed by one of the glycosylase family, APE1 (apurinic/apyrimidinic endonuclease 1) cleaves the 5’ bond to the site and recruits DNA polymerase (Pol β-δ-ε) to fill in the gap that is ligated by Ligase 3 (Lig3)/XRCC1 (X-ray cross-complementing protein 1) complex. When the AP site is generated by hydrolytic glycosylases, or by spontaneous hydrolysis, repair usually proceeds through the LP-BER sub-pathway. APE1 cleaves the 5’ phosphodiester bond, and the RF-C/PCNA-Poly complex carries out repair synthesis and nick translation, displacing several nucleotides. The flap structure is cleaved off by FEN1 (flap endonuclease) and the LP-BER is ligated by Ligase 1. PARP1/ poly(ADP-ribose) polymerase 1; PCNA/ proliferating cell nuclear antigen; OH/ hydroxyl group; dRP/ deoxyribose phosphate. Adapted from (Abbotts and Madhusudan, 2010).
1.2.1.3. Nucleotide excision repair

Nucleotide excision repair (NER) recognises and repairs base lesions associated with DNA helical distortion, including those produced by UV light or platinum chemotherapy. More than 25 proteins/complexes have been identified in eukaryotic cells which are required to complete NER. This pathway can be divided into two main sub pathways; global genome repair (GGR) and transcription-coupled repair (TCR), depending on the complexes that initiate repair. TCR removes lesions from the transcribed DNA strand of transcriptionally active genes when encountered by RNA polymerase II, restoring transcriptional activity and preventing apoptosis. GGR performs this process with poor efficiency, instead removing lesions on non-transcribed strands and transcriptionally inert genes to avoid replication fork stalling and chromosomal breakages. The key event in this pathway is incision of the damaged strand from each side of a lesion, releasing up to 32 nucleotides of the damaged DNA strand (Wood, 1997, Sugasawa, 2010).

The clinical consequences of defective NER are demonstrated by three syndromes associated with defects of NER proteins; xeroderma pigmentosum (XP), cockayne syndrome (CS) and the photosensitive form of trichothiodystrophy (TTD). Individuals with inherited XP syndrome exhibit severe sun-sensitivity, freckling and a 1000-fold increase in skin cancer including squamous and basal cell cancers as well as melanomas. Mutations in XP genes (XPA, XPB, XPC, XPD, XPE, XPF or XPG) can be identified in these patients. 20% of all XP patients, those with more than one mutation in XP genes, exhibit neurological abnormalities in addition to their XP symptoms. These symptoms include deafness, speach and walking disability
because of primary neuronal degradation. CS patients with mutation in CSA and CSB genes may exhibit sun-sensitivity and skin cancer though less than XP patients. CS symptoms include growth retardation, cachexia, neurological and mental developmental delays, cataracts, retinopathy, deafness, dental caries, and characteristic facies with a thin face, flat cheeks, and prominent tapering nose. TTD patients with mutations in TTD-A, XPB and XPD genes also show sun-sensitivity though no increased skin cancer risk. TTD symptoms are erythema, ichthyosis-like skin changes, nail and other neuroectodermal dysplasias, and short, brittle sulphur-deficient hair (Robbins et al., 1991, Thoms et al., 2007, Kraemer et al., 2007).

1.2.1.4. Mismatch repair

Mismatch repair (MMR) is the main pathway that recognises and repairs lesions/errors introduced during DNA replication. This pathway is initiated when a complex of MSH2 and MSH6 recognizes the mismatch. A range of combinations of MSH2 and either MSH6 or MSH3 are formed, which specify the type of mismatch recognized. After that, MSH proteins recruit MLH1 and its binding partners, post–meiotic-segregation increased 1 (PMS1) protein and post–meiotic-segregation increased 2 (PMS2) protein. An exonuclease removes the DNA lesion, a DNA polymerase synthesizes a new strand, and finally, a DNA Ligase completes the repair. A defect in MMR brings on a mutator phenotype, which causes a predisposition to cancer. MMR status also affects meiotic and mitotic recombination, DNA damage signalling and apoptosis (Jiricny, 2006, Li, 2008, Modrich and Lahue, 1996).
MMR deficiency is associated with three diseases, Hereditary Nonpolyposis Colorectal Cancer (HNPCC) syndrome, Muir – Torre syndrome, and Turcot syndrome. HNPCC patients often develop colon cancer and other visceral tumours such as endometrial, ovarian, stomach, kidney, and small intestinal cancers. In 80% of all HNPCC patients mutations in the \textit{hMLH1}, \textit{hMSH2}, \textit{hMSH6}, and \textit{hPMS2} MMR genes can be identified. Muir – Torre syndrome patients present with sebaceous carcinomas, colon cancer, and other visceral tumours such as endometrial, ovarian, stomach, kidney, and small intestinal cancers. Mutations in \textit{hMSH1} and \textit{hMSH2} genes can be identified in these patients. Turcot syndrome is caused by mutations in \textit{hMSH1} and \textit{hPMS2} genes. Turcot syndrome is associated with gliomas, lymphomas, colon cancer and other visceral tumours such as endometrial, ovarian, stomach, kidney, and small intestinal cancers (Peltomaki, 2003, Chung and Rustgi, 2003, Thoms et al., 2007).

1.2.1.5. Double strand break repair

DNA double strand breaks (DSBs) are the most lethal class of damage in DNA. DSBs, if left unrepaired or inappropriately repaired, can lead to cell death or to a variety of genetic mutations, translocations and chromosome loss (Hoeijmakers, 2001). This kind of lesion could be generated directly, following exposure to ionizing radiation, or indirectly during the processing of other DNA adducts or at stalled replication forks. DSB damage is repaired by one of the DSB repair sub-pathways illustrated in Figure 1.4, homologous recombination (HR) which is error-free or Non-homologous end joining (NHEJ) which is error-prone. The balance between these two sub pathways
differs between cell types of a single species and also differs between different cell cycle phases within a single cell type (O’Driscoll and Jeggo, 2006, Brandsma and Gent, 2012). In addition to classical HR and NHEJ there is an alternative DSB repair pathway:

Single Strand Annealing (SSA) pathway is activated when HR or NHEJ pathways cannot complete the repair. After resection of the break complementary stretches in the single strand DNA anneal, followed by the deletion of the intervening sequence and one of the repeat sequences. Some researchers categorize this pathway as an alternative for HR and others for NHEJ (Ivanov et al., 1996, Gottlich et al., 1998, Stark et al., 2004, Brandsma and Gent, 2012).

Microhomology-mediated end joining (MMEJ) is a subset of Alternative Non-homologous end joining (A-NHEJ). Repair by this pathway starts by resection of the 5’ – 3’ by MRN complex and then RPA binds to the DNA strand ends for protection. Where microhomologous regions are uncovered the DNA overhangs anneal, followed by DNA polymerase filling the gap and DNA Ligase sealing the strand. MMEJ is always associated with DNA deletions (McVey and Lee, 2008).

Alternative Non-homologous end joining (A-NHEJ) is an error prone pathway considered as a major source of genetic instability. Like SSA, A-NHEJ is thought to be activated when HR or NHEJ pathways cannot complete the repair. This pathway is not fully understood yet, however several proteins have been found to play a role in A-NHEJ including; PARP1, MRN complex, Ligase 3, Ligase 1 and XRCC1 (Brandsma and Gent, 2012, Dueva and Iliakis, 2013).
Homologous recombination (HR): (I) Following DSB, MRN complex and Rad51 localises to DSB via BRCA2. (II) Rad51 and related proteins assemble the pre-synaptic filament. (III) Rad51 and Rad54 identify homologous sequence of DNA and facilitate invasion into break site and new sequence of DNA is synthesised. (IV) Exchanged ends, (Holliday junctions), are rejoined by resolvases. Non-Homologous end joining (NHEJ): (V) Heterodimer of Ku70/Ku80 binds to exposed DNA initiating NHEJ. (VI) The active protein kinase DNA-PK complex (Ku70/Ku80 + DNA-PKcs) brings the ends of DSB for processing prior to religation. (VII) MRN complex, FEN1 and Artemis remove excess DNA ends. New DNA is synthesised by a DNA polymerase and ligation occurs via XRCC4 and DNA Ligase 4. DSB = double strand break, MRN = Mre11-Rad50-NBS1, XRCC2/3/4 = X-ray cross complementation group 2/3/4, V(D)J = Variable, diversity and joining, BRCA1/2 = breast cancer susceptibility 1/2. Adapted from Hoeijmakers, 2001, Copyright© 2001 by Nature Publishing Group with permission conveyed through Copyright Clearance Center Inc.
1.2.1.5.1. Homologous recombination

Homologous recombination (HR) pathway provides an error free repair of DSB. HR pathway takes place during late S phase and G2/M phase when homologous chromatids are available. HR involves multiple steps as presented in Figure 1.4, recognition of the DSB damage by the ataxia telangiectasia mutated (ATM) is the first sensor of DSB. ATM in turn activates multiple genes involved in HR repair, cell cycle arrest and apoptosis. Substrates for ATM phosphorylation include; BRCA1, p53, CHK2, histone H2AX and MRN complex (MRE11/RAD50/NBS1). The MRN complex then resects the DNA strand 5’ to 3’ generating 3’ ends of single stranded DNA (ssDNA) on either side of the DSB (Bakkenist and Kastan, 2004). The resulting SSB ends are then coated by Replication Protein A (RPA) protein which in turn recruits other proteins (including; Rad17/Rfc2-5, Rad9/Hus1/Rad1 and ATR/ATRIP) which form foci at the damaged site. RAD52 proteins then replace RPA proteins forming a ring to protect the SSB from exonucleolytic digestion. At this stage RAD52 competes with the KU complex of the NHEJ pathway and may determine sub-pathway selection between HR and NHEJ. RPA displacement is accomplished through the action of two proteins, BRCA1 and BRCA2. RAD51 and its related proteins (RAD51B, RAD51C, RAD51D, RAD54, XRCC2 and XRCC3) along with RPA, stimulate the activity of DNA strand exchange, with the complementary strand forming a heteroduplex while unpaired recipient DNA forms a D-loop. Conversely BRCA2 binds to RAD51, this complex binds to the exposed SSB and enables the loading of RAD51 to the SSB. Then DNA polymerases synthesize a new strand. The final step of the
HR repair is the sealing of the strands by DNA Ligase (Takata et al., 1998, Christmann et al., 2003, Li and Heyer, 2008).

1.2.1.5.2. Non-homologous end joining

The Non-homologous end joining (NHEJ) pathway is more commonly used for DSB repair by mammalian cells (Figure 1.4). NHEJ functions in all phases of cell cycle, especially in the G1 phase. It is an error-prone pathway as it does not rely on sister chromatids (homologous template) for repair. This results in short deletions or insertion of incorrect base pairs. After DSB induction, the first step towards repair through NHEJ is the binding of a heterodimeric complex consisting of the proteins KU70 and KU80 to the DNA damaged ends forming a ring shaped structure to protect the DNA from exonuclease digestion. Following DNA binding, the KU heterodimer recruits the catalytic subunit of the DNA dependent protein kinase (DNA-PKcs) thereby forming the active DNA-PK complex. DNA-PKcs is activated by interaction with a ssDNA at the site of DSB. In the following step, DNA-PKcs recruits XRCC4 which forms a stable complex with Ligase 4. The XRCC4/Ligase 4 complex binds to the ends of DNA molecules and brings both ends together with complementary but non-ligatable ends. This complex cannot directly re-ligate most of the DSBs before both ends are processed. Processing of DSBs is performed by the MRN complex, which has the role of removing the excess of DSB strands at the 3’ flaps. While flap endonuclease 1 (FEN1) is responsible for removal of the 5’ overhangs. Another protein involved in processing overhangs is the protein Artemis, which acts in a complex with DNA-PK.
Artemis acquires endonuclease activity, degrading single-strand overhangs and hairpins, which seems to be necessary for processing 5’ and 3’ overhangs during NHEJ. This pathway is sometimes associated with gain or loss of a few nucleotides if internal microhomologies are used for annealing before sealing. This implies the involvement of DNA polymerases or Artemis to create compatible ends. Finally, ligation is achieved by the actions of XRCC4/Ligase 4 complex (Takata et al., 1998, Christmann et al., 2003, Burma et al., 2006, Khanna and Jackson, 2001).

1.3. Ataxia telangiectasia mutated (ATM)

The *Ataxia Telangiectasia Mutated (ATM)* gene product was identified in 1995, based on its role in a rare, inherited disorder Ataxia Telangiectasia. The gene responsible for this disorder was mapped to chromosome 11q22-23. ATM protein is around 370 kDa and contains about 3056 amino acid residues. The 400 amino acids residues of its C-terminal region highly matched to the catalytic subunit of phosphatidylinositol-3 kinases (PI-3 kinases) (Savitsky et al., 1995, Rotman and Shiloh, 1998).

The catalytic site of ATM is within the C-terminal PI-3 domain, although it is believed that the rest of this large protein is responsible for receiving signals (working as a DNA damaging sensor) to regulate downstream targets. Several proteins interact with ATM including CHK2 and p53 (Rotman and Shiloh, 1998).
1.3.1. Role of ATM in the cellular response

ATM is a nuclear protein, the levels of which do not change when cells are exposed to ionizing radiation (IR). However, the protein kinase activity of ATM increases 2 to 3 fold following cellular exposure to IR, as determined by immunoprecipitation kinase assays (Canman et al., 1998, Banin et al., 1998, Chan et al., 1998). Bakkenist et al demonstrated that ATM containing a serine to alanine mutation at amino acid 1981 failed to support IR-induced phosphorylation of p53 or cell cycle arrest, suggesting that serine 1981 phosphorylation is critical for the function of ATM. Moreover, serine 1981-phosphorylated ATM localizes to nuclear foci in response to DNA damage (Bakkenist and Kastan, 2003).

The ATM signalling pathway is activated following DSBs. ATM is recruited to the damaged sites, however it is not clear if ATM interacts with damaged DNA directly or indirectly.

It was previously thought that ATM activation primarily requires direct binding to MRN complex. However recent studies has shown that PARP1 may control ATM phosphorylation and mediates the earliest recruitment of MRE11 and NBS1 to DNA damage (Rupnik et al., 2008, Haince et al., 2008, Haince et al., 2007).

Once ATM is activated, it may be released from the site of the DSB, enabling phosphorylation of several downstream substrates, such as chromatin bound histone H2AX and BRCA1 which lead to a variety of effects on DNA repair, cell cycle and apoptosis (Kurz and Lees-Miller, 2004). H2AX gets phosphorylated immediately after DNA damage by ATM on its C-terminal
serine residue 139 and is referred to as (γH2AX) forming foci, which can be observed by using immunofluorescence (Rogakou et al., 1999). In 2001 Burma et al showed that once ATM is activated at a DSB, it immediately activates γH2AX at the damaged site. This very early event could then initiate the recruitment of several DNA repair proteins (Burma et al., 2001). In terms of its role in cell cycle regulation, ATM interacts with a number of downstream substrates including proteins that activate the G1 (p53 and Nbs1), S (CHK2, BRCA1 and MRN) or G2/M (CHK1, CHK2 and Rad17) checkpoints. Activation of ATM-dependent cell cycle checkpoints would be expected to allow DNA repair, thereby enabling cells to survive. Studies proposed that considerable cross-talk exists between different ATM substrates and that several of them act in more than one cellular processes (Kurz and Lees-Miller, 2004).

Since ATM functions at the centre of a signalling network that controls different cellular responses, modulation of ATM function has become an area of interest for the treatment of cancer. Studies illustrated that ATM signalling pathways can be targeted in several ways to treat cancer. For instance, inhibition of ATM by the small molecule inhibitor (KU55933) increases the cytotoxicity of chemotherapeutic agents and IR in breast, colon and lung carcinoma cell lines (Crescenzi et al., 2008). Similarly, targeting p53 mutated cancer cells by inhibiting ATM increases cytotoxicity compared with proficient p53 cell lines (Smith et al., 2010).
1.4. DNA dependent protein kinase catalytic subunit (DNA-PKcs)

The DNA-dependent protein kinase (DNA-PK) holoenzyme consists of two subunits: the first component is the heterodimer complex KU, comprising of two enzymes KU70 (approximately 70 kDa) and KU80 (80 kDa). The second component of DNA-PK is a large polypeptide that corresponds to DNA-PK catalytic subunit (DNA-PKcs) (Hartley et al., 1995). The DNA-PKcs gene was mapped to chromosome 8q11 (Connelly et al., 1998). DNA-PKcs protein is around 460 kDa with approximately 3500 amino acid residues. The carboxy-terminal end of DNA-PKcs consists of 380 amino acid residues that fall into the phosphatidylinositol-3 kinase (PI-3 kinase) superfamily (Poltoratsky et al., 1995, Hartley et al., 1995, Kurimasa et al., 1999).

1.4.1. Role of DNA-PKcs in the cellular response

DNA-PKcs is a nuclear protein and recruitment to the DSB results in translocation of the heterodimer KU inward on the dsDNA allowing DNA-PKcs to directly interact with DSB ends (Yoo and Dynan, 1999, Calsou et al., 1999). DNA-PKcs was found to have unlimited kinase activity in the absence of KU heterodimer and DNA DSBs (Hammarsten and Chu, 1998, West et al., 1998). The kinase activity of DNA-PKcs is essential for NHEJ and the DNA damage response. DNA-PKcs can phosphorylate several nuclear proteins including KU heterodimer (Chan et al., 1999), XRCC4 (Leber et al., 1998), Ligase 4 (Wang et al., 2004), Artemis (Weterings and Chen, 2008), H2AX (Stiff et al., 2004) and p53 (Shieh et al., 1997). Recently it was found that
DNA-PKcs phosphorylates NHEJ Werner syndrome protein (WRN) that is required for efficient DSB repair (Kusumoto-Matsuo et al., 2014).

Following induction of a DSB, DNA-PKcs is phosphorylated at more than 40 sites. The best characterized DNA-PKcs phosphorylation cluster is the threonine 2609. Phosphorylation of threonine 2609 cluster is important for NHEJ. Recently Zhang et al demonstrated that knock-in mutant mice with human DNA-PKcs lacking a functional threonine 2609 are highly sensitive to replication stress agents which confer early lethality. In addition, the threonine 2609 phosphorylation cluster has been found to associate with other DNA repair proteins. For example, DNA-PKcs phosphorylation at the threonine 2609 is important for the coordination between ATM and Artemis in DNA DSB repair (Zhang et al., 2011, Davis et al., 2014a).

Regarding DNA-PKcs phosphorylations role in DNA repair, it is proposed that DNA-PK holoenzyme functions as a scaffolding protein to bring the broken DNA ends together and assists in the localisation of repair factors (Davis et al., 2014a). Moreover, DNA-PKcs phosphorylation has a role in regulation of cell cycle arrest in response to DSB. Chen et al.’s illustrated that DNA-PKcs functions primarily in the G1 phase (Chen et al., 2005). Conversely, DNA-PKcs plays a direct role in the suppression of apoptosis. Le Romancer et al demonstrated that cleavage and inactivation of DNA-PK during apoptosis is expected to switch off the function of kinase in DNA repair. In mammalian cells, loss of DNA-PK due to mutations increases sensitivity to treatments that induce DSBs (Le Romancer et al., 1996).
1.5. The development of ATM and DNA-PKcs inhibitors

To date, several DSB repair inhibitors have been described. Wortmannin is a non-specific inhibitor of PI-3 kinase. It is known to block ATM as well as DNA-PKcs (Rosenzweig et al., 1997). In 2004, the first small molecule to target ATM kinase KU55933 (2-morpholin-4-yl-6-thianthren-1-yl-pyran-4-one) was developed. KU55933 potentiates the cytotoxicity of IR and chemotherapeutic agents in cell lines (Hickson et al., 2004). In 2008 CP466722, a specific inhibitor of ATM [2-(6, 7-dimethoxyquinazolin-4-yl)-5-(pyridin-2-yl)-2H-1, 2, 4-triazol-3-amine], was identified. CP466722 inhibited ATM dependent phosphorylation events and resulted in G2/M cell cycle arrest. CP466722 shows similarity to KU55933 inhibitor in rapidly and potently inhibiting ATM over a period of several hours and demonstrates reasonable stability in tissue culture (Rainey et al., 2008). In 2009, KU60019 (2-[(2R, 6S)-2, 6-dimethylmorpholin-4-yl]-N-[5-(6-morpholin-4-yl-4-oxo-4H-pyran-2-yl)-9H-thioxanthen-2-yl]-acetamide), another specific inhibitor of ATM kinase was developed. KU60019 is an improved analogue of KU55933, and has been shown to be around 10 times more potent than KU55933 at radiosensitizing human glioma cells (Golding et al., 2009).

After initial use of Wortmannin for DNA-PKcs inhibition, LY294002 (2-(4-morpholinyl)-8-phenylchromone) was developed. LY294002 is non-specific DNA-PKcs inhibitor and more toxic than Wortmannin (Vlahos et al., 1994, Davidson et al., 2013). A number of inhibitors have been generated using LY294002 as a template. NU7026 (2- (morpholin-4-yl)-benzo [h]chomen-4-one), is over 50 fold more selective for DNA-PKcs than to other PI-3Ks.
including ATM and ATR. NU7026 alone had no effect on cell cycle distribution although it increases DSB levels (Willmore et al., 2004). **NU7441** inhibitor (2-N-morpholino-8-dibenzothiophenyl-chromen-4-one) is more potent and specific to DNA-PKcs, (IC$_{50}$ of only 14 nmol/L) and has at least 100 fold selectivity for DNA-PKcs compared with other PI-3K family kinases. NU7441 inhibitor has proved to be not only a potent chemo- and radio-sensitiser but also a powerful tool to study the biology of DNA-PKcs (Leahy et al., 2004, Zhao et al., 2006, Tavecchio et al., 2012, Davidson et al., 2013).

### 1.6. Breast Cancer

Every day, thousands of women around the world are diagnosed with breast cancer. In 2008 almost 1.4 million women were diagnosed with breast cancer worldwide and around 459,000 deaths were recorded. Incidence rates were much higher in more developed countries with 71.7 per 100,000 compared to 29.3 per 100,000 in less developed countries, whereas the corresponding mortality rates were 17.1 per 100,000 and 11.8 per 100,000 respectively. The highest incidence rates were recorded in Western Europe, Australia/New Zealand and Northern Europe, whilst rates were lowest in Eastern Africa and Middle Africa (Youlden et al., 2012). There is a 0.5% overall increase in incidence annually. However, observed improvements in breast cancer survival especially in developed countries over recent decades have been attributed to the introduction of population-based screening using mammography, improved education and the systemic use of adjuvant therapies (Rosso et al., 2010, Gunsoy et al., 2014).
There are a number of factors that contribute to an increase risk of breast cancer, such as:

(1) Age, about 2 out of 3 invasive breast cancers are found in women over the age of 50. (2) Diet, there is an increase in breast cancer risk by 13% with the higher level of fat intake. (3) Tobacco exposure in all its forms (smoking, and including second-hand smoke). (4) Alcohol consumption, alcohol has been described as being one of the most constant enhancers of breast cancer risk. (5) Body size, obesity if associated with an increased risk of breast cancer (Youlden et al., 2012, McCormack and Boffetta, 2011, Gunsoy et al., 2014). (6) Family history, having a first degree relative with breast cancer can increase a woman’s risk by two fold. This risk is greater if the relative has been diagnosed before the age of 50 (Pharoah et al., 1997). Family history accounts for 5-10% of the total number of cases of breast cancer and often involves germline mutations in breast cancer susceptibility genes such as BRCA1 and BRCA2, or others like p53, PTEN, ATM, CHK2 and RAD51 (Lose et al., 2006, Flanagan et al., 2010).

A wide variety of clinical and pathological factors are regularly used to categorize breast cancer patients in order to assess prognosis and determine the appropriate therapy. These factors include;

(1) Age at diagnosis, women under the age of 40 often have a more aggressive cancer with higher mortality and recurrence rates in comparison with older women. Younger patients are more likely to have poor clinical characteristics, e.g. larger tumour size and higher histological grade, lymphatic vessel invasion and the involvement of lymph nodes. This is why this group is classified as
high risk and approximately 80% of these patients are offered adjuvant cytotoxic treatment (Kroman et al., 2000, van der Sangen et al., 2008, Schnitt, 2010). (2) Tumour size, it has been established that the rate of distant recurrence increases in patients with large tumours as opposed to smaller tumours (Koscielny et al., 2009). However, tumour size alone is an unreliable prognostic factor in breast cancer (Foulkes et al., 2009). (3) Lymph node status, it is thought to be the second most prominent factor in gauging the probability of survival, after tumour size (Vorgias et al., 2001). (4) Hormone receptor expression, estrogen receptors (ER) and progesterone receptors (PR) located within the cell nucleus have prognostic value in breast cancer patients (Fisher et al., 1986, Garicochea et al., 2009). The prognostic and predictive importance of assessment of ER expression in breast cancer is well established; e.g. patients with ER-positive tumours have more favourable prognostic characteristics than patients with ER-negative tumours which relapse earlier. Patients with ER positive breast cancer will usually receive hormonal therapy; however the added value of PR assessment is controversial. PR status in ER- positive breast cancer might be used to help guide clinical management, as high levels of PR expression may identify a subset of ER-positive patients most likely to benefit from hormonal therapy (Hefti et al., 2013). (5) Histological type, the histological type of invasive breast cancer presents valuable prognostic information. A few histologic types of invasive carcinoma have better prognostic value than others. Ductal carcinoma of no special type, accounting for up to 70% of all breast cancers, has the worst prognosis, whereas tubular breast cancer demonstrates the best prognosis (Martinez and Azzopardi, 1979, Fisher et al., 1975, Ellis et al., 1992, Rakha et
Molecular classification, gene expression profiling has been used to define breast cancer subtypes based on the molecular characteristics and its association with clinical outcome. This classification groups tumours into five molecular subtypes; (a) Luminal A (breast cancer with the highest expression of ER receptor and expression of keratin 18 and 19 in luminal mammary cells, this class is the most common subtype, representing 50–60% of the total). (b) Luminal B (similar to luminal A but has moderate expression of ER receptor, luminal B molecular profile makes up between 10–20% of all breast cancers). (c) Basal like (ER-negative and high expression of cytokeratin 5, cytokeratin 17, the basal-like subtype represents 10–20% of all breast carcinomas). (d) Normal breast-like (expression of genes known to be expressed in normal mammary tissue, account for about 5–10% of all breast carcinomas). (e) HER2 positive (high expression of the HER2 gene and other genes associated with the HER2 pathway and/or HER2 amplicon located in the 17q12 chromosome, 15-20% of all breast cancers correspond to this molecular subtype) (West et al., 2001, Eroles et al., 2012).

1.7. The BRCA1 tumour suppressor gene

The Breast Cancer susceptibility 1 (BRCA1) gene was identified by Mary King’s group in 1990; this gene was mapped to chromosome 17q21 then cloned by Mark Skolnick's group in 1994. BRCA1 gene contains 24 exons, 22 coding exons and 2 non-coding exons. The identification of BRCA1 gene was a significant breakthrough in the management of breast and ovarian cancers (Hall et al., 1990, Miki et al., 1994).
BRCA1 protein is around 220 kDa and is a highly phosphorylated nuclear protein. BRCA1 contains nuclear import and export signals, allowing movement between the nucleus and cytoplasm to enable BRCA1 to perform its functions. BRCA1 contains about 1863 amino acid residues (Figure 1.5) including an N-terminal RING domain, two nuclear localization signals (NLSs), and two C-terminal BRCA1 Carboxyl Terminal (BRCT) domains of around 110 amino acid residues (Miki et al., 1994, Venkitaraman, 2002, Silver and Livingston, 2012). These domains interact with several proteins and contribute to multiple functions of BRCA1, including in DNA damage response and repair, cell cycle regulation, and transcriptional regulation.

![Figure 1.5. The Structure Domain of BRCA1.](image)

Two domains of BRCA1 that are frequently mutated in families with hereditary breast and ovarian cancer are the N-terminal RING domain and the C-terminal BRCT repeats. Both could contribute to tumour suppressing functions of BRCA1. NLS = nuclear localisation signal, BRCT = BRCA1 carboxyl terminal. Adapted from (Silver and Livingston, 2012). Copyright© 2012 by American Association for Cancer Research with permission conveyed through Copyright Clearance Center Inc.
1.7.1. Role of BRCA1 in DNA damage response and repair

One of the first links between BRCA1 and DNA repair was the functional interaction between BRCA1 and RAD51 suggesting a role for BRCA1 in genomic instability (Scully et al., 1997). Since then many studies have shown that BRCA1 plays a vital role in the HR pathway (Moynahan et al., 1999). In addition BRCA1 is also involved in the initial detection of DNA damage. Moreover, BRCA1 has roles in other DNA repair pathways such as:

(1) **NER**: BRCA1 has shown direct control of global genomic repair (GGR) a sub-pathway of NER pathway through \textit{XPC}, \textit{DDB2} and \textit{GADD45} genes, suggesting that deficiency in BRCA1 is followed by deficiency in GGR/NER (Hartman and Ford, 2002).

(2) **NHEJ**: the role of BRCA1 in the NHEJ pathway is uncertain, although several studies suggest a major role of BRCA1 in NHEJ and the maintenance of genomic integrity (Zhong et al., 2002, Bau et al., 2004), while others did not (Moynahan et al., 1999, Merel et al., 2002).

(3) BRCA1 has been suggested to stimulate the activity of three BER enzymes; OGG1 (8-oxoguanine DNA glycosylase), NTH1 (homolog of endonuclease III), and APE1. These enzymes mediate repair of three signature oxidative lesions, 8-oxoguanine, thymine glycol and abasic sites, respectively.

BRCA1 regulation of BER enzymes was found to be through a transcriptional mechanism involving the transcription factor OCT1 (Octamer-binding transcription factor 1). The exact mechanism by which the BRCA1/OCT1 complex regulates the BER pathway, and whether or not there is a physiologically important defect in BER that can be therapeutically targeted in
BRCA1 deficient cells, remains to be answered (Saha et al., 2010). A previous study has shown that BRCA1 in collaboration with APE1 down regulates levels of reactive oxygen species, oxidized DNA and nitrated proteins (Saha et al., 2009).

1.7.2. Role of BRCA1 in cell cycle regulation

BRCA1 is involved in cell cycle regulation, a function that may allow adequate time for DNA repair to occur in cells. The phosphorylation of BRCA1 is required to activate cell cycle arrest after DNA damage. For instance, in response to ionizing irradiation, ATM phosphorylates BRCA1 on serine 1387 which is required to activate S phase arrest. On the other hand, BRCA1 phosphorylation on serine 1423 has a direct role in G2/M cell cycle arrest (Xu et al., 2002). Cell cycle regulation is controlled by a family of cyclin dependent kinases and their endogenous inhibitors acting at a variety of cell cycle checkpoints. For example, BRCA1 phosphorylation has been shown, through p53 dependent and independent pathways, to stimulate the transcription of the p21\textsuperscript{CIPI/WAF1} leading to G1/S cycle arrest (Chai et al., 1999, Mullan et al., 2006). BRCA1 participates in the G1/S checkpoint response indirectly through CHK2 and p53 phosphorylation (Foray et al., 2003). Recently, DNA-PKcs was shown to activate the CHK2–BRCA1 pathway during mitosis to ensure chromosomal stability (Shang et al., 2014).
1.7.3. Role of BRCA1 in transcriptional regulation

BRCA1 may regulate transcription by interacting with a core component of transcription RNA called polymerase II, BRCA1 is also a co-activator or co-repressor of a number of known transcription factors (Irminger-Finger et al., 1999). For example, BRCA1 has been reported to regulate the oestrogen receptor alpha (ERα). The ability of BRCA1 to induce expression of ERα is dependent on the transcription factor OCT1, which is required to recruit BRCA1 to the ERα promoter (Gorski et al., 2009). In addition, BRCA1 has been shown to interact with the C-terminus of p53 and alter the transcriptional activity of p53 by re-directing it from activation of pro-apoptotic target genes to other genes involved in cell cycle arrest and DNA repair (Chai et al., 1999). Interestingly, mutations in p53 are common in BRCA mutated cancers and that this loss of p53 function may be important in tumourgenesis (Mullan et al., 2006).

1.8. BRCA1 and cancer

Germ-line mutations in BRCA1 predispose to breast and ovarian cancer development. BRCA1 mutation carriers have an 80% lifetime risk of developing breast cancer and 40-50% risk of developing ovarian cancer (Risch et al., 2001). In addition, BRCA1 mutations increase the risk of other types of cancer, such as pancreatic and prostate cancers. Children (both male and female) of a carrier have a 50% chance of inheriting the mutation in their germline, where one defective allele is sufficient to predispose to cancer (Thompson and Easton, 2002). If there is somatic loss of function of the
second allele then a tumour is likely to develop. This is known as the Knudson two hit hypothesis for tumour suppressor genes (Knudson, 1971).

1.9. Synthetic lethality

Synthetic lethality is defined as a genetic combination of mutations in two or more genes/pathways that leads to cell death, whereas a mutation in only one of them does not (Figure 1.6). Synthetic is used here for its ancient Greek meaning: the combination of two entities to form something new. This phenomenon was first discovered from the investigation of fruit flies in 1922 by Calvin Bridge. Sturtevant replicated these results in *Drosophila pseudoobscura* in 1956 (Rehman et al., 2010, Kaelin, 2005, Nijman, 2011). Synthetic lethality represents a new approach of targeting DNA repair pathways with cancer therapies based on the genetic background of the tumour.

![Diagram of Synthetic Lethality](image)

**Figure 1.6. Synthetic Lethality.**

Deletion of either gene A or gene B does not affect cell viability whereas inactivation of both at the same time is synthetically lethal. Adapted from (Nijman, 2011).
1.9.1. Synthetic lethality between BRCA and PARP

The well known synthetic lethality relationship to date is targeting BRCA mutation with PARP1 inhibitors (Farmer et al., 2005, Bryant et al., 2005). PARP1 plays a role in single strand break repair (SSBR), a pathway related to BER. Inhibition of SSBR is associated with accumulation of DSBs, which can be exploited in a subset of cancers possessing defects in DSB repair. BRCA gene products have a role in the HR pathway (Venkitaraman, 2002). BRCA deficient cells have impaired HR repair (Moynahan et al., 1999). The PARP1/BRCA synthetic lethality model proposes that targeting BRCA deficient tumours, which are HR deficient, with PARP inhibitor leads to accumulation of SSBs. This is followed by the formation of DSBs during DNA replication, leading to cell death (Farmer et al., 2005, Bryant et al., 2005). A study by Farmer et al demonstrated reduced survival of BRCA1 and BRCA2 deficient cell lines after treatment with PARP1 inhibitors (KU0058684 and KU0058948). Both treated BRCA deficient cell lines showed G2/M cell cycle arrest and apoptosis. In addition, Bryant et al, demonstrated that PARP1 inhibitors (NU1025 and AG14361) were more effective in BRCA2 deficient cell lines as compared to BRCA2 proficient cell lines. Treatment of BRCA2 deficient cell lines resulted in induction of γH2AX foci formation (representing DNA DSBs). Both of these studies were reported independently and concluded that BRCA deficient cells were selectively sensitive to PARP1 inhibition.

The PARP1/BRCA synthetic lethality model was then translated into clinical trials. A Phase I clinical trial of PARP inhibitor was conducted with Olaparib (known previously as AZD2281 and then KU0059436) in a cohort of patients
with BRCA deficient tumours. The study demonstrated that Olaparib has antitumor activity in BRCA mutation carriers. In addition, a Phase II clinical trial of Olaparib in BRCA deficient tumours showed positive proof of the PARP1/BRCA synthetic lethality concept (Fong et al., 2010, Balmana et al., 2011, Audeh et al., 2010, Tutt et al., 2010). The outcomes of both Phase I and II clinical trials provided evidence that PARP inhibitors could be used as a new treatment in BRCA deficient tumours (Chan and Mok, 2010, Balmana et al., 2011).

Recent studies have, however, illustrated some problems with PARP inhibition, including a lack of specificity. The PARP family of enzymes consists of at least 17 members and each one of them has a different structure and function (Rouleau et al., 2010, Mangerich and Burkle, 2011, Underhill et al., 2011). PARP1 for instance is involved in: telomere length, organizing the spindle apparatus, localization and activation of p53 and also the SSBR pathway (Mangerich and Burkle, 2011). Recent studies have evaluated a series of 185 inhibitors of PARP, including the best-known inhibitors being tested clinically such as Olaparib, ABT-888 and Rucaparib, for the specificity to bind to the catalytic domains of 13 of the 17 human PARP family members. In this study they determined that majority of the inhibitors bind to multiple targets and are therefore not specific (Wahlberg et al., 2012). Resistance to PARP inhibition is also an emerging problem. Potential resistance mechanisms include: altered NHEJ capacity and restoration of, or increase in, HR capacity (Montoni et al., 2013). Moreover, although BRCA1 deficient tumours are sensitive to PARP1, BRCA1 hypomorphic mutant tumours can restore BRCA1
expression and are therefore insensitive to PARP inhibition (Drost et al., 2011). Another potential resistance mechanism involves Tumor Suppressor p53 Binding Protein 1 (53BP1), which binds to the central DNA-binding domain of p53. Bunting et al., demonstrated that loss of 53BP1 can restore the HR pathway in BRCA1 deficient cells. The mechanism for this is the promotion of ATM function and therefore this may be overcome by the use of ATM inhibitors (Bunting et al., 2010). On the other hand, because most agents that would be combined with a PARP inhibitor are already administered at or near a maximum dose, the addition of another potentiating agent is likely to increase toxicity as well as efficacy. For instance, a phase II trial was conducted involving 40 patients with metastatic malignant melanoma treated with PARP inhibitor (AG014699) in combination with temozolomide. Results showed that 18% of the study subjects demonstrated partial responses, with notable side effects such as, temozolomide-related myelosuppression and one toxic death (Plummer et al., 2006, Pacher and Szabo, 2007). More recently a phase I trial was terminated; this study was designed to determine the maximum tolerated dose and safety profile of Olaparib in combination with topotecan in solid tumours. The most common side effects reported were fatigue and gastrointestinal adverse events similar to the ones previously reported in Olaparib monotherapy studies. However, unexpected high haematological toxicities were observed even with substantially lower doses of both agents (Samol et al., 2012).
1.9.2. Alternative synthetic lethality targets

The discovery of the synthetic lethality (SL) relationship between BRCA and PARP has become the paradigm for a new class of rational cancer therapies. Several novel SL candidates have also been recently investigated. For instance, RAD51D deficiency showed sensitivity to PARP inhibitor suggesting a possible SL target (Loveday et al., 2011). Similarly, Mantle Cell Lymphoma (MCL) deficient in both ATM and p53 are more sensitive to the PARP inhibitor (Olaparib) than cells lacking ATM function alone. Suggesting that inhibition of both ATM and PARP may have efficacy in targeting p53 deficient malignancies (Williamson et al., 2012). Similar sensitivity is observed in MRE11 deficient cells with PARP1 inhibitor (ABT-888) treatment (Vilar et al., 2011).

The emerging problems with PARP inhibition led our group to investigate alternative SL targets. For example, we have shown that targeting Apurinic/apyrimidinic endonuclease (APE1), a central component of the BER pathway, with a small molecule inhibitor induces SL in HR deficient cell lines (Mohammed et al., 2011, Sultana et al., 2012). Similarly, targeting APE1 in Phosphatase and tensin homolog (PTEN) deficient melanoma cell lines induces SL (Abbotts et al., 2014a). In addition, targeting X-ray repair cross complementing gene 1 (XRCC1) deficient cell lines with DSB repair inhibitors (ATM or DNA-PKcs) or SSB break repair inhibitor (ATR) results in SL (Sultana et al., 2013a, Sultana et al., 2013b).
1.10. **Hypothesis and Aims**

The base excision repair (BER) pathway is critical for processing DNA damage caused by alkylation, oxidation, single strand breaks and base deamination (Dianov and Hubscher, 2013). The well known synthetic lethality relationship to date is targeting *BRCA* mutated tumours (HR deficient) with PARP1 inhibitors. However with the emerging problems using PARP inhibition, several novel synthetic lethality candidates have been recently investigated within the BER pathway. On the other hand, studies suggest a cross talk between BRCA1 and BER (Alli et al., 2009). In a more recent study, a functional interaction between Pol β and BRCA1 was demonstrated (Masaoka et al., 2013), implying a potential role for Pol β in BRCA1 mediated DSB repair. In addition, BRCA1 has also been shown to be involved in the transcriptional regulation of BER factor such as OGG1, NTH1 and APE1 (Saha et al., 2010). As a result, the hypothesis of this study is that with the essential role of BRCA1 in regulating several BER factors, such as APE1, Pol β, OGG1 and NTH1, then BRCA1 deficient cell lines may have impaired BER factors which leads to accumulation of SSBs followed by the formation of DSBs during the DNA replication process. Therefore, it is suggested that BRCA1 deficient cells are reliant on the DSB pathway for cellular survival. Accordingly targeting the DSB pathway by ATM or DNA-PKcs inhibition in BRCA1-BER deficient cell lines may induce synthetic lethality (Figure 1.7).

The aims of this research project are as follows:

1. To investigate DNA repair mRNA and protein expression in BRCA1 deficient and proficient cell lines.
2. To investigate if targeting BRCA1 deficiency by ATM inhibitors (KU55933 & KU60019) is synthetically lethal.

3. To investigate if targeting BRCA1 deficiency by DNA-PKcs inhibitors (NU7441 & NU7026) is synthetically lethal.

4. To investigate if targeting BRCA1 deficiency by ATM or DNA-PKcs inhibitor in combination with cisplatin is synthetically lethal.

**Figure 1.7. Hypothetical synthetic lethality in BRCA1/ BER deficient cells, by ATM or DNA-PKcs inhibitors.**

Impaired BER factors in BRCA1 deficient cell lines leads to accumulation of SSBs followed by the formation of DSBs during the DNA replication process. Therefore, it is suggested that cells are reliant on the DSB pathway for cellular survival. Targeting the DSB pathway by ATM or DNA-PKcs inhibition may induce synthetic lethality leading to cell death.
Chapter 2

Materials and Methods
2. Materials and Methods

2.1. Materials

KU55933 [(2R,6S-rel)-2,6-Dimethyl-N-[5-(4-morpholinyl)-4-oxo-4H-pyran-2-yl]-9H-thioxanthen-2-yl-4-morpholineacetamide], KU60019 [(2R,6S-rel)-2,6-Dimethyl-N-[5-(4-morpholinyl)-4-oxo-4H-pyran-2-yl]-9H-thioxanthen-2-yl-4-morpholineacetamide], NU7441 [8-(4-Dibenzothienyl)-2-(4-morpholinyl)-4H-1-benzopyran-4-one], NU7026 [2-(4-Morpholinyl)-4H-naphthol[1,2-b]pyran-4-one], and NU1025 [8-Hydroxy-2-methyl-4(3H)-quinazolinone] were purchased from Tocris Bioscience (UK), while (3-AB) 3-aminobenzamide was purchased from Sigma (UK). All compounds were dissolved in 100 % v/v dimethyl sulphoxide (DMSO) to give 5 or 10 mM stock solutions, and stored at -20°C. Cisplatin solution was obtained from Pharmacy, Nottingham City Hospital, UK and stored at room temperature.

2.2. Cell lines and culture media

2.2.1. HeLa SilenciX® cells

Human cervical adenocarcinoma adherent cell lines, BRCA1 HeLa SilenciX (Adherent HeLa cells silenced for BRCA1 (Accession Number: NM_007295)) and Control HeLa SilenciX (Adherent HeLa cells transfect with control shRNA), were purchased from Tebu-Bio (www.tebu-bio.com) and were grown in DMEM medium (with L-Glutamine 580mg/L, 4500 mg/L D-Glucose, with 110mg/L Sodium Pyruvate) (Invitrogen, UK) supplemented with 10% FBS, 1% penicillin/streptomycin and 125 μg/ml Hygromycin B (Invitrogen, UK). BRCA1 HeLa SilenciX cells were used between passage 20
and 34; Control HeLa SilenciX cells were used between passage 48 and 60. All materials were obtained from (Sigma/PAA, UK) unless otherwise stated. To make it easier for discussion of the results, BRCA1 deficient HeLa SilenciX cells are referred to as **BRCA1 HeLa** and Control BRCA1 proficient HeLa SilenciX cells are referred to as **Control HeLa**.

### 2.2.2. Breast cancer cells

Human breast adenocarcinoma adherent cell lines, MCF7 (hemizygous for BRCA1 wild type used between passage 25 and 38), and MDA-MB-436 (*BRCA1* mutated used between passage 17 and 29), were used in this study. MDA-MB-436 cells are homozygous for a 5396 + 1G>A mutation in the splice donor site of exon 20 resulting in loss of the wild-type *BRCA1* allele. Analysis has identified two *BRCA1* transcript lengths; one transcript had skipped exon 20 (predicting an in-frame deletion of 28 amino acids in the encoded BRCA1 proteins) and the second transcript had spliced at a cryptic splice site in intron 20 (5396 + 88/89) (predicting an insertion of 7 amino acids encoded by intron sequences followed by a termination codon) (Merajver et al., 1995, Elstrodt et al., 2006).

MCF7 cells were obtained from ATCC® and were maintained with RPMI-1640 supplemented with 10% FBS and 1% penicillin/streptomycin. MDA-MB-436 cells were obtained from CLS and were grown in DMEM: Ham’s F12 supplemented with 2 mM L-glutamine, 10% FBS and 1% penicillin/streptomycin. All materials were obtained from Sigma/PAA (UK).

### 2.3. Methods
2.3.1. Subculture of cell lines

All cell lines were cultured at 37°C in a humidified incubator with 5% CO₂ and 95% air. To subculture, all cells were handled separately with their own reagents and under sterile conditions. Medium was removed and cells washed twice with phosphate buffered saline (PBS) (without Ca²⁺ and Mg²⁺). Cells were detached from the flask by adding 5 ml or less of 0.5mg/ml trypsin-EDTA (Sigma/PAA, UK) and incubated for 5 minutes at 37°C. Medium was used to deactivate the trypsin and the suspension was centrifuged at 1000 rpm for 5 minutes to remove residual trypsin. The pellet was resuspended in medium, mixed well and split into a new tissue culture flask. All cell lines were routinely confirmed to be mycoplasma negative.

2.3.2. Cryopreservation of cell lines

Cells were cryopreserved at a low passage number for future use. Prior to freezing cells were trypsinised and counted using a haemocytometer. 1 x 10⁶ cells were resuspended in 1 ml of freezing medium, stored in a cryovial at -80°C overnight, and then transferred to liquid nitrogen for long term storage. Freezing medium consisted of complete medium for the cell line + 10% DMSO, with the exception of HeLa cells where the freezing medium was 50% FBS + 40% PBS +10% DMSO. Cells were recovered from liquid nitrogen by thawing rapidly in a water bath at 37°C. The cells were then resuspended in 9 ml fully supplemented culture medium and centrifuged at 1000 rpm for 5 minutes in order to remove traces of DMSO. The medium was then removed
and the pellets were resuspended in medium and transferred to tissue culture flasks.

2.3.3. AQueous non-radioactive cell proliferation assay (MTS assay)

2.3.3.1. Background principles

MTS assay (Promega, UK) is frequently used for screening different compounds to determine if they have effects on cell proliferation or show direct cytotoxic effects that eventually lead to cell death.

MTS solution is composed of a tetrazolium compound [3-(4, 5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium, inner salt; MTS] and an electron coupling reagent phenazine methosulfate (PMS). Cells reduce MTS into a formazan product that is soluble in tissue culture medium; formazan is produced by dehydrogenase enzymes that are only found in metabolically active cells (Riss et al., 2011, Cory et al., 1991).

2.3.3.2. Assay

To evaluate the cytotoxicity of varying doses of inhibitors (KU55933, NU7441 or NU1025), MTS assays were performed according to the manufacturer’s recommendation. 500 - 2000 cells, in 200 µl of medium, were seeded into each well of a 96-well plate. Cells were allowed to adhere for 24 hours, followed by incubation with varying concentrations of inhibitors. The MTS assay was performed on day 6.
2.3.3.3. Data analysis

On day 6, 20 µl of MTS reagent was added to each well, and the plate was incubated for 3-4 hours at 37ºC in the dark. Formazan absorbance was measured at 490 nm using a plate reader (FLUOstar OPTIMA, UK/ Infinite® F50, UK). Percentage of the absorbance was determined by comparison to a control population of untreated cells. This assay was performed three times, each time in triplicate. Graphs were produced and statistical analysis performed using the Microsoft Excel 2010 and GraphPad prism software version 6.02.

2.3.4. Clonogenic survival assay

2.3.4.1. Background principles

The clonogenic survival assay allows a determination of cell survival after exposure to potentially cytotoxic agents. The assay measures viable cells that have undergone at least 5-6 rounds of replication to produce a colony and have therefore survived the drug exposure (Munshi et al., 2005).

2.3.4.2. Plating efficiency

Cell lines were plated into 6-well plates at varying densities. Plates were incubated for 12-14 days under normal incubator conditions, then media was removed and wells were fixed (with methanol and acetic acid mixture) for 10 minutes. Plates were stained with crystal violet and colonies counted manually.
2.3.4.3. Clonogenic assay

HeLa SilenciX cells (200-500 cells/well) and Breast cancer cells (300-900 cells/well) where seeded in a 6-well plate. Cells were allowed to adhere for 24 hours. Compounds (KU55933, KU60019, NU7441, NU7026 or 3-Aminobenzamide) were added at the indicated concentrations, with a control for each cell line that did not receive drug. The plates were left in the incubator for 14 days. After incubation, the media was discarded, cells were fixed (with methanol and acetic acid mixture) and colonies stained with crystal violet and counted manually.

To evaluate cisplatin chemopotentiation cells were seeded as above and after 24 hours cells were exposed to cisplatin for 16 hours. Plates were then washed twice with PBS, fresh media added and plates placed in an incubator with or without inhibitors 14 days then they were fixed and stained as above. All assays were performed in duplicate and repeated three times.

A second clonogenic survival assay method was also employed to determine validity of the previously described protocol. In this method, cells lines were plated as previously described, cells were allowed to adhere for 24 hours, after which inhibitory compound was added at varying concentrations. Cells were incubated for 24 hours, after which the plates were gently washed with PBS and the medium replaced without inhibitor. Plates were again incubated for 14 days prior to staining as previously described.
2.3.4.4. Data analysis

Blue stained visible colonies with ≥ 50 cells were counted manually and surviving fraction was calculated as follows:

Surviving Fraction (SF) = \([\text{No. of colonies formed} / (\text{No. of cells seeded} \times \text{Plating efficiency})]\).

Plating efficiency was calculated by:

Plating Efficiency (PE) = (\text{No. of colonies formed} / \text{No. of cells seeded}) \times 100.

When comparing between the two clonogenic protocols, and in the clonogenic combination study, the inhibitory concentration of an inhibitor that result in the death of 50% of the cell population (IC\(_{50}\)) was calculated using curve fitting with nonlinear regression analysis. This assay was performed three times. Graphs were produced and statistical analysis performed using the Microsoft Excel 2010 and GraphPad prism software version 6.02.

2.3.5. γH2AX immunofluorescence microscopy assay

2.3.5.1. Background principles

The histone protein H2AX is one of three types of histone H2A found in eukaryotic cells and has been shown to be involved in DNA repair and is a central component of signalling pathways activated in response to DNA double strand breaks. Within minutes of a DNA double strand break, H2AX becomes rapidly phosphorylated by ATM, ATR or DNA-PK, on a serine 139 residue forming γH2AX foci, each foci is thought to correspond to one double strand break within the nucleus (Rogakou et al., 1998, Rogakou et al., 1999, Paull et
This focus can be detected using antibody staining and fluorescence microscopy.

2.3.5.2. Assay

100,000 cells per well were seeded onto sterile coverslips in 6-well plates. Cells were allowed to adhere for 24 hours, and then cells were incubated in medium containing ATM inhibitors (KU55933/KU60019) or DNA-PKcs inhibitors (NU7441/NU7026). After 24 and 48 hours treatment, cells were washed with PBS and fixed with ice-cold 100% methanol for 10 minutes. After rehydrating in PBS, cells were permeabilized in blocking buffer [KCM buffer (120mM potassium chloride, 20mM sodium chloride, 10mM Tris-HCl, 1mM EDTA, 0.1% Triton X-100) with 2% bovine serum albumin, 10% milk powder, and 10% normal goat serum] for one hour. Cells were incubated with primary anti phospho-histone H2AX (Ser139) antibody (mouse monoclonal antibody, Millipore Corp., dilution 1:200) in blocking buffer at room temperature for 1 hour. After incubation, the cells were washed in KCM buffer and incubated with secondary anti-mouse antibody (polyclonal goat anti-mouse immunoglobulins, DAKO, dilution 1:200) at room temperature for 1 hour in the dark, followed by 2-3 washes in KCM buffer. Coverslips were air dried at room temperature, stained with DAPI (Vectashield Hard Set, Vector Laboratories Inc., Burlingame, USA) and stored overnight at 4°C before analyses.

To evaluate cisplatin chemopotentiation, cells were seeded as above and after 24 hours cells were exposed to cisplatin for 16 hours. Plates were then washed twice with PBS, fresh media added and plates placed in an incubator
with or without inhibitors for 24 and 48 hours then washed and fixed as above. Immunofluorescence microscopy was then performed as described below.

2.3.5.3. Data analysis

Images were obtained using Olympus BX40 microscope and the images captured by cellSens (Vers 1.4) Imaging Software and camera (Olympus). Cells containing γH2AX foci were determined in 100 cells per slide in three separate experiments. Nuclei containing more than six γH2AX foci were considered positive. This method (> 6 γH2AX foci) of quantification has been shown to be reliable. Moreover, the basal level of γH2AX foci for the cell lines used in the present study (Breast cancer and HeLa cell lines) has previously been shown to be a maximum of 6 (Farmer et al., 2005, Bryant et al., 2005, Sultana et al., 2012). The student t-test was used to calculate if there is a significant difference in sensitivity between mutant/knockdown cells before and after treatment with inhibitors. A p-value ≤0.05 was defined as a significant relationship. Graphs were produced and statistical analysis performed using the Microsoft Excel 2010 and GraphPad prism software version 6.02.

2.3.6. Flow cytometric analyses for cell cycle

2.3.6.1. Background principles

Distribution of cell cycle by flow cytometry is a widely used procedure. This assay is based on analysis of the cellular DNA content after staining cells with the fluorescent DNA binding dye, propidium iodide (PI). This method
reveals distribution of cells in three major phases of the cycle (G1, S and G2/M) (Ormerod and Kubbies, 1992, Pozarowski and Darzynkiewicz, 2004).

2.3.6.2. Assay

100,000 cells were plated in to T25 flasks, and were allowed to adhere for 24 hours. Inhibitory compound was added and after 48 hours treatment, cells were collected by trypsinization and centrifugation (1000 rpm for 5 minutes). The cell pellets were fixed in 70% ethanol in PBS, incubated at 4°C overnight to allow fixation and then stored under these conditions until FACS analyses. Prior to FACS analysis, fixed cells were centrifuged and the pellet was resuspended in 500 µl PBS containing propidium iodide (PI) (2 µg/ml) and DNAse-free RNase A (10 µg/ml). After incubation at 37°C for 1 hour, samples then were analysed.

To evaluate cisplatin chemopotentiation cells were seeded as above and after 24 hours cells were exposed to cisplatin for 16 hours. Plates were then washed twice with PBS, fresh media added and placed in an incubator with or without inhibitors 48 hours. Cell cycle analysis was then performed.

2.3.6.3. Data analysis

Samples were analysed on a Cytomics F500 flow cytometer (Beckman Coulter, USA) using a 488nm laser for excitation and a 620nm bandpass filter for collection of data. Data was analysed using FlowJo 7.6.1 (Tree Star, Ashland, USA). At least 50,000 cells from each cell suspension were analysed. This assay was performed three times; the student t-test was used to calculate if there is a significant difference in sensitivity between mutant/knockdown
cells before and after treatment with inhibitors. A p-value ≤0.05 was defined as a significant relationship. Graphs were produced and statistical analysis performed using the Microsoft Excel 2010 and GraphPad prism software version 6.02.

2.3.7. Annexin V flow cytometric analyses for Apoptosis

2.3.7.1. Background principles

Programmed cell death (Apoptosis), is a common form of cell death in eukaryotes and during embryonic development. It is characterized by chromatin condensation, altering in cell volume and translocation of membrane phosphatidylserine (PS) from the inner side of the plasma membrane to the surface (Figure 2.1). Annexin V is a Ca$^{2+}$-dependent phospholipid-binding protein with a high affinity for PS. Staining with Annexin V is used in conjunction with a second dye such as propidium iodide (PI) for identification of early and late apoptotic cells (Figure 2.2). Viable cells with intact membranes are negative (reject) to PI, whereas dead cells with damaged membranes are positive (permeable) to PI. Therefore, viable cells are both Annexin V and PI negative, early apoptotic cells are Annexin V positive and PI negative, late apoptotic cells are Annexin V and PI positive, finally necrotic cells are Annexin V negative and PI positive (Koopman et al., 1994, Riccardi and Nicoletti, 2006).
Figure 2.1. Annexin V flow cytometry mechanism (V-FITC).

In early apoptosis, phosphatidylserine, which is located inside the plasma membrane, transposes to the cell outer surface. Annexin V is a human vascular anticoagulant, with high affinity for PS. Annexin V-FITC/PI staining method is widely used in the detection of early apoptosis. Adapted from (www.flow-cytometry.us).

Figure 2.2. Annexin V and PI Staining.

Viable cells are both Annexin V and PI negative, early apoptosis cells are Annexin V positive and PI negative, late apoptosis cells are Annexin V and PI positive, finally necrotic are Annexin V negative and PI positive.
2.3.7.2. Assay

Cells were grown to sub-confluency (70-80%) and harvested. 100,000 cells were plated in to T25 flasks and were allowed to adhere for 24 hours. Inhibitory compound was added and cells were collected after 24 and 48 hours exposure. Cells were trypsinized and centrifuged at 1000 rpm for 5 minutes. The cell pellets were washed twice with ice-cold PBS and then resuspend in Binding buffer (BioLegend, UK) at a concentration of $1 \times 10^6$ cell/ml. 5 µl of FITC Annexin V (ImmunoTools, Germany) and 5 µl PI were added to 100 µl of the cell solution and incubated for 15 minutes at room temperature in the dark. 400 µl of the Binding buffer was then added and the sample analysed by FACS.

To evaluate cisplatin chemopotentiation cells were seeded as above and after 24 hours cells were exposed to cisplatin for 16 hours. Plates were then washed twice with PBS, fresh media added and placed in an incubator with or without inhibitors for 24 and 48 hours. Cell cycle analysis was then performed.

2.3.7.3. Data analysis

Samples were analysed on a Cytomics F500 flow cytometer (Beckman Coulter, USA) using a 488nm laser for excitation and a 575nm bandpass filter for collection of data. Data was analysed using FlowJo 7.6.1 (Tree Star, Ashland, USA). At least 20,000 cells from each cell suspension were analysed. This assay was performed three times; the student t-test was used to calculate if there is a significant difference in sensitivity between mutant/knockdown cells before and after treatment with inhibitors. Percentage of the absorbance was determined by comparison to a control population of untreated cells. A p-
value $\leq 0.05$ was defined as a significant relationship. Graphs were produced and statistical analysis performed using Microsoft Excel 2010 and GraphPad prism software version 6.02.

2.3.8. DNA repair gene expression profiling

2.3.8.1. Background principles

To determine the human DNA repair gene profile in BRCA1 deficient cell lines, RT-qPCR array was performed using the DNA repair RT² Profiler™ PCR Array (Qiagen, UK). This array represents 84 key genes involved in the Base excision, Nucleotide excision, Mismatch, Double-strand break, and other DNA repair pathways. Additionally, quantitative real time PCR (RT-qPCR) using QuantiTect® Primers (Qiagen, UK) was performed to confirm up/down gene expression.

2.3.8.2. RNA extraction

Total cellular RNA was isolated using the RNeasy mini kit (Cat No: 74104, Qiagen) according to the manufacturer’s instructions. Briefly, 350 $\mu$l of Buffer RLT was added to the $2 \times 10^6$ pelleted cells, and then vortexed. 350 $\mu$l of 70% ethanol was mixed with the lysate and then 700 $\mu$l of the lysate was transferred to an RNeasy spin column. The sample was then centrifuged for 15 seconds at 10,000 rpm. The flow through was discarded and 350 $\mu$l of Buffer RW1 was added to the RNeasy spin column to wash the membrane. The sample was centrifuged for 15 seconds at 10,000 rpm, and the flow through was discarded.
DNA digestion was performed to eliminate genomic DNA contamination using the DNase digestion kit (Cat No: 79254, Qiagen). 70 μl of Buffer RDD and 10 μl of DNase I stock solution was added into a 0.5 ml eppendorf. This DNase I incubation mix was centrifuged for 15 seconds at 10,000 rpm. 80 μl of the DNase I mix was then added directly to the RNeasy spin column membrane, and incubated at room temperature for 15 minutes. 350 μl of Buffer RW1 was added to the RNeasy spin column, and the sample was centrifuged for 15 seconds at 10,000 rpm. The flow through was discarded and 500 μl of Buffer RPE was added to the RNeasy spin column to wash the membrane. The sample was centrifuged for 15 seconds at 10,000 rpm. 500 μl of Buffer RPE was added to the RNeasy spin column to wash the membrane, and the sample was centrifuged for 2 minutes at 10,000 rpm. Following centrifugation, the RNeasy spin column was carefully placed in a new 1.5 ml eppendorf, and 40 μl of RNase-free water was added directly to the spin column membrane. The sample was centrifuged for 1 minute at 10,000 rpm to elute the RNA.

The concentration (µg/µl) of the eluted RNA was performed by measuring absorbance ratio at 260:280 nm using a spectrophotometer system (Nanodrop 2000c, Thermo Scientific, USA) and RNA aliquots were then stored at -80ºC.

2.3.8.3. cDNA Synthesis

Equal amounts of RNA (0.5 µg) were reverse transcribed to form cDNA using RT² First Strand Kit (Cat No; 79254330401, Qiagen). For each RNA sample 2 µl GE (DNA elimination buffer) and nuclease-free water was added to bring the total volume of genomic DNA elimination mixture to 10 µl. The genomic DNA elimination mixture was centrifuged for 10 seconds at 10,000
rpm. The mixture was incubated at 42°C for 5 minutes. Following incubation, the genomic DNA elimination mixture was chilled on ice for at least one minute. For preparation of the room temperature cocktail, 4 μl of 5X RT Buffer 3 (BC3), 1 μl of Primer & External Control Mix (P2), 2 μl of RE3 (RT Enzyme Mix 3), and 3 μl of nuclease free water were added to a sterile 1.5 mL eppendorf. The 10 μl of RT cocktail was added to the genomic DNA elimination mixture. The reaction was incubated at 42°C for 15 minutes and the reaction was stopped by heating at 95°C for 5 minutes. Following heating, 91 μl of nuclease free water was added to the cDNA and held on ice until the array was prepared or the cDNA was stored at -80°C.

2.3.8.4. RT² Profiler™ DNA Repair PCR Array

To evaluate the expression of 84 DNA repair genes, cDNA samples were mixed with RT² SYBR Green Mastermix (Qiagen) according to the supplier’s instructions. Briefly, the PCR component mix was prepared as follow:

**Table 2.1. Reaction mix for RT² Profiler™ DNA Repair PCR Array**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>RT² SYBR Green Mastermix</td>
<td>1350 μl</td>
</tr>
<tr>
<td>cDNA synthesis reaction</td>
<td>102 μl</td>
</tr>
<tr>
<td>RNase-free water</td>
<td>1248 μl</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>2700 μl</strong></td>
</tr>
</tbody>
</table>

25 μl of the mix was added to each well of the PCR array plate (Cat No: PAHS-042ZC). The plate was sealed using optical adhesive film and centrifuged for 1 minute to remove bubbles. RT PCR was performed on an Applied Biosystems 7500 FAST cycler, the cycler conditions were; initial 1:10
minutes at 95ºC; followed by 2-41:15 seconds at 95ºC, then 1 minute at 60ºC. Quality controls included within the array plates confirmed the lack of DNA contamination and successfully tested for RNA quality and PCR performance.

2.3.8.5. Quantitative real time PCR (RT-qPCR)

To confirm up/down regulation of gene expression as shown by RT² Profiler™ DNA Repair PCR Array, quantitative real time PCR (RT-qPCR) using QuantiTect® Primers (Qiagen, UK) was performed according to the supplier’s instructions. Briefly, the PCR component mix was prepared as follow;

Table 2.2. Reaction mix for Quantitative real time PCR (RT-qPCR).

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>cDNA from RT² First Strand synthesis</td>
<td>2 μl</td>
</tr>
<tr>
<td>10x QuantiTect primer</td>
<td>1 μl</td>
</tr>
<tr>
<td>SYBR Green Mastermix</td>
<td>5 μl</td>
</tr>
<tr>
<td>RNase-free water</td>
<td>2 μl</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>10 μl</strong></td>
</tr>
</tbody>
</table>

PCR mix was added to each well of the PCR plate (Applied Biosystems®, UK). The plate was sealed using optical adhesive film and centrifuged for 1 minute to remove bubbles. RT qPCR was performed on an ABI prism 7700 (Applied Biosystems) using SYBR green detection (Applied Biosystems®, UK). Thermal cycler conditions included incubation at 95ºC for 10 minutes followed by 40 cycles of 95ºC for 15 seconds and 60ºC for 1 minute. Following the 40 cycles, the products were heated from 60ºC to 95ºC over 20 minutes to allow melting curve analysis to be done. This allowed the
specificity of the products to be determined (single melting peak) and confirmed the absence of primer-dimers. The housekeeping gene GAPDH was used to standardise the samples and the relative expression of BRCA1, XRCC1, APE1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs were therefore calculated as the ratio between the expression of test gene and the expression of housekeeping gene. Positive and negative controls (no template) were included in each experiment and all reactions were performed three times.

<table>
<thead>
<tr>
<th>Gene Symbol</th>
<th>Source</th>
<th>Catalog no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>APE1</td>
<td>Qiagen</td>
<td>QT00012474</td>
</tr>
<tr>
<td>XRCC1</td>
<td>Qiagen</td>
<td>QT00016688</td>
</tr>
<tr>
<td>SMUG1</td>
<td>Qiagen</td>
<td>QT00002317</td>
</tr>
<tr>
<td>Pol β</td>
<td>Qiagen</td>
<td>QT00088655</td>
</tr>
<tr>
<td>Lig3</td>
<td>Qiagen</td>
<td>QT00017115</td>
</tr>
<tr>
<td>ATM</td>
<td>Qiagen</td>
<td>QT00061593</td>
</tr>
<tr>
<td>DNA-PKcs</td>
<td>Qiagen</td>
<td>QT00086828</td>
</tr>
<tr>
<td>BRCA1</td>
<td>Qiagen</td>
<td>QT00039305</td>
</tr>
<tr>
<td>GAPDH</td>
<td>Qiagen</td>
<td>QT00079247</td>
</tr>
<tr>
<td>β2M</td>
<td>Applied Biosystems</td>
<td>Forward /GAGTATGCCTGCCGTGTG Reverse /AATCCAAATGCGGCATCT</td>
</tr>
</tbody>
</table>

2.3.8.6. Data analysis

Raw data from both real-time PCR methods were uploaded using a PCR array data analysis template available at http://pcrdataanalysis.sabiosciences.com/pcr/arrayanalysis.php. The integrated Web-based software package for the PCR array system automatically performed all comparative threshold cycle (ΔΔCt)–based fold-change calculations from the uploaded data. For these calculations, GAPDH was used for normalization of the data. After normalization, the relative expression of each gene was averaged for the two samples in each cell line. Fold changes in average gene expression were expressed as the difference in expression of
BRCA1 deficient cells compared with BRCA1 proficient cells (control). Statistical significance determined for the RT² Profiler™ Array using the Bonferroni correction for multiple testing, with p values less than 0.05 (threshold p < 0.000595 = 0.05/84). RT-qPCR statistical significance determined using t test, with p values less than 0.05. Analysis performed using GraphPad prism software version 6.02.

2.3.9. Western Blot analysis

2.3.9.1. Background principles

Western blot is a method developed more than 30 years ago (Burnette, 1981). This process involves separation of specific proteins in given cells or tissue extract on a sodium dodecyl sulphate (SDS)-polyacrylamide gel, followed by electrophoresis transfer to a nitrocellulose membrane and detected using specific antibodies (primary antibodies) that bind to the target protein. Horseradish peroxidise (HRP) enzyme conjugated secondary antibodies were used to bind with their specific primary antibodies which can be detected using ECL™ chemiluminescent system. Visualisation of the immune complex was performed using X-ray film.

2.3.9.2. Preparation of cell lysate

70-80% confluent cells were washed with PBS, trypsinised and counted using a haemocytometer. The cell pellet was resuspended in media, 1 x 10⁶ or 2 x 10⁶ cells were lysed using 100 µl of RIPA buffer (20 mM Tris, 150 mM NaCl, 1% Nonidet p-40, 0.5% sodium deoxycholate, 1 mM EDTA, 0.1% SDS; Sigma) containing 10% of protease inhibitor (Sigma) and 1% of phosphatase
inhibitor cocktails 2 and 3 (Sigma). The lysate was incubated on ice for 1 hour, then repeatedly aspirated with a fine-gauge needle and incubated overnight at -20°C. Next day lysates were centrifuged at 13000 rpm for 20 minutes at 4°C and the supernatant was stored at -20°C.

2.3.9.3. Protein quantification (Bradford assay)

Bradford assay (Bio-Rad, UK) is a method based on binding Coomassie dye to protein (Bradford, 1976). Differential colour change occurs in response to the protein concentration. Briefly, 10 µl of protein were mixed with 250 µl assay solution in 96-well plate. The absorbance value was measured at 595 nm by plate reader (FLUOstar OPTIMA, UK/ Infinite® F50, UK) with in 1 hour. A protein standard curve was generated using bovine serum albumin (BSA) against corresponding protein concentrations. The determination of the protein concentration, in respective cell lysates, was carried out to ensure equal loading of protein in the wells of western blot gel.

2.3.9.4. Preparation of cell lysates for electrophoresis

Equal volumes (20 – 50 µg) of protein were mixed with 5 µl lithium dodecyl sulphate (LDS) sample buffer (Expedeon, UK), 1 µl Dithiothreitol (DTT) (Invitrogen, UK) and dH₂O. The mix was incubated at 70°C for 10 minutes, and then placed on ice.

2.3.9.5. Polyacrylamide gel electrophoresis and Protein transfer

Samples were loaded onto precast SDS 4-12% gels (Expedeon, UK) and were run in 3-(N-morpholino) propanesulphonic acid-sodium dodecyl sulphate
(MOPS-SDS) running buffer (Expedeon), at a constant voltage of 125-200 V for 60-120 minutes.

Using the Xcell II Blot Module (Invitrogen), proteins were transferred to a nitrocellulose membrane (Whatman, GE Healthcare) in transfer buffer [20% (v/v) methanol, 50 mM Tris-HCl, and 380 mM glycine] by application of a 25V current for 60-120 minutes.

2.3.9.6. Immunoblotting

Membranes were blocked by incubation with PBS-T (PBS, 0.05% Tween 20) containing 5% non-fat milk for 60 minutes at room temperature. Membranes were incubated with primary antibody diluted in blocking buffer for 60 minutes at 37°C or overnight at 4°C. The membranes were washed 3 times (5 minutes each) with PBS-T. Protein expression was examined by application of HRP-labelled secondary antibody (Dako, Denmark) at 1:1000 dilution in blocking buffer for 60 minutes at room temperature. The membranes were washed 3 times (5 minutes each) with PBS-T and detected on photographic film using ECL reagent (GE Healthcare).
2.3.9.7. Primary antibodies

<table>
<thead>
<tr>
<th>Antigen</th>
<th>Antibody</th>
<th>MW</th>
<th>Source</th>
<th>Catalog no.</th>
<th>Dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td>APE1</td>
<td>Rabbit</td>
<td>37 kDa</td>
<td>Novus</td>
<td>NB 100-101</td>
<td>1:1000</td>
</tr>
<tr>
<td>XRCC1</td>
<td>Mouse</td>
<td>85 kDa</td>
<td>Neomarkers</td>
<td>MS-434-PO</td>
<td>1:150</td>
</tr>
<tr>
<td>SMUG1</td>
<td>Goat</td>
<td>37 kDa</td>
<td>Acris</td>
<td>APO884PU-N</td>
<td>1:1000</td>
</tr>
<tr>
<td>Pol β</td>
<td>Rabbit</td>
<td>38 kDa</td>
<td>Abcam</td>
<td>Ab26343</td>
<td>1:1000</td>
</tr>
<tr>
<td>Lig3</td>
<td>Rabbit</td>
<td>113 kDa</td>
<td>Abcam</td>
<td>Ab96576</td>
<td>1:1000</td>
</tr>
<tr>
<td>BRCA1</td>
<td>Rabbit</td>
<td>220 kDa</td>
<td>Santa Cruz</td>
<td>Sc-642</td>
<td>1:150</td>
</tr>
<tr>
<td>ATM</td>
<td>Goat</td>
<td>350 kDa</td>
<td>Abcam</td>
<td>ab2631</td>
<td>1:1000</td>
</tr>
<tr>
<td>DNA-PKcs</td>
<td>Rabbit</td>
<td>460 kDa</td>
<td>Abcam</td>
<td>ab32566</td>
<td>1:1000</td>
</tr>
<tr>
<td>β-actin</td>
<td>Mouse</td>
<td>42 kDa</td>
<td>Sigma</td>
<td>A2228</td>
<td>1:10000</td>
</tr>
</tbody>
</table>

2.3.9.8. Data analysis

Photographic films were scanned and analysed using Image Studio Lite (Ver 3.1) (Li-Cor, USA). β-actin was used for normalization of the data. This assay was performed in two or three independent experiments; the student t-test was used to calculate the significant difference between control cells and mutant/knockdown cells. A p-value ≤0.05 was defined as significant. Graphs were produced and statistical analysis performed using the GraphPad prism software version 6.02.
Chapter 3

DNA repair profiling in BRCA1 deficient and proficient cells
3. DNA repair profiling in BRCA1 deficient and proficient cells

3.1. Introduction

Germ-line mutations in the \textit{BRCA1} gene are one of the most common causes of hereditary forms of breast and ovarian cancers. \textit{BRCA1} mutation carriers have 80\% lifetime risk of developing breast cancer, and 40-50\% risk of developing ovarian cancer (Risch et al., 2001). In addition, \textit{BRCA1} mutations increase the risk of other types of cancer, such as pancreatic and prostate cancers (Thompson and Easton, 2002). Moreover, in sporadic breast cancer, epigenetic mechanisms of gene inactivation are now well recognized as a potential alternative to genetic mutation in the silencing of the \textit{BRCA1} promoter in up to 11-14\% of tumours (Turner et al., 2004). Dysfunctional BRCA pathway may also contribute to a ‘BRCAness’ phenotype in about 25\% of cancers (Turner et al., 2004), where breast cancers do not harbour germline \textit{BRCA} mutations but display similar phenotypes including HR deficiency. The term ‘BRCAness’ is a phenotype which refers to molecular and histopathological characteristics observed in a subset of breast cancers that exhibit similarity to BRCA deficient tumours, including ‘basal phenotype’ and ‘triple negative’ (oestrogen receptor, progesterone receptor and HER2 negative) cancers (Turner et al., 2004, Matros et al., 2005). This chapter investigated, whether BRCA1 deficient cells compared to BRCA1 proficient cells exhibit altered profile of DNA repair expression.
3.1.1. BRCA1 mutation in breast cancer cell lines

Four established breast cancer cell lines with *BRCA1* mutation have been reported, HCC1937 was the first BRCA1 deficient cell line reported in 1998. HCC1937 was derived from a germline *BRCA1* mutation carrier (Tomlinson et al., 1998). In 2006 Elstrodt and colleagues screened 41 human breast cancer cell lines and found three to be *BRCA1* mutant (summarised in Table 3.1), the same results were confirmed by analysis of 51 human breast cancer cell lines in 2013 (Elstrodt et al., 2006, Riaz et al., 2013).

<table>
<thead>
<tr>
<th>Cell line</th>
<th>BRCA1 Mutation status</th>
<th>BRCA1 transcript expression</th>
<th>Exon</th>
<th>ER</th>
<th>PGR</th>
<th>P53</th>
<th>EGFR</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDA-MB-436</td>
<td>5396 + 1G&gt;A</td>
<td>++*</td>
<td>20</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>SUM1315M02</td>
<td>185delAG</td>
<td>+</td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>SUM149PT</td>
<td>2288delT</td>
<td>+</td>
<td>11</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>HCC1937</td>
<td>5382insC</td>
<td>++</td>
<td>20</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

Table 3.1. BRCA1 mutations in breast cancer cell lines.
ER, estrogen receptor; EGFR, epidermal growth factor receptor; PGR, progesterone receptor. For transcript expression: (+, low transcript levels); (++, normal transcript levels); (++*, Two transcript lengths that both differ from the wild type sequence). For protein expression: (+, expression) and (-, no expression) (Tomlinson et al., 1998, Elstrodt et al., 2006, Riaz et al., 2013).
3.1.2. Rationale for the study

As discussed previously (section 1.7.1) BRCA1 plays a role in many DNA repair pathways including, BER, NER, MMR, HR and NHEJ. Therefore in this chapter, a comprehensive analysis of the DNA repair pathway genes in BRCA1 deficient cells compared to proficient cells will be described using the RT² Profiler™ DNA Repair PCR Array (as described in section 2.3.8.4.). The RT² Profiler PCR Array has the advantage of combining RT-PCR performance with the ability of microarrays to detect the expression of 84 DNA repair genes simultaneously.

BRCA1 has been suggested to stimulate the activity of three BER enzymes (see section 1.7.1); OGG1, NTH1 and APE1. These enzymes mediate repair of three signature oxidative lesions; 8-oxoguanine, thymine glycol and abasic sites, respectively (Saha et al., 2010). A more recent study supports these findings and has confirmed down-regulation of the same enzymes in BRCA1 deficient cells, it also suggested XRCC1 to be regulated by BRCA1 (De Summa et al., 2014). XRCC1 protein associates with Pol β, and Lig3, to form a complex that repairs the single strand DNA breaks generated during the BER process. In addition, recent studies suggest a cross talk between BRCA1 and other BER genes. For instance, a functional interaction between Pol β and BRCA1 (Masaoka et al., 2013) has been described, and an association between low SMUG1 protein expression and loss of BRCA1 expression (Abdel-Fatah et al., 2013). For this reason in the current study BER genes (APE1, XRCC1, SMUG1, Pol β and Lig3) will be investigated for a potential link with BRCA1 deficiency. In addition, as the main aim of this study is to investigate synthetic lethality in a BRCA1 deficient system by targeting the DSB pathway, ATM (a
key enzyme in HR) and DNA-PKcs (a key enzyme in NHEJ) genes will be investigated as well. Therefore, the selected genes (BRCA1, APE1, XRCC1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs) will be investigated using the RT² Profiler™ Array and then further assessment of the results will be carried out by quantitative real time PCR (RT-qPCR) using QuantiTect® Primers (as described in section 2.3.8.5.). Finally, the mRNA expression of the selected genes will be compared with protein expression using western blot analysis.

### 3.1.2.1. Aims

The aims of this chapter were as follows:

1. To explore DNA repair expression in BRCA1 proficient and deficient cell lines using RT² Profiler™ DNA Repair PCR Array.

2. To confirm up or down regulation of selected DNA repair genes in a panel of BRCA1 proficient and deficient cell lines using Quantitative real time PCR (RT-qPCR).

3. To investigate alteration in protein expression of selected DNA repair factors in BRCA1 proficient and deficient cell lines using western blot analysis.
3.2. Materials and Methods

Materials and methods for the assays listed are described in full in Chapter 2. BRCA1 deficient MDA-MB-436 breast cancer cells and BRCA1 proficient MCF7 breast cancer cells were used in this study (Table 3.2). A second set of BRCA1 proficient and deficient cell line model was also used; BRCA1 deficient HeLa SilenciX cells and Control BRCA1 proficient HeLa SilenciX cells.

To make it easier for discussion of the results, BRCA1 deficient HeLa SilenciX cells are referred to as BRCA1 HeLa and Control BRCA1 proficient HeLa SilenciX cells are referred to as Control HeLa.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>BRCA1 Mutation status</th>
<th>BRCA1 transcript expression</th>
<th>ER</th>
<th>PGR</th>
<th>P53</th>
<th>EGFR</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDA-MB-436</td>
<td>5396 + 1G&gt;A</td>
<td>+¹</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>MCF7</td>
<td>Wild type</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 3.2. Breast cancer cell lines used in this project.

ER, estrogen receptor; EGFR, epidermal growth factor receptor; PGR, progesterone receptor. For transcript expression: (+, normal transcript levels); (+¹, Two transcript lengths that both differ from the wild type sequence). For protein expression: (+, expression) and (-, no expression) (Elstrodt et al., 2006, Riaz et al., 2013).
3.3. Results

3.3.1. Determination of BRCA1 expression in cell lines

3.3.1.1. Western blot analysis demonstrates reduced BRCA1 protein expression

The protein expression of BRCA1 was determined in all cell lines by western blotting. Briefly, cell lysates were prepared from 70-80% confluent cells and loaded onto polyacrylamide gels as described in section 2.3.9. BRCA1 was resolved on precast SDS 4-12% gels. Following electrophoresis, proteins were electrotransferred onto nitrocellulose and probed for target protein using specific antibody (Table 2.4). Recombinant BRCA1 protein (Sigma/UK, 0.02µl) was also run as control on the gel, moreover as a loading control; β-actin antibody was used.

![Image of western blot results]

Figure 3.1. BRCA1 protein expression in the cell lines studied by western blot

Whole cell protein lysates were collected for each cell line and 50 µg samples were separated by electrophoresis on denaturing polyacrylamide gels as described in section 2.3.9. Recombinant BRCA1 protein was also run as control on the gel. Blot was probed using specific BRCA1 antibody with predicted molecular weight 220 kDa. As a loading control, β-actin antibody (42 kDa) was used.
As expected, MDA-MB-436 and BRCA1 HeLa cell lines are BRCA1 deficient (Figure 3.1), when compared to MCF7 and Control HeLa cell lines. Quantification of protein expression is shown in Figure 3.2. Significant under-expression of BRCA1 protein was confirmed in BRCA1 HeLa (fold change mean = 0.09, p < 0.0001) and MDA-MB-436 cells (fold change mean = 0.07, p = 0.0001) compared to Control HeLa and MCF7 cells respectively.

Figure 3.2. Quantification of BRCA1 protein expression

(A) Relative BRCA1 expression in BRCA1 HeLa cells was calculated in comparison to Control HeLa cells. (B) Relative BRCA1 expression in MDA-MB-436 cells was calculated in comparison to MCF7 cells. Values plotted are means ± SD of the fold-change (ratio of protein/β-actin normalized to control) *** p<0.001 and **** p<0.0001.

3.3.1.2. RT- qPCR demonstrate reduced BRCA1 mRNA expression

After confirming low expression of BRCA1 protein in BRCA1 HeLa and MDA-MB-436 cells, mRNA expression was determined by RT-qPCR as described in section 2.3.8.5. Relative expression were calculated for each cell line and compared to their respective control cell lines. The mRNA expression
of housekeeping genes \( \beta 2 \)-microglobulin (\( \beta 2M \)) and GAPDH (primer details in Table 2.3) were used to standardise the samples. The relative expression of BRCA1 mRNA was calculated as the ratio between the expression of BRCA1 gene and the expression of the housekeeping gene. Negative control was included in each experiment. As both housekeeping genes showed similar results, GAPDH was used for all subsequent experiments. Data as expected (Figure 3.3) demonstrated low mRNA expression of BRCA1 in BRCA1 HeLa cells compared to Control HeLa (mean fold change = 0.16; \( p = 0.0015 \)). Low BRCA1 mRNA expression was also seen in MDA-MB-436 cells compared to MCF7 cells (mean fold change = 0.09; \( p = 0.0026 \)).

![Figure 3.3](image)

**Figure 3.3. Quantification of BRCA1 mRNA expression**

(A) Relative BRCA1 expression in BRCA1 HeLa cells was calculated in comparison to Control HeLa cells. (B) Relative BRCA1 expression in MDA-MB-436 cells was calculated in comparison to MCF7 cells. The housekeeping gene GAPDH was used to standardise the samples and the relative expression level of BRCA1 was therefore calculated as the ratio between the level of gene of interest and the level of housekeeping gene. Negative control (no template) was included in each experiment and all reactions were run in triplicate \( ** \) \( p<0.01 \).
3.3.2. RT² Profiler DNA Repair PCR Array

Real-time (RT-PCR) is a highly sensitive and reliable method for gene expression analysis. RT² Profiler PCR Arrays has the advantage of combining RT-PCR performance and the ability of microarrays to detect the expression of many genes simultaneously. RT² Profiler PCR array from Qiagen is a high throughput assay which allows analysis of the gene expression profiles in chosen cell populations. It consists of two step reverse transcription RT-PCR reaction: firstly isolated RNA and then converted to cDNA. Secondly the cDNA in combination with gene specific primers and a PCR master mix which contains a SYBR Green (which fluoresces when bound to double-stranded DNA (dsDNA)), subjected to thermal cycling.

Baseline fluorescence represents the background ‘noise’ fluorescence, which refers to the signal level during the initial cycles and it is automatically determined by the PCR cycler software. RT-PCR phases can be broken up into three sections; Exponential phase; each PCR cycle results in a doubling of dsDNA product which increases fluorescent signal until a pre-specified threshold is reached. This threshold cycle (C_t) is the cycle number at which fluorescent signal of the reaction crosses the baseline threshold, and is dependent upon the initial concentration of the primer target within the sample.

Linear phase; the reaction components are being consumed, gradually reducing reaction efficiency and products are starting to degrade. Plateau phase; the reaction has stopped, no more products are being made and if left long enough, the PCR products will begin to degrade. Graphical representation of C_t can be observed using amplification plots (Figure 3.4). By normalising the C_t of the gene of interest against the C_t of a housekeeping gene using \( \Delta \Delta C_t \) method.
gene expression can be compared in different samples (LifeTechnologies, 2012, Qiagen, 2011).

The Qiagen RT² Profiler PCR array is performed in a 96-well format (Figure 3.5), allowing analysis of 84 genes simultaneously. In addition to 5 housekeeping genes, the genomic DNA control (GDC) that specifically detects non-transcribed genomic DNA contamination with a high level of sensitivity. The reverse-transcription control (RTC) that tests the efficiency of the reverse-transcription reaction performed with the RT² First Strand Kit by detecting a template synthesized from the kit’s built-in external RNA control. The positive PCR control (PPC) consists of a pre-dispensed artificial DNA sequence and the assay that detects it. This control tests the efficiency of the polymerase chain reaction itself. Genes assessed in the RT² Profiler DNA PCR Array are listed in APPENDIX B.
Fluorescent reference dye (Rn), in this case SYBR Green, is normalised against ROX which is a passive reference dye contained within the PCR master mix. In amplification plots ΔRn scale, which is plotted against cycle number on a Log10, is a subtraction of baseline fluorescence from Rn. To demonstrate if the amplification was successful, post-amplification melting curve analysis can be performed. After thermo-cycling is completed, fluorescence of the PCR product is recorded while temperature is increased. When the melting point (Tm) is reached it is followed by a sudden decrease in fluorescent signal. Tm may be affected by multiple factors like (DNA length, and the presence of base mismatches); therefore different PCR products display different melting characteristics. By plotting change in Fluorescence

Figure 3.5. RT² Profiler PCR Array plate layout.

Wells A1 to G12 each contain a real-time PCR assay for a DNA repair pathways related gene. Wells H1 to H5 contain a housekeeping genes panel to normalize array data (HK1-5). Well H6 contains a genomic DNA control (GDC). Wells H7 to H9 contain replicate reverse-transcription controls (RTC). Wells H10 to H12 contain replicate positive PCR controls (PPC). Adapted from (Qiagen, 2011).
against change in Temperature (−ΔF/ΔT) to give a melting profile, it is possible to detect the presence of nonspecific product or ‘primer-dimers’ (binding of homologous primers to self rather than template) as additional peaks on the melting curve. To avoid presence of non-specific product, optimise the concentration of RNA was needed to find the concentration that gives successful amplification with less background noise (lifetechnologies, 2012, Qiagen, 2011).

3.3.3. Optimisation of the purified RNA for RT-PCR

High-quality RNA is essential for obtaining good, RT-PCR results. The most important prerequisite for any gene expression analysis experiment is consistently high-quality RNA from every experimental sample. So the concentration (µg/µl) of the purified RNA was performed by measuring absorbance ratio at 260:280 nm using a spectrophotometer system (Nanodrop 2000c, Thermo Scientific, USA) before every assay.

Optimisation of purified RNA for RT-PCR was carried out at different concentrations; starting with 1 µg total RNA per array. Amplification plot for the RT² Profiler array demonstrated successful amplification (Figure 3.6), however some background noise were detected. Melting curve analysis for both HeLa cell lines is summarised in Figure 3.7. Three nonspecific products (melting peaks) detected in Control HeLa cells, and one nonspecific product was detected with BRCA1 HeLa cells.
Figure 3.6. Optimisation of RNA concentration for RT² PCR.

Amplification plot for RT² Profiler DNA Repair Array, using 1 µg total RNA per array. Product is indicated by exponential fluorescence accumulation with increasing cycle number. Plot demonstrates successful amplification, though some background noise is detected.
As mentioned above, using 1 µg total RNA per array showed successful amplification, however some background noise in amplification plot along with nonspecific product detected in melting curves. Therefore the total RNA was reduced using 0.5 µg which shows successful amplification of all 84 primers for each of the cell lines (see Figure 3.8). Successful melting curve analysis for all the cell lines is summarised in Figure 3.9 with every specific product for all the 84 target genes.

**Figure 3.7. Melting curve analysis for HeLa cells with 1 µg total RNA per array.**

The presence of nonspecific product is observed as additional peaks to the left of the melting curve. Three nonspecific products (melting peaks) are seen in Control HeLa cells, and one nonspecific product was seen with BRCA1 HeLa.
Figure 3.8. Amplification plots for RT² Profiler DNA Repair Array.

Amplification of PCR product is indicated by exponential fluorescence accumulation with increasing cycle number.
Figure 3.9. Melting curves for RT$^2$ Profiler PCR Array.

Single melting peaks are observed for all the 84 primers assayed with all the cell lines, indicating high specificity.
3.3.4. BRCA1 deficiency is associated with multiple altered DNA repair gene expression profile

To investigate whether BRCA1 deficiency is associated with an altered DNA repair gene expression, both breast cancer (MDA-MB-436 and MCF7) and HeLa cell lines (BRCA1 HeLa deficient cells and Control HeLa proficient cells) were used to investigate mRNA expression by using RT² PCR DNA repair profile assay as described in section 2.3.8.4. Data analysis as described in section 2.3.8.6 was performed by importing the raw RT² PCR DNA repair profile assay data into a Qiagen analysis template that can compare multiple experimental replicates for one cell line against multiple experimental replicates of the control cells. The integrated Web-based software package for the PCR array system automatically performed all comparative threshold cycle (ΔΔCₜ)–based fold-change calculations from the uploaded data. For these calculations, GAPDH was used for normalization of the data. Scatter plots were charted to compare gene expression in BRCA1 wild type MCF7 cells to MDA-MB-436 and Control HeLa cells to BRCA1 HeLa cells as illustrated in Figure 3.10 and Figure 3.11.

A number of genes were initially observed to be under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cells (though not statistically significant), including: BER factors (Table 3.3) such as (APE1, Pol β, SMUG1, NTHL1, UNG, FEN1, OGG1, NEIL1 and XRCC1); HR factors (Table 3.4) such as (RAD21, RAD51, RAD54L and ATM); NHEJ factors (Table 3.5) such as (DNA-PKcs, LIG4, XRCC5 and XRCC6); NER factors (Table 3.6) such as (ERCC1, ERCC2, ERCC3, ERCC4 and ERCC5) and MMR factors
(Table 3.7) such as (MLH1, MSH2, MLH3, MSH3, MSH4 and MSH5). Other related DNA repair gene expressions results are shown in APPENDIX C.

Figure 3.10. Scatter plot comparing DNA repair gene expression in BRCA1 HeLa deficient cells against Control HeLa BRCA1 proficient cells. The area outside the grey lines indicates two-fold change in gene expression in BRCA1 HeLa deficient cells compared to Control HeLa proficient cells. Green circles indicate gene under-expression. Data are from 4 independent experiments.
Figure 3.11. Scatter plot comparing DNA repair gene expression in BRCA1 deficient MDA-MB-436 cells against Control BRCA1 proficient MCF7 cells.

The area outside the grey lines indicates two-fold change in gene expression in BRCA1 deficient MDA-MB-436 cells compared to Control BRCA1 proficient MCF7 cells. Green circles indicate gene under-expression. Data are from 3 independent experiments.
<table>
<thead>
<tr>
<th></th>
<th>Fold Change</th>
<th>P Value</th>
<th></th>
<th>Fold Change</th>
<th>P Value</th>
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<tr>
<td><strong>Base Excision Repair pathway</strong></td>
<td></td>
<td></td>
<td><strong>BRCA1 HeLa cells compared to Control HeLa cells.</strong></td>
<td><strong>MDA-MB-436 cells compared to MCF7 cells</strong></td>
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<td>0.019016</td>
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<td>CCNO</td>
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<td>0.3013</td>
<td>0.065909</td>
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Table 3.3. Base excision repair genes under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

Statistical significance determined using the Bonferroni correction for multiple testing, with p values less than 0.05 (threshold p < 0.000595 = 0.05/84).
### Table 3.4. Homologous recombination genes under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

<table>
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<tr>
<th>Gene</th>
<th>Fold Change</th>
<th>P Value</th>
<th>Fold Change</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
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<td>0.1102</td>
<td>0.000774</td>
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<td>BRCA2</td>
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<td>0.602818</td>
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<td>0.010779</td>
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<td>RAD21</td>
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<tr>
<td>RAD50</td>
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<td>0.221385</td>
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<td>RAD51D</td>
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### Table 3.5. Non-homologous end joining genes under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

<table>
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Statistical significance determined using the Bonferroni correction for multiple testing, with p values less than 0.05 (threshold p < 0.000595 = 0.05/84).
<table>
<thead>
<tr>
<th>Gene</th>
<th>BRCA1 HeLa cells compared to Control HeLa cells</th>
<th>MDA-MB-436 cells compared to MCF7 cells</th>
</tr>
</thead>
<tbody>
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<td>XPC</td>
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Table 3.6. Nucleotide excision repair genes under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

Statistical significance determined using the Bonferroni correction for multiple testing, with p values less than 0.05 (threshold p < 0.000595 = 0.05/84).
Table 3.7. Mismatch repair genes under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

Statistical significance determined using the Bonferroni correction for multiple testing, with p values less than 0.05 (threshold p < 0.000595 = 0.05/84).

3.3.5. DNA repair protein expression is down-regulated in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

The RT² Profiler™ Array suggested reduced expression of several mRNAs involved in BER, HR, NHEJ, NER and MMR in BRCA1 deficient cell lines as compared to BRCA1 proficient cell lines. As discussed previously (section 3.1.2.) selected DNA repair factors (APE1, XRCC1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs) will be investigated. To determine the protein expression of these factors, western blots were performed as described in section 2.3.9. Western blots results of BER, HR and NHEJ factors are shown in Figure 3.12 and Figure 3.13. Reduced expression of XRCC1, APE1, SMUG1, Pol β and lig3, along with both ATM and DNA-PKcs proteins were
observed in BRCA1 deficient cell lines as compared to BRCA1 proficient cell lines.

![Image showing western blot analysis of BER proteins in BRCA1 deficient and proficient cell lines.]

**Figure 3.12. Representative western blot analysis of Base Excision Repair proteins in BRCA1 deficient and proficient cell lines.**

BER Proteins (XRCC1, APE1, SMUG1, Pol β and Lig3) expression shows down-regulation in BRCA1 deficient cell lines (BRCA1 HeLa and MDA-MB-436) compared to BRCA1 proficient cell lines (Control HeLa and MCF7). Whole cell protein lysates were collected for each cell line and 25 or 50 µg samples were separated by electrophoresis on denaturing polyacrylamide gels as described in section 2.3.9. Blot was probed using specific antibodies with predicted molecular weight (XRCC1, 80 kDa; APE1, 37 kDa; SMUG1, 37 kDa; Pol β, 38 kDa and Lig3, 113 kDa). As a loading control, β-actin antibody (42 kDa) was used.
Figure 3.13. Representative western blot analysis of ATM and DNA-PKcs expression.

ATM and DNA-PKcs are down-regulated in BRCA1 deficient cell lines (BRCA1 HeLa and MDA-MB-436) compared to BRCA1 proficient cell lines (Control HeLa and MCF7). Whole cell protein lysates were collected for each cell line and 25 or 50 µg samples were separated by electrophoresis on denaturing polyacrylamide gels as described in section 2.3.9. Blot was probed using specific antibodies with predicted molecular weight (ATM, 350 kDa and DNA-PKcs, 460 kDa). As a loading control, β-actin antibody (42 kDa) was used.

3.3.6. Quantitative real time PCR (RT-qPCR) in BRCA1 deficient and BRCA1 proficient cells.

Quantitative real time PCR (RT-qPCR) using QuantiTect® Primers (Qiagen, UK) was performed as described in section 2.3.8.5 for the selected DNA repair factors; BRCA1, APE1, XRCC1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs (discussed previously section 3.1.2.). mRNA expression is shown in Table 3.8.
for breast cancer cell lines and mRNA expression for HeLa cell lines is shown in Table 3.9.

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Table 3.8. Quantitative real time PCR (RT-qPCR) for multiple DNA repair genes in BRCA1 deficient cells.

DNA repair mRNA expression in MDA-MB-436 compared to MCF7 cells. Statistical significance determined using t test, with p values less than 0.05.

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</tr>
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</table>

Table 3.9. Quantitative real time PCR (RT-qPCR) for multiple DNA repair genes in BRCA1 deficient cells.

DNA repair mRNA expression in BRCA1 HeLa cells compared to Control HeLa cells. Statistical significance determined using t test, with p values less than 0.05.
3.3.7. Quantification of DNA repair mRNA and protein expression in BRCA1 deficient and BRCA1 proficient cells.

Relative quantification of mRNA and protein expression for XRCC1, APE1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs was performed. BRCA1 HeLa cells, expressed relatively low DNA repair factors compared to Control HeLa cells, as demonstrated in Figure 3.14 and Figure 3.16. MDA-MB-436 cells expressed relatively low DNA repair factors compared to MCF7 cells, as demonstrated in Figure 3.15 and Figure 3.17.

Western blot results were analysed using Image Studio (Li-Cor, Lincoln, USA) as described in section 2.3.9.8. RT-qPCR results were analysed as described in section 2.3.8.6. These assays were performed in two or more independent experiments. The student t-test was used to investigate the difference in expression of DNA repair proteins between BRCA1 proficient and BRCA1 deficient cell lines.
Figure 1: Bar charts showing the fold change in protein and mRNA expression levels for XRCC1, APE1, and Lig3 in control and BRCA1 Hela cells.

(A) Protein expression

(B) mRNA expression
Figure 3.14. Quantification of Base Excision Repair proteins expression (left) and quantitative real time PCR (RT-qPCR) showing mRNA expression (right) in BRCA1 deficient and proficient HeLa cell lines.

(A) Showing protein expression, values plotted are means ± SD of the fold-change (ratio of protein/ß-actin normalized to control) and (B) mRNA expression, values plotted are means ± SD of the fold-change (ratio of mRNA/GAPDH normalized to control). * p<0.05, ** p<0.01 and *** p<0.001, p value was assessed by t-test compared to Control HeLa cell line.

Graphs were produced and statistical analysis performed using GraphPad prism.
Figure 3.15. Quantification of Base Excision Repair proteins expression (left) and quantitative real time PCR (RT-qPCR) showing mRNA expression (right) in BRCA1 deficient and proficient breast cancer cell lines.

(A) Showing protein expression, values plotted are means ± SD of the fold-change (ratio of protein/β-actin normalized to control) and (B) mRNA expression, values plotted are means ± SD of the fold-change (ratio of mRNA/GAPDH normalized to control). * p<0.05, ** p<0.01 and *** p<0.001, p value was assessed by t-test compared to MCF7 cell line. Graphs were produced and statistical analysis performed using GraphPad prism.
Figure 3.16. Quantification of ATM and DNA-PKcs protein (left) and quantitative real time PCR (RT-qPCR) showing mRNA expression (right) in BRCA1 deficient and proficient HeLa cell lines.

(A) Showing protein expression, values plotted are means ± SD of the fold-change (ratio of protein/ß-actin normalized to control) and (B) mRNA expression, values plotted are means ± SD of the fold-change (ratio of mRNA/GAPDH normalized to control). * p<0.05, p value was assessed by t-test compared to Control HeLa cell line. Graphs were produced and statistical analysis performed using GraphPad prism.
Figure 3.17. Quantification of ATM and DNA-PKcs protein expression (left) and quantitative real time PCR (RT-qPCR) showing mRNA expression (right) in BRCA1 deficient and proficient breast cancer cell lines.

(A) Showing protein expression, values plotted are means ± SD of the fold-change (ratio of protein/β-actin normalized to control) and (B) mRNA expression, values plotted are means ± SD of the fold-change (ratio of mRNA/GAPDH normalized to control). * p<0.05, ** p<0.01 and *** p<0.001, p value was assessed by t-test compared to MCF7 cell line. Graphs were produced and statistical analysis performed using GraphPad prism.
3.3.8. Growth inhibition MTS assay in response to MMS

BRCA1 deficient cells demonstrate low BER factors as shown. To test whether low BER expression confers phenotypic consequence, MMS (alkylating agent) in BRCA1 proficient and deficient cells was investigated. MTS assay was used to determine cell growth inhibition in response to MMS treatment (0.00 µM to 800 µM) for 72 hours in BRCA1 mutated MDA-MB-436 cells and control MCF7 cells. As shown in Figure 3.18, BRCA1 mutated MDA-MB-436 cells were more sensitive to MMS than control MCF7 cells. The data concurs with a previous study by Masaoka et al, who demonstrated similar MMS sensitivity in BRCA1 deficient cells (Masaoka et al., 2013).

**Figure 3.18.** MMS treatment in BRCA1 deficient and proficient cells.

BRCA1 deficient cells exhibited decreased cell growth after exposure for 72 hours to MMS, compared to BRCA1 proficient cell line. Data are shown as the mean and SD values for each concentration from 2 independent experiments.
3.4. Discussion and conclusions

3.4.1. Loss of BRCA1 expression links to a global reduction in mRNA and proteins involved in Base excision repair pathway

In a recent publication, BRCA1 has been suggested to stimulate the activity of three BER enzymes; OGG1 (8-oxoguanine DNA glycosylase), NTH1 (homolog of endonuclease III), and APE1 (apex nuclease 1). These enzymes mediate repair of three signature oxidative lesions; 8-oxoguanine, thymine glycol and abasic sites, respectively (Saha et al., 2010). A more recent study supports these findings, and has confirmed down-regulation of APE1, NTH1, OGG1 as well as XRCC1 in BRCA1 mutated and sporadic breast cancers (De Summa et al., 2014). XRCC1 protein associates with Pol β, and Lig3, to form a complex that repairs the single strand DNA breaks generated during the BER process (Saha et al., 2010). Other studies suggest a cross talk between BRCA1 and other BER genes e.g. Pol β and SMUG1 (Masaoka et al., 2013, Abdel-Fatah et al., 2013), highlighting additional BER factors that may be associated with BRCA1 expression. For these reasons in the current study BER factors; APE1, XRCC1, SMUG1, Pol β and Lig3 were investigated. To achieve this, the RT² Profiler™ Array was conducted first to investigate 84 DNA repair genes in BRCA1 deficient cell lines as compared to BRCA1 proficient cell lines. However, the establishment of a statistical significance threshold that adequately controls the proportion of false-positive findings is not a trivial issue. Therefore, due to the sample size and number of arrays statistical significance was determined using a Bonferroni correction to adjust for multiple testing, with corrected p values set at less than 0.05 (threshold p <
0.000595 = 0.05/84). Thus, the current RT² Profiler array results did not always reach statistical significance and further independent replicates are required. Further assessment of the results was carried out using quantitative real time PCR (RT-qPCR) for the selected DNA repair genes as discussed above. Finally, the mRNA expression of the selected genes was compared with protein expression using western blot.

In the current study, expression of several BER mRNAs were found to be down-regulated in both BRCA1 mutant/knockdown cell lines (see Table 3.3); low mRNA expression was demonstrated for APE1, OGG1 and NTHL1 genes which was consistent with the results published by Saha et al. mRNAs of the selected BER genes also showed down-regulation. Interestingly many other BER mRNAs showed down-regulation including; PARP1, PARP2, PARP3, NEIL1, NEIL3, UNG, FEN1, MUTYH and CCNO. Targeted assessment in individual primer assays may be useful to further assess these genes. Conversely, individual RT-qPCR analysis for XRCC1, APE1, LIG3, SMUG1 and Pol β mRNA expression demonstrated significant reduction in both mutant/knockdown BRCA1 cell lines as compared to proficient BRCA1 cell lines (see Table 3.8 and Table 3.9). This reduction was linked to reduction in protein expression (see Figure 3.14 and Figure 3.15). These results suggest that there may be impairment in key BER factors in BRCA1 deficient cells, which may cause different types of unrepaired single strand breaks.

3.4.1.1. BRCA1 deficient cell lines are hypersensitive to MMS

In this section of the study the efficacy of the alkylating agent in DNA damage formation in BRCA1 deficient and proficient cells was investigated.
Methyl methane sulphonate (MMS) is an alkylating agent, damage induced by MMS is repaired by BER pathway. MMS acts by methylating DNA on N7-deoxyguanosine and N3-deoxyadenosine, creating adducts that are substrates for BER. At high concentrations of MMS, the frequency of methylation events overwhelms BER capacity and induce replication fork stalling, resulting in cytotoxic levels of DSB formation (Stevnsner et al., 1993, Masaoka et al., 2013). A number of studies have shown that BRCA1/BER deficient cell lines were hypersensitive to MMS compared to their wild type matched controls (Bridge et al., 2005, Masaoka et al., 2013). In this study MMS was confirmed to be selectively toxic to BRCA1/BER mutated human cancer cell line (MDA-MB-436) compared to MCF7 proficient cell line by using MTS assay. This result suggests impairment in BER. However, measuring BER efficiency should be conducted in BRCA1 deficient cell lines in the future studies.

3.4.2. Loss of BRCA1 expression is associated with reduction in expression of Homologous recombination factors

The first report of a role for BRCA1 in genomic stability and double strand break repair, by Scully and colleagues, suggested a functional interaction between BRCA1 and RAD51 in both the meiotic and mitotic cell cycles. This in turn suggests an important role for BRCA1 in controlling genomic stability (Scully et al., 1997). This association was corroborated by a later report by Chen and colleagues, who observed the presence of both BRCA1 and RAD51 on the damaged DNA site in meiotic cells (Chen et al., 1998). This finding was followed by the first functional evidence for the role of BRCA1 in HR repair pathway through its direct interaction with RAD51 (Moynahan et al., 1999).
Subsequent observations that agents causing recombinogenic lesions like cisplatinum and ionizing radiation, do not induce formation of RAD51 foci in BRCA1 deficient cells, suggesting that disruption of BRCA1 was associated with markedly reduced expression of RAD51 (Bhattacharyya et al., 2000, Bindra et al., 2004). In the current study, reduced BRCA1 expression in mutant/knockdown cell lines was associated with a reduction in RAD51 expression at mRNA (see Table 3.4) in both mutant/knockdown cell lines. Several other HR mRNA expressions were also observed to be down-regulated in both mutant/knockdown cell lines; low mRNA expression was demonstrated for RAD51 related family (RAD51B, RAD51C and RAD51D), RAD50, RAD52, XRCC3, BRCA2 and DMC1. In addition, RAD54L and XRCC2 were also down-regulated in both mutant/knockdown cell lines. Recently, RAD21 has been suggested as a potential BRCA1 mutation status dependent predictive and prognostic marker in familial breast cancer (Yan et al., 2012). In the present study low mRNA expression of RAD21 in both mutant/knockdown cell lines was demonstrated. ATM is believed to be the first sensor in DSB and a key player in HR pathway. ATM activates multiple genes involved in HR repair, cell cycle arrest and apoptosis. Substrates for ATM phosphorylation include; BRCA1, p53, CHK2, histone H2AX and MRN complex (Bakkenist and Kastan, 2004). In this study low mRNA expression of ATM in BRCA1 mutant/knockdown cell lines was demonstrated using RT²-PCR array. Then ATM was assessed in individual RT-qPCR which showed significant mRNA reduction in both mutant/knockdown BRCA1 cell lines compared to proficient BRCA1 cell lines (see Table 3.8 and Table 3.9). Gene expression was associated with protein expression for ATM (see Figure 3.16 and Figure 3.17).
Taken together, these results suggest that BRCA1 deficiency may associate with a global defect in HR repair expression. However, individual primer assays may be useful to further assess these genes.

3.4.3. Loss of BRCA1 expression is associated with reduction in expression of Non-homologous end joining factors

BRCA1 interacts with the Rad50/Mre11/Nbs1 complex that occupies a central role in DNA double-strand break repair mediated by HR and NHEJ. Zhong and colleagues suggested a BRCA1 role in NHEJ. They investigated Brca1-deficient mouse embryonic fibroblasts cell extracts and found that it exhibit reduced end-joining activity independent of the endogenous protein amounts of DNA LIG4, KU80 (known as XRCC5), and KU70 (known as XRCC6) (Zhong et al., 2002). In the current study, reduced BRCA1 expression in mutant/knockdown cell lines was associated with a reduction in LIG4, XRCC5 and XRCC6 expression at mRNA confirming Zhong et al results (see Table 3.5). In addition, XRCC4 show down-regulation at the mRNA expression in mutant/knockdown cell lines. DNA-PKcs is known to have a critical role in NHEJ pathway. Recently, Shang and colleagues reported that DNA-PKcs is the upstream regulator of the CHK2–BRCA1 pathway (Shang et al., 2014). In this study low mRNA expression of DNA-PKcs was demonstrated in mutant/knockdown cell lines by RT²-PCR array (see Table 3.5). Then DNA-PKcs was assessed in individual RT-qPCR which showed significant mRNA reduction in both mutant/knockdown BRCA1 cell lines compared to proficient BRCA1 cell lines (see Table 3.8 and Table 3.9). Also,
gene expression was consistent with protein expression in DNA-PKcs (see Figure 3.16 and Figure 3.17). Taken together, these results suggest that BRCA1 deficiency is associated with impairment in NHEJ expression.

3.4.4. Loss of BRCA1 expression is associated with reduction in expression of other DNA repair factors

BRCA1 has been suggested to control Nucleotide excision repair (NER), through the p53 protein which regulates NER by transcriptional regulation of genes involved in the recognition of adducts in genomic DNA. Loss of p53 function results in deficient global genomic repair (GGR), a subset of NER that targets and removes lesions from the whole genome. BRCA1 specifically enhances the GGR pathway, independent of p53, and can induce p53-independent expression of the NER genes XPC, DDB2, and GADD45 (Hartman and Ford, 2002). In the current study, reduced BRCA1 expression in mutant/knockdown cell lines was associated with a reduction in XPC and DDB2 expression at mRNA consistent with the data published by Hartman and Ford (see Table 3.6). In addition, a panel of NER factors also shows down-regulation at the mRNA expression. For instance; CCNH, RPA3, ERCC1, ERCC4, ERCC6, ERCC8, BRIP1, LIG1, ATXN3, RAD23A, RPA1, CDK7, DDB1, DDB2, PNKP, POLL, SLK, ERCC2, ERCC3, ERCC5, XAB2 and XPA.

The role of BRCA1 in MMR pathway is still not understood, however recently a report by Song and colleagues demonstrated that mutation in BRCA1 was followed by mutation in MSH6 and MSH2 (Song et al., 2014). In the current
study, reduced BRCA1 expression in mutant/knockdown cell lines was associated with a reduction in $MSH6$ and $MSH2$ expression at mRNA supporting Song et al results (see Table 3.7). Moreover, a panel of MMR mRNA expressions were observed to be down-regulated in both mutant/knockdown cell lines including; $POLD3$, $PMS1$, $PMS2$, $MLH1$, $MLH3$, $MSH3$, $MSH4$, $MSH5$, $MPG$ and $TREX1$. These results suggest that BRCA1 deficiency may associate with a global defect in NER and MMR expression. However, further investigation is needed to measure the efficiency of these pathways.

Taken together, the main aim of this chapter was to investigate a potential impairment in DNA repair pathways within BRCA1 deficient cells. To achieve this, RT² Profiler™ Array was conducted first to investigate 84 DNA repair genes in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines. Data demonstrated down-regulation in several DNA repair mRNAs however; the results did not always give statistical significance. Further assessment of the results was carried out using quantitative real time PCR (RT-qPCR) for selected DNA repair genes ($APE1$, $XRCC1$, $SMUG1$, $Pol \beta$, $Lig3$, $ATM$ and $DNA-PKcs$) which had previously been shown to have a potential link with BRCA1 (Abdel-Fatah et al., 2013, Masaoka et al., 2013, De Summa et al., 2014). Data for individual primers demonstrated statistically significant down-regulation of the selected DNA repair genes. This highlights the difference between the two PCR methods, where both showed consistent down-regulation but data was not statistically significant with the RT² Profiler™ Array. This can have two explanations. Firstly the efficiency, specificity and sensitivity of
primers can differ. Secondly, and more importantly, the average threshold cycle (Ct) should be in the range of 15 to 35 which is considered as a strong positive reaction. Ct values greater than 35 are considered as a weak reaction. Ct value is negatively associated with the concentration of nucleic acids detected. Ct values are classified in subsequent categories of Ct < 25, Ct 25-35 and Ct 35-40. In the present study both Ct values for both PCR methods were in the range of 18 to 33. However the percentage distributions of Ct values were different. In the RT² Profiler™ Array 66% of Ct values were < 25 and 34% between 25 and 35, whereas by RT-qPCR around 90% of Ct values were 25-35 and 10% were < 25. This suggests that the reason for having consistent but not statistically significant results in RT² Profiler™ Array was because of the distribution of the Ct values. It is worth mentioning that a previous study demonstrated similar results when they compared a quantitative RT-PCR array (qPCR-array) with an oligonucleotide microchip (microarray) (Chen et al., 2009). For future work this problem can be avoided by further dilution of the cDNA.

3.4.5. Conclusions

These in vitro studies suggested that BRCA1 has been linked to a possible defect in BER, HR, NHEJ, NER and MMR, via mechanisms that are not fully understood. This chapter described the assessment of DNA repair factors expressions in BRCA1 mutant/knockdown cell lines. Selected DNA repair factors (APE1, XRCC1, SMUG1, Pol ß, Lig3, ATM and DNA-PKcs) were observed to be down-regulated at the mRNA and/or protein expression. Taken together, these results suggest that BRCA1 deficiency may be associated with a global defect in DNA repair.
Chapter 4

Targeting BRCA1 deficiency for personalized therapy using ATM inhibitors (KU55933 & KU60019)
4. Targeting BRCA1 deficiency in breast cancer for personalized therapy using ATM inhibitors (KU55933 & KU60019)

4.1. Introduction

BRCA1 facilitates the efficient resolution of DNA double-strand breaks (DSBs) through homologous recombination (HR) (Huen et al., 2010, Caestecker and Van de Walle, 2013). Cells lacking functional BRCA1 protein have impaired HR, and thus are suggested to depend on the more error-prone non-homologous end joining (NHEJ) pathway leading to chromosomal instability that drive breast cancer development (Huen et al., 2010). BER pathway is critical for processing DNA damage caused by alkylation, oxidation, ring saturation, single strand breaks and base deamination (Dianov and Hubscher, 2013).

PARP1 (poly [ADP-ribose] polymerase 1) plays a role in single strand break repair (SSBR), a BER related pathway (Langelier and Pascal, 2013). The DNA repair intermediates generated during SSBR/BER, if unrepaired, may get converted to toxic DSBs (Dianov and Hubscher, 2013). PARP1 inhibition is synthetically lethal in BRCA deficient cells (Farmer et al., 2005, Bryant et al., 2005). Although it should be noted that recent studies illustrated some problems with PARP inhibition (Edwards et al., 2008, Montoni et al., 2013, Fojo and Bates, 2013). For instance, PARP family of enzymes consist of at least 17 members, each one of them has a different structure and function (Rouleau et al., 2010, Mangerich and Burkle, 2011, Underhill et al., 2011). So there are issues with specificity of current inhibitions under development. Moreover, a recent study evaluated a series of 185 inhibitors of PARP,
including the best-known inhibitors being tested clinically such as Olaparib, ABT-888 and Rucaparib, for the specificity to bind to the catalytic domains of 13 of 17 human PARP family members. In this study they determined that the majority of the inhibitors bind to multiple targets (Wahlberg et al., 2012).

Emerging studies suggest a cross talk between BRCA1 and BER factors. BRCA1 mutated and basal-like breast cancer cells were found to be sensitive to oxidative DNA damage induced by hydrogen peroxide (H$_2$O$_2$) treatment. The increased sensitivity was associated with defective BER as assessed by cell based BER assay in BRCA1 deficient cells (Alli et al., 2009). In a more recent study, BRCA1 deficient cells were sensitive to alkylating agent Methyl methane sulphonate (MMS) and functional interaction between Pol β and BRCA1 was demonstrated in that study (Masaoka et al., 2013) implying a potential role for Pol β in BRCA1 mediated DSB repair. In addition, BRCA1 has also been shown to be involved in the transcriptional regulation of BER factor such as OGG1, NTH1, XRCC1 and APE1 (Saha et al., 2010, De Summa et al., 2014).

Ataxia telangiectasia mutated (ATM) is a key component of HR pathway. ATM is a serine/threonine kinase that activates over a hundred proteins involved in DNA damage response, DNA repair, cell cycle regulation, apoptosis and other pathways (Kurz and Lees-Miller, 2004, Chaudhary and Al-Baradie, 2014). The ATM signalling pathway is activated following DSBs. ATM is recruited to the damaged sites, however it is not clear if ATM interacts with damaged DNA directly or indirectly. It was previously thought that ATM activation primarily requires direct binding to MRN complex which is a sensor of DSBs (Chaudhary and Al-Baradie, 2014). However recent studies have
shown that PARP1 may control ATM phosphorylation and mediates the earliest recruitment of MRE11 and NBS1 to DNA damage (Rupnik et al., 2008, Haince et al., 2008, Haince et al., 2007).

4.1.1. Rationale for study

Data presented in Chapter 3 of this thesis suggested that BRCA1 deficiency is associated with reduced expression of several factors involved in BER. In this chapter it was investigated if such BRCA1-BER deficient cells could be targeted by blockade of ATM for personalized therapy.

4.2. Aims

The aims of this study were as follows:

1. To investigate the effect of ATM inhibitors (KU55933 and KU60019) on cell viability and growth using clonogenic cell survival and MTS assays.

2. To confirm the effect of PARP inhibitors (NU1025 and 3-Aminobenzamide) on cell viability and growth using clonogenic cell survival and MTS assays.

3. To test the hypothesis that DNA double strand breaks accumulate in cells after exposure to ATM inhibitors in BRCA1 deficient cells.

4. To determine the effect of ATM inhibitors on cell cycle in BRCA1 deficient cells.

5. To explore induction of apoptosis in BRCA1 deficient cells after treatment with ATM inhibitors.
4.3. Results

4.3.1. Clonogenic cell survival and Growth inhibition in response to ATM inhibitors (KU55933 & KU60019)

The plating efficiencies (which is a measure of a cell's ability to form colonies) of the HeLa cell lines were around 60% to 98% in this study, this is consistent with the previous published literature which showed the high plating efficiency of HeLa cells (Puck and Fisher, 1956, Rafferty, 1985). On the other hand, previous studies showed plating efficiency of breast cancer cell lines around 29% to 55% (Mukherjee et al., 2005, Mukherjee and Martin, 2006, Ware et al., 2007, Zhang et al., 2014, Karasawa et al., 2014, Duangmano et al., 2012). In this present study plating efficiencies for MCF7 and MDA-MB-436 cells in clonogenic assay were around (45% to 58%) and (24% to 54%), respectively. It is possible that one of the reasons for this slight increase/decrease in plating efficiencies is due to the specific culture conditions (e.g. cell culture media and FBS concentrations) and cell seeding number in different studies.

Clonogenic survival assays were performed to test whether BRCA1 deficient cells are sensitive to ATM inhibition. BRCA1 mutated (MDA-MB-436) and knockdown cell lines (BRCA1 HeLa cells) along with BRCA1 proficient MCF7 cells and Control HeLa cell lines were treated with increasing concentrations of KU55933 ($\text{C}_2\text{H}_1\text{H}_7\text{NO}_3\text{S}_2$) which is a potent, selective and competitive ATM kinase inhibitor. Results are shown in Figure 4.1. IC$_{50}$ values were calculated using nonlinear regression, with the proviso that in BRCA1 proficient cell lines (Control HeLa and MCF7) cytotoxicity did not
approach 50% in some cases, thus impairing extrapolation. Clonogenic assay demonstrated that BRCA1 HeLa cells, are more sensitive to KU55933 compared to Control HeLa cells (mean IC₅₀= 2.911 ± 0.21 vs. mean IC₅₀= 9.160 ± 1.6). Similarly, increased sensitivity was also demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells (mean IC₅₀= 6.39 ± 0.43 vs. mean IC₅₀= 9.23 ± 0.48).

Figure 4.1. Clonogenic survival assay of ATM inhibitor (KU55933).

Cell lines with reduced BRCA1 expression exhibited decreased colony forming ability after treatment with KU55933 inhibitor for 14 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 85.83% ± 3.433%; BRCA1 HeLa- 74.10% ± 4.054% (B) MCF7- 45.71% ± 4.69%; MDA-MB-436- 36.73% ± 3.67%. Clonogenic data are shown as the mean and SD values for each concentration from ≥ 3 independent experiments.

In general, increasing concentrations of KU55933 inhibitor reduced the colony forming ability in BRCA1 deficient cells compared to the BRCA1 proficient cells. To validate further, an alternative clonogenic assay protocol was performed, where cells were treated with ATM inhibitor for 24 hours then
culture media was replaced with fresh media and the remaining assay was conducted as described in 2.3.4.3. Results are illustrated in Figure 4.2.

![Figure 4.2. Clonogenic survival assay of inhibitor KU55933 (24 hours treatment protocol).](image)

Cell lines with reduced BRCA1 expression exhibited decreased colony forming ability after treatment with KU55933 inhibitor for 24 hours. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 83.30% ± 3.30%; BRCA1 HeLa- 86.80% ± 6.80% (B) MCF7- 48.78% ± 4.33%; MDA-MB-436- 53.62% ± 1.94%. Clonogenic data are shown as the mean and SD values for each concentration from 2 independent experiments.

The alternative clonogenic assay results demonstrated that BRCA1 HeLa cells are more sensitive to KU55933 than Control HeLa cells (mean IC₅₀= 5 ± 0.24 vs. IC₅₀ above the maximum level tested). Similarly, sensitivity was demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells (mean IC₅₀= 4.45 ± 0.23 vs. IC₅₀ above the maximum level tested). Both the 24 hour KU55933 treatment protocol and the 14 day treatment protocol, provide evidence that increasing concentration of KU55933
inhibitor reduces the colony forming ability of BRCA1 deficient cell lines compared to the BRCA1 proficient cell lines. The 14 day treatment protocol was used for all further clonogenic assays described in the rest of the thesis.

For additional validation, the proliferation assay (MTS assay) was used to determine cell growth in response to KU55933. As shown in Figure 4.3, BRCA1 deficient HeLa cells were more sensitive to KU55933 than the Control HeLa cells. Similar sensitivity was demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells.

Figure 4.3. MTS cell growth inhibition assays of ATM inhibitor (KU55933).
Cell lines with reduced BRCA1 expression exhibited decreased cell growth after treatment with KU55933 inhibitor for 6 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
For further validation, KU60019 \((C_{30}H_{33}N_{3}O_{5}S)\), another ATM potent kinase inhibitor, was investigated (Golding et al., 2009). As shown in Figure 4.4 clonogenic cell survival assays, BRCA1 HeLa cells are more sensitive to KU60019 than Control HeLa cells. Similar sensitivity was also demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells. Taken together, these data provide strong evidence that BRCA1 deficient cells are sensitive to ATM inhibition.

**Figure 4.4. Clonogenic survival assay of ATM inhibitor (KU60019).**

Cell lines with reduced BRCA1 expression exhibited decreased colony forming ability after treatment with KU60019 inhibitor for 14 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 87.75\% \pm 5.75\%; BRCA1 HeLa- 67.00\% \pm 17.00\%; (B) MCF7- 61.36\% \pm 4.21\%; MDA-MB-436- 46.00\% \pm 4.25\%. Clonogenic data are shown as the mean and SD values for each concentration from \(\geq 2\) independent experiments.
4.3.2. Growth inhibition and clonogenic cell survival in response to PARP inhibitors

MTS cell growth assays were performed using NU1025 (C₉H₈N₂O₂) a well characterised inhibitor of PARP (Bowman et al., 1998, Griffin et al., 1998, Sultana et al., 2012), as a positive control drug to induce lethality in this model system. Figure 4.5 demonstrates that BRCA1 HeLa cells were more sensitive to NU1025 than the Control HeLa cells. Similar sensitivity was also demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells. This result is consistent with previous studies that demonstrated similar sensitivity in BRCA1 deficient cells to NU1025 (Bryant et al., 2005, Sultana et al., 2012).

![Graph A: MTS cell growth inhibition assays of PARP inhibitor (NU1025).](image1)

![Graph B: MTS cell growth inhibition assays of PARP inhibitor (NU1025).](image2)

Figure 4.5. MTS cell growth inhibition assays of PARP inhibitor (NU1025). Cell lines with reduced BRCA1 expression exhibited decreased cell growth after treatment with NU1025 inhibitor for 6 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 experiments.
For additional validation, 3-Aminobenzamide (C₇H₈N₂O) another PARP inhibitor, was investigated (Purnell and Whish, 1980). As shown in Figure 4.6, BRCA1 HeLa cells are more sensitive to 3-Aminobenzamide compared to Control HeLa cells. Similar sensitivity was also demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells. This result is consistent with a previous study that showed sensitivity in BRCA1 deficient cells to 3- Aminobenzamide (Bryant et al., 2005). Functional analyses in cells were then carried out to investigate ATM inhibition in BRCA1 deficient cells.

![Figure 4.6. Clonogenic survival assay of PARP inhibitor (3-Aminobenzamide).](image)

Cell lines with reduced BRCA1 expression exhibited significantly decreased colony forming ability after treatment with 3-Aminobenzamide inhibitor for 14 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cell line compared to Control HeLa. (B) MDA-MB-436 compared to MCF7. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 98.17% ± 1.83%; BRCA1 HeLa- 98.33% ± 0.33% (B) MCF7- 64.06% ± 0.94%; MDA-MB-436- 64.61% ± 0.72%. Clonogenic data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
4.3.3. γH2AX focus immunofluorescence microscopy assay

To determine whether DSBs accumulate after treatment with KU55933, DSBs were measured using the γH2AX immunofluorescence microscopy assay. As described in Chapter 2 section 2.3.5, cells were treated with 10 µM of KU55933 for 24 hours and 48 hours. As shown in Figure 4.7, after 48 hours of treatment, a significant increase in foci formation was observed in BRCA1 deficient cells. Cells were then treated with 5 µM of KU55933 for 48 hours as shown in Figure 4.8, but no significant accumulation of γH2AX foci in BRCA1 deficient cells was observed. The data support a dose response effect of ATM inhibition in BRCA1 deficient cells.

Figure 4.7. Representative photomicrographic of Immunofluorescence microscopy assay following KU55933 treatment for 24 hours and 48 hours.

BRCA1 HeLa cells and Control HeLa cells, were treated with 10 µM for 24 and 48 hours, then fixed and stained with γH2AX primary and FITC-conjugated secondary antibodies. DAPI was used as a nuclear stained showing in blue and γH2AX foci detected fluoresce in green.

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γH2AX formation in response to ATM inhibitors

BRCA1 proficient and BRCA1 deficient cell lines following treatment with 10 µM of KU55933 for 48 hours are shown in Figure 4.9. The data demonstrates a statistically significant increase in the accumulation of γH2AX foci in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (mean 66.50% ± 6.5 vs. mean 20.50% ± 3.5; p = 0.02) and compared to corresponding treated Control HeLa cells (mean 8.00% ± 3.00; p = 0.014).
Similar results were also observed in treated MDA-MB-436 cells in which accumulation of γH2AX foci were significantly higher compared to untreated MDA-MB-436 cells (mean 88.00% ± 4.0 vs. 5.50% ± 0.50; p = 0.002) and compared to corresponding treated MCF7 cells (mean 4.50% ± 1.50; p = 0.0026).

For further validation KU60019, another ATM inhibitor, was investigated. Treatment with 1µM of KU60019 produced similar results as with KU55933. As shown in Figure 4.10, a statistically significant increase in the accumulation of γH2AX foci in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (mean 41.50% ± 4.5 vs. 10% ± 1.0; p = 0.02), and compared to corresponding treated Control HeLa cells (mean 5.00% ± 1.00; p = 0.015)

**Figure 4.9. Immunofluorescence microscopy staining for γH2AX foci formation following KU55933 treatment for 48 hours.**

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 3 independent experiments. * p<0.05 and ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
were observed. Similar results were also observed in treated MDA-MB-436 cells in which accumulation of γH2AX foci was significantly higher compared to untreated MDA-MB-436 cells (mean 71.50% ± 2.5 vs. 17.50% ± 1.5; p = 0.002), and compared to the corresponding treated MCF7 cells (mean 37.50% ± 1.5; p = 0.007). Although MDA-MB-436 cells were more sensitive to KU60019 inhibitor, MCF7 cells showed a similar change (from 10% to 37.50%). This result infers that KU60019 inhibitor maybe toxic to MCF7.

Figure 4.10. Immunofluorescence microscopy staining for γH2AX foci formation following KU60019 treatment for 48 hours.

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05 and ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
4.3.4. Cell cycle analysis by propidium iodide flow cytometry after treatment with ATM inhibitors

DSB formation induces cell cycle checkpoint activation to trigger cell cycle arrest to allow DNA repair, if the damage is extensive then this process can initiate apoptosis (Kurz and Lees-Miller, 2004). To evaluate cell cycle progression, Propidium iodide (PI) flow cytometry was used as described in Chapter 2, section 2.3.6.

Cells were treated for 24 hours and 48 hours with 10 µM or 20 µM of KU55933 (Figure 4.11). Although no significant difference was observed after 24 hours, at 48 hours in BRCA1 deficient cells treated with 10 µM KU55933 exhibited a significant increase in G2/M phase arrest compared to control cells was observed. On the other hand, at 20 µM concentration conducts apoptosis was evident, illustrated by the loss of cells count.
Results for BRCA1 proficient and BRCA1 deficient cells following treatment with 10 µM (KU55933) for 48 hours are shown in Figure 4.12. The data demonstrate a statistically significant G2/M arrest in treated BRCA1 HeLa cells compared with untreated BRCA1 HeLa cells (mean 26.60% ± 0.40 vs. 20.50% ± 1.50; p = 0.05), and compared to corresponding treated Control

Figure 4.11. Representative graphs of flow cytometric cell cycle analysis following KU55933 treatment.

BRCA1 HeLa cells and Control HeLa cells were treated with 10 µM and 20 µM for 24 hours and 48 hours. Y axis represents cell count and propidium iodides detected fluorescence is shown on the X axis.
HeLa cells (mean 21.50% ± 0.50; p = 0.0154). Results of treated MDA-MB-436 cells demonstrate an increase in G2/M arrest compared to MDA-MB-436 untreated cells, though this is not significant (mean 32.45% ± 2.75 vs. mean 27.60% ± 2.9; p = 0.3). The observed G2/M arrest was significantly higher in treated MDA-MB-436 cells compared to treated MCF7 cells (mean 20.30% ± 0.50; p = 0.0491). Conversely, Control HeLa and MCF7 cells did not show the same increase (i.e. ~5%) as with BRCA1 deficient cells. (Control HeLa and MCF7 cells, G2/M arrest mean 21.50% to 22.55% and 19% to 20.30%, respectively).

Figure 4.12. Flow cytometric cell cycle analysis following KU55933 treatment for 48 hours.

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 3 independent experiments. * p≤ 0.05, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.

For further validation KU60019, another ATM inhibitor, was investigated. Treatment with 1µM of KU60019 produced similar results as with KU55933.
As shown in Figure 4.13, a statistically significant G2/M arrest in treated BRCA1 HeLa cells was demonstrated as compared to untreated BRCA1 HeLa cells (mean 31.55% ± 0.55 vs. 26.9% ± 0.2; p = 0.015), and as compared to the corresponding treated Control HeLa cells (mean 23.80% ± 0.10; p = 0.005). A similar, although not statistically significant, G2/M arrest was also observed in treated MDA-MB-436 cells compared to untreated MDA-MB-436 cells (mean 34.60% ± 0.40 vs. 28.30% ± 1.7; p = 0.06). A statistically significant G2/M was observed in treated MDA-MB-436 cells as compared to treated MCF7 cells (mean 25.15% ± 1.15; p = 0.0162). In contrast, Control HeLa cells and MCF7 did not show the same increase (G2/M arrest mean 19.5% to 23.8% and 25.70% to 25.15%, respectively).

Figure 4.13. Flow cytometric cell cycle analysis following KU60019 treatment for 48 hours.

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05 and ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
4.3.5. Apoptosis detection by annexin V-FITC flow cytometry after treatment with ATM inhibitors

Annexin V was used in conjunction with a propidium iodide (PI) for identification of late apoptotic cells as described in Chapter 2, section 2.3.7. Viable cells with intact membranes are negative (reject) to PI, whereas dead cells with damaged membranes are positive (permeable) to PI. Therefore, viable cells are both Annexin V and PI negative, early apoptosis cells are Annexin V positive and PI negative, late apoptosis cells are Annexin V and PI positive, necrotic cells are Annexin V negative and PI positive (Koopman et al., 1994, Riccardi and Nicoletti, 2006).

Cells were treated for 24 hours and 48 hours with 10 μM of KU55933. As shown in Figure 4.14, although no significant increase in apoptosis after 24 hours was evident, at 48 hours in BRCA1 deficient cells treated with 10 μM of KU55933 showed significant increase in apoptosis compared to control cells was observed.
Results for BRCA1 proficient and BRCA1 deficient cells following treatment with 10 µM (KU55933) for 48 hours are shown in Figure 4.15, data were normalised against baseline apoptotic fraction (untreated cells) to determine the percentage increase in apoptosis. The data demonstrate a statistically significant increase in apoptosis with treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (mean 11.12% vs. 3.6%; p =
Similar results were also observed in treated MDA-MB-436 cells, in which a significant increase in apoptosis as compared to untreated MDA-MB-436 cells was observed (mean 13.1% vs. 4.8%; p < 0.0001). On the other hand, Control HeLa and MCF7 cells did not show the same rate of increase (mean 3.9% to 5.7% and 3.17% to 7.5%, respectively).

![Graph](image)

**Figure 4.15. Apoptosis detection by annexin V-FITC flow cytometry in KU55933 treated cells.**

Treatment with KU55933 inhibitor was associated with a significant increase in apoptotic cells. Data were normalised against baseline apoptotic fraction to determine the percentage increase in apoptosis. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 3 experiments. * p<0.05 and ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.

For further validation KU60019, another ATM inhibitor, was investigated. Treatment with 1µM of KU60019 produced similar results as with KU55933. As shown in Figure 4.16, the data demonstrates a statistically significant increase in apoptosis in treated BRCA1 HeLa cells compared to the
corresponding untreated BRCA1 HeLa cells (mean 9.56% vs. 5.53%; p = 0.037). A similar increase was also observed in treated MDA-MB-436 cells as compared to untreated MDA-MB-436 cells (mean 8.99% vs. 4.68%; though was not significant). Conversely, Control HeLa and MCF7 cells did not show the same rate of increase (mean 4% to 3.8% and 3.97% to 5.9%, respectively).

Figure 4.16. Apoptosis detection by annexin V-FITC flow cytometry in KU60019 treated cells.

Treatment with KU60019 inhibitor was associated with an increase in apoptotic cells. Data were normalised against baseline apoptotic fraction to determine the percentage increase in apoptosis. (A) BRCA1 HeLa cells compared to Control HeLa cells. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 3 experiments. ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
4.4. Discussion and conclusions

BER is a highly conserved pathway, responsible for repairing the vast majority of endogenous DNA damage including alkylation, oxidation, ring saturation, single strand breaks and base deamination. Defects in BER have been genetically linked to cancer risk. For instance, Pol β was found to play an important role in suppressing carcinogenesis since 30% of human tumors investigated to date carry Pol β mutations (Sobol et al., 2002, Starcevic et al., 2004, Wallace, 2014). More recently in our group, a study was undertaken in breast cancer and found that low Pol β is associated with poor survival (Abdel-Fatah et al., 2014). In addition, XRCC1 another key factor in BER, recently shown that loss of XRCC1 was associated with poor survival in breast cancer. Moreover, XRCC1 deficiency was highly associated with loss of other DNA repair factors, such as BRCA1 and p53 (Sultana et al., 2013a). Recent data indicate a cross talk between BRCA1 and BER factors. Saha et al study for instance showed that BRCA1 is involved in the transcriptional regulation of OGG1, NTH1, XRCC1 and APE1 (Saha et al., 2010). In agreement with this data, Chapter 3 of this thesis has shown that reduced BRCA1 expression in mutant/knockdown cell lines was associated with a significant reduction in several BER genes at the mRNA and protein expression. As BRCA1 deficient cells were shown to be BER deficient, cells may be more reliant on backup DSB repair pathways to maintain cellular survival. Blocking DSB repair should cause selective toxicity in BRCA1-BER deficient cells. The aim of this chapter was to investigate if targeting BRCA1 deficiency by ATM inhibitors (KU55933 or KU60019) is synthetically lethal.
Clonogenic survival assays (inhibitor exposure for total assay period) were conducted first to evaluate the cytotoxicity of ATM inhibitors against BRCA1 knockdown HeLa cells and BRCA1 mutated MDA-MB-436 cells, using Control HeLa and MCF7 cells as BRCA1 proficient controls, respectively. Then an alternative clonogenic assay protocol was performed (inhibitor exposure for 24 hours of assay period). For additional validation, the proliferation assay (MTS assay) was used to determine cell growth in response to ATM inhibitor. Results of all assays showed both KU55933 and KU60019 inhibitors induced significant cytotoxicity in BRCA1 deficient cells compared to BRCA1 proficient cells. These data suggest that BRCA1 deficient cells are reliant on ATM to maintain cellular survival.

To validate the cell line system used in the current study, clonogenic survival assay and MTS assay using PARP inhibitors (NU1025 and 3-Aminobenzamide) against BRCA1 deficient and proficient cell lines were performed. As predicted, both inhibitors induced significant cytotoxicity in BRCA1 deficient cells. These results support previously published data showing selective sensitivity to PARP inhibitors (in particular; NU1025 and 3-Aminobenzamide) in BRCA1 deficient models (Bryant et al., 2005, Sultana et al., 2012). Several functional studies were conducted to understand the mechanism behind this selective cytotoxicity of ATM inhibitors to BRCA1 deficient system. BRCA1 deficient cell lines were previously demonstrated to express reduced expression of several DSB repair factors (Chapter 3). The baseline level of DSBs damage present in these cells was assessed using γH2AX immunofluorescence microscopy assay, which is a well validated marker of DSBs. BRCA1 deficient cell lines treated with ATM inhibitors
(KU55933 or KU60019) for 48 hours were associated with a statistically significant increase in the accumulation of γH2AX foci compared to untreated BRCA1 deficient cells. Interestingly MCF7 cells showed a slight increase after KU60019 inhibitor treatment but not with KU55933, this result infers that KU60019 may be toxic to MCF7 (Figure 4.10B); this could be linked to the slight increase in apoptosis as well (Figure 4.16B).

These results support the hypothesis that ATM inhibition in BRCA1-BER deficient cells causes DSB accumulation which cannot be repaired due to an actual defect in DSB pathway. Moreover accumulation of γH2AX foci in cells treated with ATM inhibitors has been reported in previous publications (Shaheen et al., 2011, Sultana et al., 2013a). Although phosphorylation of H2AX is ATM dependent, accumulation of γH2AX foci in BRCA1 deficient cells was observed. One possible explanation for this finding is that other factors such as ATR or DNA-PKcs may be involved. Further mechanistic studies are required to clarify additional pathways.

DSB formation induces cell cycle checkpoint activation to trigger cell cycle arrest to allow DNA repair, or if the DNA damage is extensive can initiate apoptosis (Kurz and Lees-Miller, 2004). Cell cycle progression was arrested significantly at G2/M phase in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (after treatment with ATM inhibitors for 48 hours). BRCA1 mutated MDA-MB-436 cells showed a modest increase in G2/M arrest, though it was not significant (Figure 4.12 and 4.13). It should be noted that the flow cytometry assay was conducted after 24 and 48 hours, and to confirm G2/M arrest in MDA-MB-436 cells different time points should be tested in the future.
These increases in G2/M arrest support previously published data showing G2/M arrest after ATM inhibition (Sultana et al., 2013a, Lee et al., 2013, Nadkarni et al., 2012, Jazayeri et al., 2006). ATM phosphorylates Ser1423 of BRCA1 which is required for G2/M checkpoint activation (Cortez et al., 1999, Xu et al., 2001, Wu et al., 2010); in agreement with previous studies the current study shows that in the absence of BRCA1 and ATM, cells arrest in G2/M phase giving the cell time to repair. This could explain the survival data since the arrest in G2 would allow more NHEJ repair in BRCA1 deficient after blocking HR. Though it is interesting that ATM inhibition leads to G2/M arrest, one possible explanation is the overlapping function between ATM/CHK2 and ATR/CHK1 pathways which share many substrates (Smith et al., 2010). This overlapping function was seen in p53 deficient cells which rely on ATM and ATR to control the checkpoint signalling to survive after DNA damage (Reinhardt et al., 2007). G2/M arrest could then be followed by apoptosis. A flow cytometry-based apoptosis detection assay was performed. Results demonstrated that BRCA1 deficient cell lines treated with ATM inhibitors for 48 hours exhibited an increase in apoptosis rate compared to untreated BRCA1 deficient cell lines. These results support those previously published showing ATM inhibitor KU55933 treatment triggers apoptosis in different cancer cell lines (Sultana et al., 2013a, Li and Yang, 2010, Korwek et al., 2012) which again highlighted the importance of ATM in DNA repair and cell survival.

The data presented here support that ATM maybe a promising alternative synthetic lethality target. This study suggests that ATM inhibition is responsible for the synthetic lethality in BRCA1 deficient cells, and also
suggests that ATM inhibitors must be evaluated further as an alternative synthetic lethality strategy in in vivo models. A model for synthetic lethality is shown in Figure 4.17. In brief, impaired expression of BER factors in BRCA1 deficient cell lines leads to an increase accumulation of unrepaired SSBs which is followed by the formation of DSBs during DNA replication process. Therefore, it is suggested that cells are reliant on DSB pathway for cellular survival. Targeting the DSB pathway by ATM inhibition would lead to the observed synthetic lethality. In contrast, in cells that are proficient in DSB repair the DSBs would be repaired and cells would survive.

A limitation of this chapter is that this synthetic lethality study was restricted to two ATM inhibitors. KU60019 is known as an improved analogue of KU55933 (Golding et al., 2009) although the current results did not show this. For example, accumulation of γH2AX foci was significantly higher in MDA-MB-436 compared to untreated cells, though KU60019 was slightly toxic to MCF7 also. As a result more tests with KU60019 are needed using different cell lines. Another limitation was with cell cycle analysis which showed G2/M arrest increase with ATM treatment in BRCA1 HeLa cells, though MDA-MB-436 cells increase was not significant. This could be because samples were analysed after 24 and 48 hours. However, more time points should be tested in the future to monitor cells response after different treatment.

4.4.1. Conclusions

These in vitro studies demonstrate that a potential synthetic lethality relationship exists between BRCA1 deficiency and targeting the DSB pathway using ATM inhibitors (KU55933 or KU60019). The data suggests that ATM
inhibitors are selectively toxic to BRCA1 deficient cells. Also DNA DSBs accumulate following treatment with these inhibitors. In addition these data have shown a significant G2/M phase arrest in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells. As MDA-MB-436 cells showed a modest increase in G2/M arrest after 48 hours, this suggests that different time points should be tested in the future. Moreover, ATM inhibitor treatment triggers apoptosis in BRCA1 deficient cell lines.

These findings represent an alternative synthetic lethality strategy in targeting BRCA1 deficiency, which can potentially overcome PARP inhibition challenges. On the other hand, the global defect in DNA repair proved in BRCA1 deficient cells in particular the impaired BER factors expression, gives a wider role for using ATM inhibitors in cancers defective in BER in the absence of germline BRCA1 mutations. However, investigations with more BER deficient cell lines are required.
Figure 4.17. Proposed model of synthetic lethality between ATM inhibitors and BRCA1-BER deficient system.
Chapter 5

Targeting BRCA1 deficiency for personalized therapy using DNA-PKcs inhibitors (NU7441 & NU7026)
5. Targeting BRCA1 deficiency in breast cancer for personalized therapy using DNA-PKcs inhibitors (NU7441 & NU7026)

5.1. Introduction

BRCA1 facilitates the efficient resolution of DNA double-strand breaks (DSBs) through homologous recombination (HR) (Huen et al., 2010, Caestecker and Van de Walle, 2013). Cells lacking functional BRCA1 protein have impaired HR, and thus are suggested to depend on the more error-prone non-homologous end joining (NHEJ) pathway which is able to function at all stages of the cell cycle and does not require the presence of a homologous template. The NHEJ pathway is suggested to lead to chromosomal instability that leads to breast cancer development (Huen et al., 2010). It has been assumed that targeting the molecular machinery driving the DNA damage response, particularly the NHEJ pathway, with small molecule inhibitors will effectively enhance the efficacy of current cancer treatments that generate DNA damage (Kashishian et al., 2003, Davidson et al., 2013).

DNA-dependent protein kinase catalytic subunit (DNA-PKcs) is a key component of NHEJ pathway. DNA-PKcs has been shown to phosphorylate a large number of proteins, including NHEJ factors such as KU70/KU80, XRCC4, Lig3, XLF, WRN and Artemis along with other DNA damage response proteins including H2AX and p53. Recruitment of DNA-PKcs to the DNA-PK complex results in translocation of a heterodimer KU70/KU80 inward on the DNA, this is more likely to allow DNA-PKcs to bind directly to the damage resulting in the activation of the catalytic activity of the enzyme (Davis and Chen, 2013). Studies show that even with the absence of
heterodimer KU70/KU80, DNA-PKcs has unlimited kinase activity (West et al., 1998, Hammarsten and Chu, 1998). In addition synthetic lethality has been observed with PARP deficient cells targeted with DNA-PKcs inhibitor NU7026 (Veuger et al., 2003).

5.1.1. Rationale for study

Chapter 3 of this thesis suggested that BRCA1 deficiency is associated with reduced mRNA and/or protein expression of several factors involved in BER, HR and NHEJ. Data presented in Chapter 4 suggested a potential synthetic lethality relationship between BRCA1 deficiency and ATM inhibitors. In this chapter the aim was to investigate if BRCA1-BER deficient cells could be targeted by blockade of DNA-PKcs for personalized therapy.

5.2. Aims

The aims of this study were as follows:

1. To investigate the effect of DNA-PKcs inhibitors (NU7441 and NU7026) on cell viability and growth using clonogenic cell survival and MTS assays.
2. To test the hypothesis that DNA double strand breaks accumulate in cells after treatment with DNA-PKcs inhibitors in BRCA1 deficient cells.
3. To determine the effect of DNA-PKcs inhibitors on cell cycle in BRCA1 deficient cells.
4. To explore induction of apoptosis in BRCA1 deficient cells after treatment with DNA-PKcs inhibitors.
5.3. Results

5.3.1. Clonogenic cell survival and Growth inhibition in response to DNA-PKcs inhibitors (NU7441 & NU7026)

Clonogenic survival assays were performed to test whether BRCA1 deficient cells are sensitive to DNA-PKcs inhibition. BRCA1 mutated (MDA-MB-436) and knockdown (BRCA1 HeLa cells) cell lines along with MCF7 cells and Control HeLa cell lines were treated with increasing concentrations of NU7441 (C25H19NO3S) which is a potent, selective and competitive DNA-PKcs kinase inhibitor. Results are shown in Figure 5.1.

IC50 values were calculated using nonlinear regression, with the proviso that in BRCA1 proficient cell lines (Control HeLa and MCF7) cytotoxicity did not approach 50% in some cases, thus impairing extrapolation. Clonogenic assay demonstrated that BRCA1 HeLa cells, are more sensitive to NU7441 compared to Control HeLa cells (mean IC50= 0.6 ± 0.16 vs. IC50 above the maximum level tested). Similarly, increased sensitivity was also demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells (mean IC50= 0.44 ± 0.12 vs. mean IC50= 0.64 ± 0.19).
In general, increasing concentrations of NU7441 inhibitor reduced the colony forming ability in BRCA1 deficient cells compared to the BRCA1 proficient cells. To validate further, an alternative clonogenic assay protocol was performed, where cells were treated with DNA-PKcs inhibitor for 24 hours then culture media was replaced with fresh media and the remaining assay was conducted as described in 2.3.4.3. Results are illustrated in Figure 5.2.

**Figure 5.1. Clonogenic survival assay of DNA-PKcs inhibitor (NU7441).**

Cell lines with reduced BRCA1 expression exhibited decreased colony forming ability after treatment with NU7441 inhibitor for 14 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 79.22% ± 1.79%; BRCA1 HeLa- 66.44% ± 5.22% (B) MCF7- 51.21% ± 3.82%; MDA-MB-436- 41.54% ± 4.33%. Clonogenic data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
Cell lines with reduced BRCA1 expression exhibited decreased colony forming ability after treatment with NU7441 inhibitor for 24 hours. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 90.30% ± 0.90%; BRCA1 HeLa- 83.60% ± 2.40% (B) MCF7- 55.58% ± 0.02%; MDA-MB-436- 54.28% ± 1.05%. Clonogenic data are shown as the mean and SD values for each concentration from 2 independent experiments.

The alternative clonogenic assay results demonstrated that BRCA1 HeLa cells are more sensitive to NU7441 than Control HeLa cells. Similarly, sensitivity was demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells. However, data did not approach 50%, thus impairing extrapolation of IC₅₀ values. Though, both the 24 hour NU7441 treatment protocol and the 14 day treatment protocol, provide evidence that increasing concentration of NU7441 inhibitor reduces the colony forming ability of BRCA1 deficient cell lines compared to the BRCA1 proficient cell lines. The 14 day treatment protocol was used for all subsequent clonogenic assays described in the rest of the thesis.
For additional validation, the proliferation assay (MTS assay) was used to determine cell growth in response to NU7441. As shown in Figure 5.3, BRCA1 deficient HeLa cells were more sensitive to NU7441 than the Control HeLa cells. Similar sensitivity was demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells.

Figure 5.3. MTS cell growth inhibition assays of DNA-PKcs inhibitor (NU7441).

Cell lines with reduced BRCA1 expression exhibited decreased cell growth after treatment with NU7441 inhibitor for 6 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
For further validation NU7026 (C₁₇H₁₅NO₃), another potent kinase inhibitor of DNA-PKcs, was investigated (Hollick et al., 2003, Veuger et al., 2003, Amrein et al., 2007). As shown in Figure 5.4 clonogenic cell survival assays, BRCA1 HeLa cells are more sensitive to NU7026 than Control HeLa cells. Similar sensitivity was also demonstrated in BRCA1 mutated MDA-MB-436 cells compared to control MCF7 cells. Taken together, these data provide strong evidence that BRCA1 deficient cells are sensitive to DNA-PKcs inhibition.

![Clonogenic survival assay of DNA-PKcs inhibitor (NU7026).](image)

**Figure 5.4. Clonogenic survival assay of DNA-PKcs inhibitor (NU7026).**

Cell lines with reduced BRCA1 expression exhibited decreased colony forming ability after treatment with NU7026 inhibitor for 14 days, compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 82.50% ± 0.50%; BRCA1 HeLa- 76.00% ± 6.50% (B) MCF7- 58.21% ± 2.21%; MDA-MB-436- 41.50% ± 2.25%. Clonogenic data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
5.3.2. γH2AX focus immunofluorescence microscopy assay

To determine whether DSBs accumulate after treatment with NU7441, DSBs were measured using the γH2AX immunofluorescence microscopy assay. As described in Chapter 2 section 2.3.5, cells were treated with 1.5 µM of NU7441 for 24 hours and 48 hours. A significant increase in foci formation was observed in BRCA1 deficient cells after 48 hours treatment. Cells were then treated with 1 µM of NU7441 for 48 hours, but no significant accumulation of γH2AX foci in BRCA1 deficient cells was observed. The data support a dose response effect of DNA-PKcs inhibition in BRCA1 deficient cells.

5.3.2.1. γH2AX formation in response to DNA-PKcs inhibitors

BRCA1 proficient and BRCA1 deficient cell lines following treatment with 1.5 µM of NU7441 for 48 hours are shown in Figure 5.5. The data demonstrates a statistically significant increase in the accumulation of γH2AX foci in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (mean 62.50% ± 7.50 vs. mean 17.50% ± 0.50; p = 0.026) and compared to corresponding treated Control HeLa cells (mean 8.00% ± 3.00; p = 0.021). Similar results were also observed in treated MDA-MB-436 cells in which accumulation of γH2AX foci were significantly higher compared to untreated MDA-MB-436 cells (mean 81.00% ± 9 vs. 7% ± 2; p = 0.015) and compared to corresponding treated MCF7 cells (mean 4.50% ± 1.50; p = 0.013).
For further validation NU7026, another DNA-PKcs inhibitor, was investigated. Treatment with 1.5 µM of NU7026 produced similar results to NU7441. As shown in Figure 5.6, a statistically significant increase in the accumulation of γH2AX foci in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (mean 46% ± 4 vs. 11% ± 1.0; \( p = 0.013 \)), and compared to corresponding treated Control HeLa cells (mean 6.00% ± 2; \( p = 0.012 \)) was observed. Similar results were also observed in treated MDA-MB-436 cells in which accumulation of γH2AX foci was significantly higher compared to untreated MDA-MB-436 cells (mean 80.50% ± 6.5 vs. 17.50% ± 1.5; \( p = 0.011 \)), and compared to the corresponding treated MCF7 cells (mean 51% ± 1; \( p = 0.046 \)). Even though MDA-MB-436 cells were more sensitive to NU7026 inhibitor, MCF7 cells showed similar changes (from 10% to 51%).

Figure 5.5. Immunofluorescence microscopy staining for γH2AX foci formation following NU7441 treatment for 48 hours.

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * \( p < 0.05 \), \( p \) value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
This result infers that NU7026 inhibitor may be toxic to MCF7, especially as this increase was not seen with NU7441.

![Figure 5.6: Immunofluorescence microscopy staining for γH2AX foci formation following NU7026 treatment for 48 hours.](image)

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.

### 5.3.3. Cell cycle analysis by propidium iodide flow cytometry after treatment with DNA-PKcs inhibitors

Cells were treated as described in Chapter 2, section 2.3.6 for 24 hours and 48 hours with 1 µM or 1.5 µM of NU7441 (Figure 5.7). Although no significant difference was observed after 24 hours, at 48 hours in BRCA1 deficient cells treated with 1.5 µM NU7441 exhibited a significant increase in G1 phase arrest compared to control was observed.
Results for BRCA1 proficient and BRCA1 deficient cells following treatment with 1.5 μM (NU7441) for 48 hours are shown in Figure 5.8. The data demonstrate an increase in G1 arrest in treated BRCA1 HeLa cells compared with untreated BRCA1 HeLa cells (mean 60.6% ± 0.60 vs. 47.2% ± 3.80; though not significant p = 0.07), and compared to corresponding treated
Control HeLa cells (mean 32.4% ± 0.42; $p = 0.0007$). Results of treated MDA-MB-436 cells demonstrate an increase in G1 arrest compared to MDA-MB-436 untreated cells, though not significant (mean 49.4% ± 0.15 vs. mean 37.8% ± 3.6; $p = 0.08$). The observed G1 arrest was significantly higher in treated MDA-MB-436 cells compared to treated MCF7 cells (mean 30.4% ± 0.50; $p < 0.0001$). On the other hand, Control HeLa and MCF7 cells did not show the same rate of increase (i.e. ~12% increase like BRCA1 deficient cells) after treatment (Control HeLa and MCF7 cells, G1 arrest mean 41% to 32.4% and 25% to 30.4%, respectively).

![Flow cytometric cell cycle analysis following NU7441 treatment for 48 hours.](image)

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. *** $p<0.001$ and **** $p<0.0001$, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
For further validation NU7026, another DNA-PKcs inhibitor, was investigated. Treatment with 10 µM of NU7026 produced similar results as with NU7441. As shown in Figure 5.9, increased G1 arrest in treated BRCA1 HeLa cells was demonstrated as compared to untreated BRCA1 HeLa cells (mean 61% ± 1 vs. 54.5% ± 0.5; p = 0.028). A modest, although not statistically significant, G1 arrest was also observed in treated MDA-MB-436 cells compared to untreated MDA-MB-436 cells (mean 40% ± 0.1 vs. 31.7% ± 3.8; p = 0.1). A statistically significant increase in G1 arrest was observed when treated MDA-MB-436 cells were compared to treated MCF7 cells (mean 27.25% ± 2.8; p = 0.04). In contrast, Control HeLa and MCF7 cells did not show the same increase.

![Flow cytometric cell cycle analysis following NU7026 treatment for 48 hours.](image)

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 3 independent experiments. * p<0.05, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
5.3.4. Apoptosis detection by annexin V-FITC flow cytometry after treatment with DNA-PKcs inhibitors

Following evaluation of cell cycle arrest in BRCA1 deficient cells treated with DNA-PKcs inhibitors, Annexin V was used in conjunction with a propidium iodide (PI) for identification of early and late apoptotic cells as described in Chapter 2, section 2.3.7. Arrest at the G1 phase of cell cycle may lead to apoptosis which could be the cause of the reduced survival observed in BRCA1 deficient cells in the clonogenic survival assay. Cells were treated for 24 hours and 48 hours with 1.5 µM of NU7441. Although no significant increase in apoptosis was evident after 24 hours, at 48 hours in BRCA1 deficient cells treated with 1.5 µM of NU7441 significant increase in apoptosis compared to control cells was observed.

Results for BRCA1 proficient and BRCA1 deficient cells following treatment with 1.5 µM (NU7441) for 48 hours are shown in Figure 5.10, data were normalised against baseline apoptotic fraction (untreated cells) to determine the percentage increase in apoptosis. The data demonstrate a statistically significant increase in apoptosis with treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells (mean 6.3% ± 0.36 vs. mean 3.6% ± 0.68; p = 0.0057). Similar results were also observed in treated MDA-MB-436 cells, in which a significant increase in apoptosis as compared to untreated MDA-MB-436 cells was observed (mean 13.5% ± 0.77 vs. 4.68% ± 0.07; p < 0.0001). On the other hand, Control HeLa and MCF7 cells did not show the same rate of increase (mean 2.44% to 3.25% and 4.06% to 4.76%, respectively).
For further validation NU7026, another DNA-PKcs inhibitor, was investigated. Treatment with 10 µM of NU7026 produced similar results as with NU7441. As shown in Figure 5.11, the data demonstrates a statistically significant increase in apoptosis in treated BRCA1 HeLa cells compared to the corresponding untreated BRCA1 HeLa cells (mean 8.71% ± 1.22 vs. mean 4.72% ± 0.34; p = 0.034). A similar increase was also observed in treated MDA-MB-436 cells as compared to untreated MDA-MB-436 cells (mean 9.13% vs. 4.67%; though was not significant p=0.06). On the other hand,
Control HeLa and MCF7 cells did not show the same rate of increase (mean 3.11% to 3.16% and 3.15% to 5.6%, respectively).

Figure 5.11. Apoptosis detection by annexin V-FITC flow cytometry in NU7026 treated cells.

Treatment with NU7026 inhibitor was associated with a significant increase in apoptotic cells. Data were normalised against baseline apoptotic fraction to determine the percentage increase in apoptosis. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
5.4. Discussion and conclusions

The aim of this chapter was to investigate whether targeting BRCA1 deficiency by DNA-PKcs, a key player in NHEJ pathway, using NU7441 or NU7026 inhibitors, would be synthetically lethal.

The role of BRCA1 in regulating the NHEJ pathway has been investigated for a number of years. The data have been conflicting with a number of studies showing a major role of BRCA1 in NHEJ and maintenance of genomic integrity (Zhong et al., 2002, Bau et al., 2004), while others did not show a role of BRCA1 in NHEJ (Moynahan et al., 1999, Merel et al., 2002). The lack of consistency between these results may be partly explained by the use of different assays, which may measure different sub-pathways of NHEJ (Bau et al., 2006). Several studies have confirmed the interaction between BRCA1 and NHEJ factors. For example, one study showed that BRCA1 interacts with a subunit of the DNA-PK complex (KU80). Another study showed that BRCA1 is rapidly recruited to DSB and is dependent on its association with KU80 (Wei et al., 2008). A more recent study has found that DNA-PKcs interacts with BRCA1 in the absence of KU70 and KU80 (Davis et al., 2014b).

To test the importance of both BRCA1 and DNA-PKcs in the maintenance of cellular survival, clonogenic survival assays (inhibitor exposure for total assay period) were conducted to evaluate the cytotoxicity in BRCA1 knockdown HeLa cells and BRCA1 mutated MDA-MB-436 cells treated with DNA-PKcs inhibitors (NU7441 or NU7026), using Control HeLa and MCF7 cells as BRCA1 proficient controls, respectively. Then an alternative clonogenic assay protocol was performed (inhibitor exposure for 24 hours of assay period).
Results of both protocols showed high sensitivity in BRCA1 deficient cells compared to BRCA1 proficient cells. For additional validation, MTS assay was used to determine cell viability in response to DNA-PKcs inhibitor. Both NU7441 and NU7026 inhibitors induced significant cytotoxicity in BRCA1 deficient cells compared to BRCA1 proficient cells. Taken together, these data suggest that BRCA1 deficient cells are reliant on DNA-PKcs to maintain cellular survival. For mechanistic insights, several functional studies were conducted. BRCA1 deficient cell lines treated with DNA-PKcs inhibitors for 48 hours demonstrate a statistically significant increase in the accumulation of γH2AX foci compared to untreated BRCA1 cell lines. This finding supports previous publications showing accumulation of γH2AX foci with inhibition of DNA-PKcs (Shaheen et al., 2011, Sultana et al., 2013a). Interestingly MCF7 showed a slight increase after NU7026 inhibitor treatment but not with NU7441, this result infers that NU7026 may be toxic to MCF7 (Figure 5.6B); this could also be linked to the slight increase in apoptosis (Figure 5.11B).

Results from Chapter 3 illustrate that ATM, ATR and DNA-PKcs are down regulated in BRCA1 deficient cells; these three enzymes are known to have a role in phosphorylation of H2AX (Podhorecka et al., 2010). Previously it had been shown that DSB deficient cells exhibited reduced phosphorylation of H2AX after ionizing radiation treatment, however they conclude that inhibitions of DNA-PKcs and ATM reduce, but do not eliminate, the accumulation of γH2AX foci (Paull et al., 2000). In the current study inhibition of DNA-PKcs or ATM in BRCA1 deficient cells have increased γH2AX foci formation around DSBs as compared to the untreated cells. The fact that γH2AX was still forming in my study can have two explanations. Firstly, with
a low expression of ATM and ATR, in the absence of DNA-PKcs, other enzymes (e.g. CHK2) may activate γH2AX, as has been shown in a previous study (Tu et al., 2013). Alternatively another signaling pathway may modulate γH2AX levels, for example DNA-PKcs has been suggested to phosphorylates H2AX directly and modulate γH2AX level indirectly through the Akt/GSK3β signaling pathway (An et al., 2010). In the absence of DNA-PKcs it may be that the Akt/GSK3β pathway can be activated by other factors. Secondly, the fact that ATM and ATR are down regulated in BRCA1 deficient cells does not mean it is completely silenced, and in the absence of DNA-PKcs low levels of ATM or ATR may be sufficient to phosphorylate H2AX. Taken together, these results support the hypothesis that DNA-PKcs inhibition in BRCA1-BER deficient cells causes DSB accumulation which cannot be repaired due to an actual defect in DSB pathway.

Monitoring cell cycle progression in BRCA1 deficient cell lines after treatment with DNA-PKcs inhibitors for 48 hours was associated with a modest increase arrest at the G1 phase compared to untreated BRCA1 deficient cell lines. However, this increase was not always significant and different time points should be tested in the future. These modest increases in G1 arrest support those previously published showing that G1 arrest increases after inhibition with DNA-PKcs (NU7441) inhibitor, in colon cancer (Zhao et al., 2006), prostate cancer (Shaheen et al., 2011, Yu et al., 2012) and breast cancer cell lines (Ciszewski et al., 2014). A more recent study showed that BRCA1 facilitates accurate repair during G1 phase through NHEJ (Jiang et al., 2013). To evaluate induction of apoptosis, a flow cytometry-based apoptosis detection assay was performed. Results demonstrated that BRCA1 deficient cell lines
treated with DNA-PKcs inhibitors for 48 hours exhibited increase in apoptosis rate compared to untreated BRCA1 deficient cell lines. These results support those previously published showing an increase in apoptosis after DNA-PKcs inhibition, in prostate cancer (Yu et al., 2012) and XRCC1 knockdown HeLa cell lines (Sultana et al., 2013a).

Taken together, the data presented here demonstrated that a potential synthetic lethality relationship which exists between BRCA1 deficiency and DNA-PKcs inhibition. Also the present study highlights the important role of DNA-PKcs in maintaining cellular survival in BRCA1 deficient cell. A model for synthetic lethality is shown in Figure 5.12. In brief, impaired expression of BER in BRCA1 deficient cell lines leads to accumulation of unrepai red SSBs followed by the formation of DSBs during DNA replication process. Therefore, it is suggested that cells are reliant on backup DSB pathway for cellular survival. Targeting the DSB pathway by DNA-PKcs inhibition would lead to the observed synthetic lethality. In contrast, in cells that are proficient in DSB repair the DSBs would be repaired and cells would survive.

It should be noted that these in vitro studies have limitations. Firstly, this investigation was restricted to two DNA-PKcs inhibitors. And a recent study has indicated that DNA-PKcs inhibitors currently available are limited by poor pharmacokinetics (Davidson et al., 2013). Secondly, cell cycle analysis showed modest G1 increase with DNA-PKcs treatment, this could be because samples were analysed after 24 and 48 hours. However, more time points should be tested in the future to monitor cells response after different treatments.
5.4.1. Conclusions

To conclude, this *in vitro* study has shown a potential synthetic lethality between BRCA1 deficiency and DNA-PKcs inhibition. Firstly, data have shown a selective toxicity in BRCA1 deficient cells to DNA-PKcs inhibitors (NU7441 or NU7026). Secondly, DNA DSB accumulation was observed after treatment with DNA-PKcs inhibitors in BRCA1 deficient cells compared to untreated cells. In addition modest increase in G1 phase arrest was detected, though this increase was not always significant and different time points should be tested in the future. Then again, DNA-PKcs inhibition was found to trigger apoptosis in BRCA1 deficient cells. Moreover, these findings present a potential new alternative synthetic lethality strategy in BRCA1 deficient cells. Finally, in agreement with previous studies this study shows the importance of DNA-PKcs in maintaining cellular survival.
Figure 5.12. Proposed model of synthetic lethality between DNA-PKcs inhibitors and BRCA1-BER deficient system.
Chapter 6

Cisplatin in combination with Ataxia telangiectasia mutated (ATM) or DNA dependent protein kinase catalytic subunit (DNA-PKcs) inhibitor in BRCA1 deficient cells
6. Cisplatin in combination with Ataxia telangiectasia mutated (ATM) or DNA dependent protein kinase catalytic subunit (DNA-PKcs) inhibitor in BRCA1 deficient cells

6.1. Introduction

Cisplatin [cis-diamminedichloroplatinum(II)] is a platinating anticancer chemotherapeutic agent that acts by covalently binding to DNA and forming DNA damaging adducts. Cisplatin is effective in the treatment of many cancers including metastatic testicular cancers, bladder, cervical, ovarian, lung, head and neck cancers (Kelland, 2007, Ratanaphan, 2011). Several preclinical studies have shown that BRCA1 deficient cell lines are hypersensitive to cisplatin treatment compared to their wild type matched controls (Bhattacharyya et al., 2000, Tassone et al., 2003, Tan et al., 2008). In addition, clinical studies have demonstrated similar hypersensitivity to cisplatin (Byrski et al., 2009, Taron et al., 2004, Quinn et al., 2007, Font et al., 2011, Silver et al., 2010).

In the previous chapters, a potential synthetic lethality was demonstrated between BRCA1 deficiency and ATM inhibition or DNA-PKcs inhibition in in vitro cell culture system. In light of these data, cisplatin was investigated in combination with ATM or DNA-PKcs inhibitor in BRCA1 proficient and BRCA1 deficient cells.
6.1.1. Rationale for study

The data presented in previous chapters suggested that BRCA1 deficiency is associated with low expression of BER. BRCA1-BER deficient cells are amenable to synthetic lethality targeting using ATM or DNA-PKcs inhibitors. As BRCA1 deficient cells are cisplatin sensitive, the potential of a combination strategy in BRCA1-BER proficient cells was investigated.

6.2. Aims

The aims of this study were as follows:

1. To investigate the effect of cisplatin in combination with ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) in clonogenic cell survival assay.

2. To test the hypothesis that DNA double strand breaks accumulate in cells after treatment with cisplatin in combination with ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) in BRCA1 deficient cells.

3. To determine the effect of cisplatin / ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) combination on cell cycle progression in BRCA1 deficient cells.

4. To quantify apoptosis in BRCA1 deficient cells after treatment with cisplatin / ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) combinations.
6.3. Results

6.3.1. Clonogenic cell survival assay in response to Cisplatin

BRCA1 mutated (MDA-MB-436) cells and knockdown (BRCA1 HeLa) cell lines along with control cells MCF7 and Control HeLa cell lines respectively, were treated with increasing concentrations of cisplatin ranging from (0.25 µM to 1.25 µM). Clonogenic survival assay results are shown in Figure 6.1. No colonies were obtained at any dose for BRCA1 HeLa and MDA-MB-436, showing high sensitivity in these cell lines and that the IC\textsubscript{50} cannot be estimated from these data but must be below the minimum value tested. On the other hand, Control HeLa, mean IC\textsubscript{50}= 0.25 µM ± 0.6 and MCF7, mean IC\textsubscript{50}= 1.23 µM ± 0.9.

![Clonogenic survival assay of cisplatin.](image)

After exposure for 16 hours to cisplatin ranging from (0.25 µM to 1.25 µM), no colonies were obtained at any dose for BRCA1 HeLa and MDA-MB-436 showing high sensitivity in these lines compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines were as follows: (A) Control HeLa- 75% ; BRCA1 HeLa- 75% (B) MCF7- 56.5% ; MDA-MB-436- 49.75%. Clonogenic data are shown as the mean and SD values for each concentration from 2 experiments.
6.3.2. Clonogenic cell survival assay in cells treated with KU55933 or NU7441 in combination with cisplatin

To evaluate the combination between cisplatin and ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441), cells were cultured with increasing concentrations of cisplatin. Cells were treated with increasing concentration of cisplatin ranging from (0.00001 μM to 0.1 μM) and KU55933 (5μM, HeLa cell lines and 7.5 μM, breast cancer cell lines) or NU7441 (0.5μM, HeLa cell lines and 0.75 μM, breast cancer cell lines). The concentration range for cisplatin in combination studies was lower as the cells were hypersensitive at higher concentrations and underwent profound cell death.

Clonogenic survival assays results for the combination of ATM inhibitor and cisplatin are shown in Figure 6.2. BRCA1 knockdown cells are more sensitive to combination treatment than cisplatin alone (mean IC₅₀ = 0.00052 ± 0.00019 vs. IC₅₀ above the maximum level tested), and compared to BRCA1 proficient Control HeLa cells. Similar sensitivity was also demonstrated in combination treated MDA-MB-436 cells compared to cisplatin alone (mean IC₅₀ = 0.0003 ± 0.00013 vs. IC₅₀ = 0.16 ± 0.026), and compared to control MCF7 cells.

Clonogenic survival assays results for the combination of DNA-PKcs inhibitor and cisplatin are shown in Figure 6.3. BRCA1 knockdown cells are more sensitive to combination treatment than cisplatin alone (mean IC₅₀ = 0.09 ± 0.098 vs. IC₅₀ above the maximum level tested), and comparing to BRCA1 proficient Control HeLa cells. Similar sensitivity was also demonstrated in
combination treated MDA-MB-436 cells compared to cisplatin alone (mean IC$_{50}$ = 0.005 ± 0.0047 vs. IC$_{50}$ = 0.16 ± 0.026), and compared to control MCF7 cells.

Figure 6.2. Clonogenic survival assay of ATM inhibitor (KU55933) in combination with cisplatin.

BRCA1 deficient cells show decreased colony forming ability after exposure for 16 hours to cisplatin ranging from (0.00001 µM to 0.1 µM) and then KU55933 (5 µM, HeLa cells lines and 7.5 µM, breast cancer cell lines), compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 59.33% ± 1.69%; BRCA1 HeLa- 65.67% ± 0.60% (B) MCF7- 44.89% ± 0.72%; MDA-MB-436- 24.53% ± 2.76%. Clonogenic data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
Figure 6.3. Clonogenic survival assay of DNA-PKcs inhibitor (NU7441) in combination with cisplatin.

BRCA1 deficient cells show decreased colony forming ability after exposure for 16 hours to cisplatin ranging from (0.00001 µM to 0.1 µM) and then NU7441 (0.5 µM, HeLa cell lines and 0.75 µM, breast cancer cell lines), compared to BRCA1 proficient cell lines. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Plating efficiencies of cell lines with standard error were as follows: (A) Control HeLa- 60% ± 1.36%; BRCA1 HeLa- 64.83% ± 1.55% (B) MCF7- 46.78% ± 2.61%; MDA-MB-436- 25.87% ± 2.36%. Clonogenic data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments.
6.3.3. γH2AX immunofluorescence microscopy in response to KU55933 or NU7441 in combination with cisplatin

To determine whether DNA double strand breaks (DSBs) accumulate after treatment with KU55933 and cisplatin, DSBs were measured using the γH2AX immunofluorescence microscopy assay. As described in Chapter 2 section 2.3.5, cells were treated with cisplatin (0.1 µM) and KU55933 (for HeLa cell lines = 5 µM or breast cancer cell lines = 7.5 µM) for 24 hours and 48 hours. Although no significant changes were observed after 24 hours, at 48 hours a significant increase in foci formation was observed in BRCA1 deficient cells. As shown in Figure 6.4, the data demonstrates a statistically significant increase in the accumulation of γH2AX foci in combination treated BRCA1 HeLa cells compared to corresponding cisplatin alone treated BRCA1 HeLa cells and untreated BRCA1 HeLa cells (76.50% vs. 18%; p = 0.003 and 76.50% vs. 11%; p = 0.0017, respectively) and compared to combination treated Control HeLa cells (15.00%; p = 0.0019).

Similar results were also observed in combination treated MDA-MB-436 cells in which accumulation of γH2AX foci were higher compared to corresponding cisplatin alone treated MDA-MB-436 cells and untreated MDA-MB-436 cells (60.50% vs. 36%; p = 0.06 and 60.50% vs. 17.50%; p = 0.012, respectively) and compared to combination treated MCF7 cells (31%; p = 0.032).
Figure 6.4. Immunofluorescence microscopy staining for γH2AX foci formation following ATM inhibitor (KU55933) in combination with cisplatin for 48 hours exposure.

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥2 independent experiments. * p<0.05 and ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
Alternatively, cells were treated with cisplatin (0.1 µM) and NU7441 (for HeLa cell lines = 0.5 µM or breast cancer cell lines = 0.75 µM) for 24 hours and 48 hours. Although no significant changes were observed after 24 hours, at 48 hours in BRCA1 deficient cells a significant increase in foci formation was observed. As shown in Figure 6.5, the data demonstrates a statistically significant increase in the accumulation of γH2AX foci in combination treated BRCA1 HeLa cells compared to corresponding cisplatin alone treated BRCA1 HeLa cells and untreated BRCA1 HeLa cells (87% vs. 18%; p = 0.0017 and 87% vs. 11%; p = 0.0009, respectively) and compared to combination treated Control HeLa cells (16%; p = 0.0026).

Similar results were also observed in combination treated MDA-MB-436 cells in which accumulation of γH2AX foci were higher compared to corresponding cisplatin alone treated MDA-MB-436 cells and untreated MDA-MB-436 cells (81.5% vs. 36%; p = 0.017 and 81.5% vs. 17.50%; p = 0.003, respectively) and compared to combination treated MCF7 cells (38.5%; p = 0.013).
Figure 6.5. Immunofluorescence microscopy staining for γH2AX foci formation following DNA-PKcs inhibitor (NU7441) in combination with cisplatin for 48 hours exposure.

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05, ** p<0.01 and *** p<0.001, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
6.3.4. Flow cytometric cell cycle analysis in response to ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) in combination with Cisplatin

For cell cycle analysis, cells were treated with cisplatin (0.1 µM) and KU55933 (for HeLa cell lines = 5 µM or breast cancer cell lines = 7.5 µM) for 48 hours. Results for BRCA1 proficient and BRCA1 deficient cells following combination treatment are shown in Figure 6.6. The data demonstrate an increase in G2/M arrest in combination treated BRCA1 HeLa cells compared to corresponding cisplatin alone treated BRCA1 HeLa cells and untreated BRCA1 HeLa cells (mean 37.6%, 29.5% and 25.8%; respectively). Similar results were also observed in combination treated MDA-MB-436 cells in which G2/M arrest were significantly higher compared to untreated MDA-MB-436 cells (mean 38% vs. 32.4%). On the other hand, cisplatin alone treated MDA-MB-436 cells showed mean of 41.4% G2/M arrest.

Conversely, cells were treated with cisplatin (0.1 µM) and NU7441 (for HeLa cell lines = 0.5 µM or breast cancer cell lines = 0.75 µM) for 48 hours. As shown in Figure 6.7, the data demonstrates an increase in G2/M arrest in combination treated BRCA1 HeLa cells compared with corresponding cisplatin alone treated BRCA1 HeLa cells and untreated BRCA1 HeLa cells (mean 34.2%, 29.5% and 25.8%; respectively). Similar results were also observed in combination treated MDA-MB-436 cells in which G2/M arrest was significantly higher compared to untreated MDA-MB-436 cells (mean 38.2% vs. 32.4%). On the other hand, cisplatin alone treated MDA-MB-436 cells showed a mean G2/M arrest of 41.4%. Control HeLa and MCF7 cells did not show the same increase as BRCA1 deficient cells with both combination treatments.
Figure 6.6. Flow cytometric cell cycle analysis following ATM inhibitor (KU55933) in combination with cisplatin

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05 and ** p<0.01, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
Figure 6.7. Flow cytometric cell cycle analysis following DNA-PKcs inhibitor (NU7441) in combination with cisplatin

(A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from ≥ 2 independent experiments. * p<0.05 and ** p<0.01, *** p<0.001, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
6.3.5. Apoptosis detection by annexin V-FITC flow cytometry in response to KU55933 or NU7441 in combination with cisplatin

Cells were treated with cisplatin (0.1 µM) and KU55933 (for HeLa cell lines = 5 µM or breast cancer cell lines = 7.5 µM) for 48 hours. Results for BRCA1 proficient and BRCA1 deficient cells following combination treatment are shown in Figure 6.8, data were normalised against baseline apoptotic fraction (untreated cells) to determine the percentage increase in apoptosis. The data demonstrate a statistically significant increase in combination treated BRCA1 HeLa cells compared to corresponding cisplatin alone treated BRCA1 HeLa cells and untreated BRCA1 HeLa cells (mean 20.6%, 15.9% and 5.3%; respectively). Similar results were also observed in combination treated MDA-MB-436 cells in which the increase in apoptosis was significantly higher compared to cisplatin alone treated MDA-MB-436 and compared to untreated MDA-MB-436 cells (mean 17.6%, 7.4% and 4.9%; respectively). On the other hand, Control HeLa and MCF7 cells did not show the same high increase (i.e. around 14% increase with BRCA1 deficient cells) after cisplatin and KU55933 treatment (mean 4.35% to 7.49% and 3.17% to 9.25%, respectively).

Conversely, cells were treated with cisplatin (0.1 µM) and NU7441 (for HeLa cell lines = 0.5 µM or breast cancer cell lines = 0.75 µM) for 48 hours. As shown in Figure 6.9, the data demonstrates a statistically significant increase in combination treated BRCA1 HeLa cells compared to corresponding cisplatin alone treated BRCA1 HeLa cells and untreated BRCA1 HeLa cells (mean 25%, 15.9% and 5.3%; respectively). Similar results were also observed in combination treated MDA-MB-436 cells in which the increase in apoptosis was significantly higher compared to cisplatin alone treated MDA-MB-436
and compared to untreated MDA-MB-436 cells (mean 19.2%, 7.4% and 4.9%; respectively). On the other hand, Control HeLa and MCF7 cells did not show the same high increase (i.e. around 17% increase with BRCA1 deficient cells) after cisplatin and NU7441 treatment (mean 4.35% to 4.13% and 3.17% to 11%, respectively).
Figure 6.8. Apoptosis detection by annexin V-FITC flow cytometry in response to ATM inhibitor (KU55933) in combination with cisplatin

Treatment with combination KU55933 and cisplatin was associated with a significant increase in apoptotic. Data were normalised against baseline apoptotic fraction to determine the percentage increase in apoptosis. (A) BRCA1 HeLa cells compared to Control HeLa cells. (B) MDA-MB-436 cells compared to MCF7 cells. Data are shown as the mean and SD values for each concentration from 3 independent experiments. * p<0.05, ** p<0.01, *** p<0.001 and **** p<0.0001, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
Figure 6.9. Apoptosis detection by annexin V-FITC flow cytometry in response to DNA-PKcs inhibitor (NU7441) in combination with cisplatin

Exposure to combination NU7441 and cisplatin was associated with a significant increase in apoptotic. Data were normalised against baseline apoptotic fraction to determine the percentage increase in apoptosis. (A) BRCA1 HeLa cell line compared to Control HeLa. (B) MDA-MB-436 compared to MCF7. Data are shown as the mean and SD values for each concentration from 3 independent experiments. ** p<0.01 and **** p<0.0001, p value was assessed by a t-test comparing BRCA1 deficient cell lines to BRCA1 proficient cell lines. Graphs were produced and statistical analysis performed using GraphPad prism.
6.4. Discussion and conclusions

In previous chapters (Chapter 4 and Chapter 5), it was suggested that there was a synthetic lethality relationship between BRCA1 deficiency and targeting DSB repair pathway by blockade of ATM or DNA-PKcs. ATM and DNA-PKcs modulation has been shown to enhance cisplatin cytotoxicity (Yoshida et al., 2008, Dejmek et al., 2009). Therefore, in this section of the study, the efficacy of combining cisplatin with ATM or DNA-PKcs inhibitor in BRCA1 proficient and BRCA1 deficient cells was investigated. To achieve this, clonogenic survival assay was conducted first to evaluate the cytotoxicity of cisplatin alone followed by combination studies using ATM (KU55933) or DNA-PKcs (NU7441) inhibitors. In agreement with previous studies, BRCA1 showed hypersensitivity to platinum therapy (Husain et al., 1998, Bhattacharyya et al., 2000, Tassone et al., 2003, Tan et al., 2008, Byrski et al., 2009, Taron et al., 2004, Quinn et al., 2007, Font et al., 2011, Silver et al., 2010). The dose of cisplatin used for the combination treatment in this study was 1/10th the dose used in previous preclinical studies. The results of both combination treatments (cisplatin/ATM inhibitor and cisplatin/DNA-PKcs inhibitor) induced hypersensitivity in BRCA1 deficient cells compared to BRCA1 deficient cells treated with cisplatin alone. Formation of γH2AX foci in BRCA1 deficient cells following combination treatments increased compared to cisplatin alone and compared to either inhibitor alone. Whereas BRCA1 knockdown cells showed 76.5% increase in the accumulation of γH2AX foci following ATM inhibitor/cisplatin treatment (Figure 6.4A) and an 87% increase with DNA-PKcs inhibitor/cisplatin treatment (Figure 6.5A), ATM inhibitor alone showed 66.5% (Figure 4.9A) and DNA-PKcs inhibitor
alone showed 62.5% (Figure 5.5A). Interestingly, this increase in γH2AX foci formation was not seen following combination treatments in the BRCA1 mutated cells (Figure 6.4B and 6.5B) compared to each inhibitor alone (Figure 4.9B and 5.5B). This observation may be because MDA-MB-436 cells were found to be hypersensitive to the combination treatments and exhibited profound cell death. It should be noted that formation of γH2AX foci was analyzed after 48 hours with both single and combination treatments and to follow the damage after combination treatments different time points should be tested in the future.

The flow cytometry assay showed that BRCA1 deficient cells arrested significantly at G2/M phase after combination treatments compared to either inhibitor alone. Whereas BRCA1 deficient cell lines showed 37.6% and 38% increase in G2/M arrest following ATM inhibitor/cisplatin treatment (Figure 6.6), ATM inhibitor alone showed 26.6% and 32.45% (Figure 4.12), respectively. A similar increase was observed following DNA-PKcs inhibitor/cisplatin treatment with 34.25% and 38.25% (Figure 6.7), compared to 21.8% and 26.55% following DNA-PKcs inhibitor alone (Figure 5.8), respectively. In previous studies, G2/M arrest was demonstrated when combining KU55933 and NU7441 with Doxorubicin or Neocarzinostatin or Ionizing radiation (Zhao et al., 2006, Yu et al., 2012, Shaheen et al., 2011, Ciszewski et al., 2014). Cisplatin has been suggested to activate CHK1 and CHK2 which then leads to G2 arrest. Previous studies have also showed the selectivity of cisplatin in the killing of p53 defective tumor cells (Fan et al., 1995, Lee et al., 1999). Interestingly, targeting DNA-PKcs alone in BRCA1 deficient cells showed G1 arrest, whereas in the combination study G2/M
arrest was observed. This current study suggests that in combination treatments cisplatin activates G2/M arrest independent of DNA-PKcs inhibition. After G2/M arrest, BRCA1 deficient cells were unable to repair the DSBs and hence initiate apoptosis. A significant accumulation of apoptosis was observed in ATM inhibitor/cisplatin combination treatment, showing 20.6% and 17.6% (Figure 6.8) compared to 11.12% and 13.1% following ATM inhibitor alone (Figure 4.15) in BRCA1 HeLa and MDA-MB-436 cells, respectively. A similar increase in apoptosis was observed following DNA-PKcs inhibitor/cisplatin treatment with 25% and 19.2% (Figure 6.9) compared to 6.3% and 13.5% following DNA-PKcs inhibitor alone (Figure 5.10) in BRCA1 HeLa and MDA-MB-436 cells, respectively.

Taken together, several important observations have emerged from this chapter: firstly, the platinum anticancer drug cisplatin is selectively toxic to BRCA1 deficient cells. Cisplatin sensitivity in BRCA1 deficient cells observed here is consistent with previous publications. Secondly, inhibition of ATM or DNA-PKcs in combination with cisplatin enhances the cytotoxicity compared to either inhibitor alone. However, it should be noted that investigating drug combination interactions is an important area to study and it is one of the limitations in this chapter. For future studies in order to evaluate the interaction between ATM inhibitors and cisplatin or DNA-PKcs inhibitors and cisplatin, number of published methodologies can be used to evaluate the nature of drug-drug combination, e.g. isobologram, combination index, curve shift analysis (Evaluation of drug interaction methods are listed with more details in APPENDIX D). Thirdly, combination treatments increase DSB accumulation in BRCA1 deficient cells. Finally, combination treatments increase G2/M cell
cycle arrest, followed by an increase in apoptosis. A model for synthetic lethality is shown in Figure 6.10. In brief, impaired BER factors in BRCA1 deficient cell lines leads to accumulation of unrepaired SSBs followed by the formation of DSBs during DNA replication process. Therefore, it is suggested that cells are reliant on backup DSB pathways for cellular survival. Targeting DNA-PKcs or ATM along with cisplatin could induce extensive DSBs and over whelm BRCA1 deficient cells. This could lead to the observed synthetic lethality. In contrast, cells that are proficient in DSB repair (BRCA1-BER proficient), the DSBs would be repaired and cells would survive.

6.4.1. Conclusions

To conclude, this chapter of the study suggests that ATM and DNA-PKcs inhibition enhances cisplatin sensitivity in BRCA1 deficient cells. Firstly, data have shown that inhibition of ATM or DNA-PKcs in combination with cisplatin enhances the cytotoxicity compared to cisplatin alone. Secondly, combination treatments increase DSBs accumulation in BRCA1 deficient cells compared to cisplatin alone. In addition, combination treatments increase G2/M cell cycle arrest, followed by an increase in apoptosis. Finally, combination data were compared with corresponding cells treated with inhibitors alone (data presented in chapter 4 and chapter 5); the results generally suggested improved outcomes in combination treatments compared to each inhibitor alone. The data implies that the combination therapy could be an attractive strategy in BRCA1 cancer cells. However, advanced preclinical investigations with more cell lines and inhibitors are required.
Figure 6.10. Working model for cisplatin and ATM or DNA-PKcs inhibitor as a synthetic lethality strategy in BRCA1-BER deficient cells.

(A) Inhibitors alone. (B) Cisplatin in combination with inhibitors. (BRCA1+/BER+) BRCA-BER proficient cells and (BRCA-/BER-) BRCA1-BER deficient cells.
Chapter 7

General discussion and suggestions for future studies
7. General Discussion and suggestions for future studies

Maintenance of genome stability is of primary importance for living organisms. However, DNA is under constant attack from numerous endogenous and exogenous DNA damaging agents. The accumulation of DNA damage can induce permanent changes that lead to carcinogenesis and could promote aggressive cancer biology. To counteract the deleterious effects of DNA damage, all organisms are equipped with DNA repair machinery to process DNA damage and maintain genomic integrity. Cancer cells on the other hand acquire defects in a certain DNA repair pathway and become dependent on a compensatory repair pathway in order to survive. More recently, inhibitions of the compensatory DNA repair (backup repair) pathway has emerged as a promising target for anticancer therapy. This strategy of selective killing of cancer cells is known as synthetic lethality (Abbotts et al., 2014b, Rassool and Tomkinson, 2010, Lord and Ashworth, 2008). Synthetic lethality is defined as a genetic combination of mutations in two or more genes/pathways that leads to cell death, whereas a mutation in only one of them does not. This represents a new approach of targeting DNA repair pathways with cancer therapies based on the genetic background of the tumour (Lord and Ashworth, 2008). The well known synthetic lethality relationship to date is targeting BRCA mutation with PARP inhibitors. BRCA/PARP synthetic lethality model proposes that targeting BRCA deficient tumours, which are HR deficient, with PARP inhibitor leads to the accumulation of SSBs. This is followed by the formation of DSBs during DNA replication, leading to cell death (Farmer et al., 2005, Bryant et al., 2005). On 23 October 2014, PARP
inhibitor (Olaparib, AstraZeneca©) has moved a step closer to being available for patients in the UK, after receiving a positive recommendation from the Committee for Medicinal products for Human Use (CHMP), which is part of the European Medicines Agency (EMA). The CHMP recommended that Olaparib (known as Lynparza™) be used as monotherapy for adult patients with relapsed, platinum-sensitive epithelial ovarian, fallopian tube or primary peritoneal cancer with mutations in one of two genes BRCA1 or BRCA2, and who have responded to platinum-based chemotherapy (EMA, 2014). This makes Lynparza™ the first in class drug depending on synthetic lethality strategy.

Clinical data provide compelling evidence that alterations in expression of DNA repair factors may have prognostic and predictive significance in patients. Germ-line mutations in BRCA1 predispose to breast and ovarian cancer development. BRCA1 mutation carriers have an 80% lifetime risk of developing breast cancer and 40-50% risk of developing ovarian cancer (Risch et al., 2001). In addition, BRCA1 mutations has been found to increase the risk of pancreatic and prostate cancers (Thompson and Easton, 2002). A more recent study suggested that BRCA1 mutation increases the risk of other types of cancer such as kidney, leukemia, lung, colorectal, myeloma, melanoma, stomach, thyroid and uterine cancers (Mersch et al., 2014). BRCA1 functions to maintain genomic stability through the assembly of multiple protein complexes involved in DNA repair pathways. BRCA deficiency on the other hand has been linked to genomic instability related to a defect in double strand break repair (Moynahan et al., 1999, Kass et al., 2013).
Alternative synthetic lethality target is an area of intense research. Our group has shown that targeting APE1, a central component of the BER pathway is synthetically lethal in HR deficient cells (Mohammed et al., 2011, Sultana et al., 2012). Similarly, targeting APE1 in PTEN deficient melanoma cell lines also shows synthetic lethality (Abbotts et al., 2014a). In addition, targeting XRCC1 deficient cell lines by ATM, DNA-PKcs or SSB repair inhibitor of ATR also induces synthetic lethality (Sultana et al., 2013a, Sultana et al., 2013b). Recently, BRCA1 has been suggested to be involved in the transcriptional regulation of three BER enzymes; OGG1, NTH1 and APE1 (Saha et al., 2010). A more recent study supports these observations and demonstrated down-regulation of APE1, NTH1, OGG1 and XRCC1 in BRCA1 mutated and sporadic breast cancers (De Summa et al., 2014). Therefore impaired BER in BRCA1 deficient cell lines may be an opportunity for novel synthetic lethality strategy. The primary aim of the work described in this thesis was to investigate whether targeting BRCA1-BER deficient cells using ATM or DNA-PKcs inhibitors will be synthetically lethal. In order to achieve the aim, firstly DNA repair gene and protein expression in BRCA1 deficient and BRCA1 proficient cells were investigated (Chapter 3). Secondly, BRCA1 deficient cells were targeted by ATM inhibitors (KU55933 and KU60019) to investigate whether they would be synthetically lethal compared to BRCA1 proficient cells (Chapter 4). Then, BRCA1 deficient cells were targeted by DNA-PKcs inhibitors (NU7441 and NU7026) to investigate if they would also be synthetically lethal (Chapter 5). Finally, BRCA1 deficient cells were targeted by ATM or DNA-PKcs inhibitor in combination with cisplatin to
investigate whether it would enhance the observed synthetic lethality (Chapter 6).

Initially the RT² Profiler™ DNA Repair PCR Array was used to scan 84 DNA repair genes. Data demonstrated down-regulation of several DNA repair mRNAs in BRCA1 mutant/knockdown cell lines as compared to BRCA1 proficient cell lines. Low mRNA expression was demonstrated for APE1, OGG1 and NTHL1, which is consistent with previous observations (Saha et al., 2010, De Summa et al., 2014). However RT² Profiler™ Array results did not always give statistical significance. Further assessments of the results were carried out using quantitative real time PCR (RT-qPCR) for a panel of selected DNA repair genes (APE1, XRCC1, SMUG1, Pol β, Lig3, ATM and DNA-PKcs) which had previously be linked to BRCA1 (Abdel-Fatah et al., 2013, Masaoka et al., 2013, De Summa et al., 2014). RT-qPCR demonstrated statistically significant down-regulation of the selected genes. Protein expression was then analyzed for this selected group by western blot, which showed down-regulation consistent with the mRNAs expression. Taken together, these results suggest that BRCA1 deficiency may associate with a global defect in BER pathway. Interestingly, BRCA1 expression was also associated with a global reduction in HR (see section 3.4.2), NHEJ (see section 3.4.3), NER and MMR pathways (see section 3.4.4). However, further investigation is needed to measure the efficiency of these pathways.

As BRCA1 deficient cells were suggested to be BER deficient, cells may be amenable to synthetic lethality approach by targeting additional DSB repair targets. To examine this, clonogenic survival assays were conducted first to evaluate the cytotoxicity of ATM inhibitors (KU55933 and KU60019) against
BRCA1 knockdown HeLa cells and BRCA1 mutated MDA-MB-436 cells, using Control HeLa and MCF7 cells as BRCA1 proficient controls, respectively. Data demonstrated significant cytotoxicity in BRCA1 mutant/knockdown cell lines. For additional validation, the proliferation assay (MTS assay) was used to determine cell growth in response to ATM inhibitor, which showed consistent results with the clonogenic assay. Several functional studies were conducted to understand the mechanism behind this selective cytotoxicity. γH2AX immunofluorescence microscopy assay showed that BRCA1 deficient cell lines treated with ATM inhibitors (KU55933 or KU60019) for 48 hours were associated with statistically significant increase in the accumulation of DSBs (γH2AX foci) compared to untreated BRCA1 deficient cells. Cell cycle progression was arrested significantly at G2/M phase in treated BRCA1 HeLa cells compared to untreated BRCA1 HeLa cells. On the other hand, MDA-MB-436 cells showed a modest increase in G2/M arrest after 48 hours treatments which suggests a disturbance in cell cycle progression, though different time points should be tested in the future. Increases in G2/M arrest supports previously published data showing G2/M arrest after ATM inhibition (Sultana et al., 2013a, Lee et al., 2013, Nadkarni et al., 2012, Jazayeri et al., 2006). A flow cytometry-based apoptosis detection assay was performed after that and results demonstrated an increase in apoptosis rate in treated BRCA1 deficient cells compared to untreated BRCA1 deficient cells. These results support those previously published showing ATM inhibitor KU55933 treatment triggers apoptosis in different cancer cell lines (Sultana et al., 2013a, Li and Yang, 2010, Korwek et al., 2012) which again highlighted the importance of ATM in DNA repair and cell survival.
Altogether, these results support the hypothesis of BRCA1/ATM synthetic lethality and suggest that ATM plays a vital and irreplaceable role in genomic stability in BRCA1 deficient cells.

In light of the observed BRCA1/ATM synthetic lethality relationship, targeting BRCA1 deficiency by DNA-PKcs inhibitors (NU7441 and NU7026) was also carried out to investigate if there was a synthetically lethal relationship. Clonogenic survival assays results in Chapter 5 demonstrated that upon DNA-PKcs inhibition, BRCA1 mutant/knockdown cell lines showed increased cellular toxicity compared to BRCA1 proficient cell lines. For additional validation, MTS assay was used to determine cell viability in response to DNA-PKcs inhibitor. Both NU7441 and NU7026 inhibitors induced consistent results with the clonogenic assay. DSBs formation increased significantly after DNA-PKcs inhibition in BRCA1 mutant/knockdown cell lines. This finding supports previous publications showing accumulation of γH2AX foci with inhibition of DNA-PKcs (Shaheen et al., 2011, Sultana et al., 2013a).

Monitoring cell cycle progression in BRCA1 deficient cell lines after treatment with DNA-PKcs inhibitors for 48 hours was associated with a modest arrest at G1 phase compared to untreated BRCA1 deficient cell lines. This suggests further investigation with different time points is required. The data demonstrate that a potential synthetic lethality relationship which exists between BRCA1 deficiency and DNA-PKcs inhibition, and highlights the role of DNA-PKcs in the maintenance of genomic stability of BRCA1 deficient cells.

Finally, combination strategy was investigated to demonstrate if targeting BRCA1 deficiency by ATM or DNA-PKcs inhibitor in combination with
cisplatin would enhance this synthetic lethality. Clonogenic combination results in Chapter 6 of both treatments (cisplatin/ATM inhibitor and cisplatin/DNA-PKcs inhibitor) induced hypersensitivity in BRCA1 deficient cells compared to BRCA1 deficient cells treated with cisplatin alone. Functional studies demonstrated that cisplatin enhanced cell death by increased accumulation of DSBs, G2/M cell cycle arrest and increased induction of apoptosis compared to either inhibitor alone. These data suggest that cisplatin can enhance the observed synthetic lethality effect.

Interestingly, it has always been assumed that cells lacking functional BRCA1 protein have impaired HR, and thus are suggested to depend on the more error-prone non-homologous end joining (NHEJ) pathway (Huen et al., 2010). In this thesis BRCA1 deficiency has been associated with a global impairment of expression of HR and NHEJ factors. Targeting NHEJ by DNA-PKcs inhibitors or HR by ATM inhibitors in BRCA1 deficient cells showed synthetic lethality. This observation supports that both DSB sub-pathways play an important role in maintaining chromosomal stability (Takata et al., 1998). Moreover, in BRCA1 proficient cells with functional BER, despite ATM or DNA-PKcs inhibition, DSBs may be rapidly repaired in backup DNA repair pathways and cells may continue to survive. The backup repair could operate through the complex overlap between HR, NHEJ and alternative non-homologous end joining (A-NHEJ) pathways (Schipler and Iliakis, 2013, Chapman et al., 2012). On the other hand, in BRCA1 deficient cells with low BER, ATM or DNA-PKcs inhibitors may lead to accumulation of DSBs which beyond a threshold, leading cells to cell cycle arrest and apoptosis. Recently our group has shown that ATM or DNA-PKcs inhibitors are synthetically lethal in XRCC1 deficient
cells (Sultana et al., 2013a). Given the potential role of XRCC1 in the A-NHEJ (Mladenov and Iliakis, 2011, Brandsma and Gent, 2012, Dueva and Iliakis, 2013), BRCA1 deficient cells with low XRCC1 may have impaired A-NHEJ in addition to BER resulting in an increased genomic instability and enhanced synthetic lethality when cisplatin was added in the current study. Moreover, ATM and DNA-PKcs modulation has been shown to enhance cisplatin cytotoxicity (Yoshida et al., 2008, Dejmek et al., 2009).

A limitation of my study is that this investigation was restricted to a few cell lines only. For future work, use of positive control drugs (e.g. several PARP inhibitors) in the functional studies will validate the results further. As with most research, the data presented here have thrown up as many questions as answers. Future in vivo studies are required to test these new findings and to evaluate the toxicity in normal tissue, which will validate the potential therapeutic benefit in clinical use. Another important area of investigation would be to evaluate potential biological mechanisms of resistance to ATM and DNA-PKcs inhibition. Moreover, the findings described here would also need further validation using pharmaceutical grade ATM and DNA-PKcs inhibitors that are currently under development. Also, to investigate if BRCA1 overexpression increases enzymatic activities related to DNA repair pathways (e.g. BER) a knock-in method using a BRCA1 expression plasmid created by cloning the BRCA1 cDNA into a vector could be used. In addition, ATM and DNA-PKcs knockdown cell lines can be targeted by BRCA1 small interfering (si) RNA to study the synthetic lethality relationship further.

This study suggests that BRCA1 deficiency may be associated with a global defect in DNA repair pathways, in particular base excision repair. The study
also provides proof that BRCA1 deficient cells could be targeted by inhibitors of the double strand break repair pathway, proposing that drug targets other than PARP can also be used for personalized cancer therapy.

7.1. Summary of key findings

✓ Loss of BRCA1 expression is associated with reduction in mRNA and protein expression of factors involved in Base excision repair.

✓ In cell lines, loss of BRCA1 expression is also associated with reduction in the expression of factors involved in Homologous recombination and Non-homologous end joining.

✓ BRCA1 mutated and knockdown cell lines are sensitive to ATM inhibitors (KU55933 & KU60019).

✓ ATM inhibitors treatment is associated with DNA double strand break accumulation and induction of apoptosis in BRCA1 deficient cells.

✓ BRCA1 mutated and knockdown cell lines are sensitive to DNA-PKcs inhibitors (NU7441 & NU7026).

✓ DNA-PKcs inhibitors treatment is associated with DNA double strand break accumulation and induction of apoptosis in BRCA1 deficient cells.

✓ Cisplatin in combination with ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) enhances cytotoxicity in BRCA1 deficient cells.

✓ ATM inhibitor (KU55933) or DNA-PKcs inhibitor (NU7441) in combination with cisplatin treatment is associated with DNA double strand break accumulation, G2/M arrests and induction of apoptosis in BRCA1 deficient cells.
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Appendices
APPENDIX A. ΔΔCt Method

ΔΔCt method is a convenient way to analyze the relative changes in gene expression from real-time quantitative PCR experiments. This involves comparing the Ct values of the samples (BRCA1 deficient cells) of interest with a control (BRCA1 proficient cells). The Ct values of both the control and the samples of interest are normalized to a housekeeping gene (GAPDH). The ΔΔCt method is also known as the comparative Ct method and 2−[delta][delta]Ct method (Livak and Schmittgen, 2001), where

\[
\Delta \text{Ct}_1 = \text{Ct} (\text{target, control sample}) - \text{Ct} (\text{ref, control sample}) \\
\Delta \text{Ct}_2 = \text{Ct} (\text{target, deficient sample}) - \text{Ct} (\text{ref, deficient sample}) \\
\Delta \Delta \text{Ct} = \Delta \text{Ct}_1 - \Delta \text{Ct}_2
\]

Where

Ct (target, control sample) = Ct value of gene of interest in BRCA1 proficient sample
Ct (ref, control sample) = Ct value of housekeeping gene in BRCA1 proficient sample
Ct (target, deficient sample) = Ct value of gene of interest in BRCA1 deficient sample
Ct (ref, deficient sample) = Ct value of housekeeping gene in BRCA1 deficient sample

Then calculate the ratio of our target gene in our deficient sample relative to our control sample by taking \(2^{\Delta \Delta \text{Ct}}\).

Example:

<table>
<thead>
<tr>
<th></th>
<th>control sample</th>
<th>deficient sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRCA1</td>
<td>31.7</td>
<td>35.50</td>
</tr>
<tr>
<td>GAPDH</td>
<td>24.14</td>
<td>21.847</td>
</tr>
</tbody>
</table>

\[
\Delta \Delta \text{Ct} = (31.7 - 24.14) - (35.50 - 21.847) \\
\Delta \Delta \text{Ct} = (7.56) - (13.653) \\
\Delta \Delta \text{Ct} = - 6.093 \\
2^{\Delta \Delta \text{Ct}} = 2^{(0 - 6.093)} = 0.02 \text{ (Fold Change)}
\]

So BRCA1 gene is decreased by 0.02 times in BRCA1 deficient sample versus BRCA1 proficient sample.
## APPENDIX B. DNA repair genes assayed in Qiagen RT² Profiler DNA Repair

<table>
<thead>
<tr>
<th><strong>Symbol</strong></th>
<th><strong>Description</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>APE1</td>
<td>APEX nuclease (multifunctional DNA repair enzyme) 1</td>
</tr>
<tr>
<td>APE2</td>
<td>APEX nuclease (apurinic/apyrimidinic endonuclease) 2</td>
</tr>
<tr>
<td>ATM</td>
<td>Ataxia telangiectasia mutated</td>
</tr>
<tr>
<td>ATR</td>
<td>Ataxia telangiectasia and Rad3 related</td>
</tr>
<tr>
<td>ATXN3</td>
<td>Ataxin 3</td>
</tr>
<tr>
<td>BRCA1</td>
<td>Breast cancer 1, early onset</td>
</tr>
<tr>
<td>BRCA2</td>
<td>Breast cancer 2, early onset</td>
</tr>
<tr>
<td>BRIP1</td>
<td>BRCA1 interacting protein C-terminal helicase 1</td>
</tr>
<tr>
<td>CCNH</td>
<td>Cyclin H</td>
</tr>
<tr>
<td>CCNO</td>
<td>Cyclin O</td>
</tr>
<tr>
<td>CDK7</td>
<td>Cyclin-dependent kinase 7</td>
</tr>
<tr>
<td>DDB1</td>
<td>Damage-specific DNA binding protein 1, 127kDa</td>
</tr>
<tr>
<td>DDB2</td>
<td>Damage-specific DNA binding protein 2, 48kDa</td>
</tr>
<tr>
<td>DMC1</td>
<td>DMC1 dosage suppressor of mck1 homolog, meiosis-specific homologous recombination (yeast)</td>
</tr>
<tr>
<td>ERCC1</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 1</td>
</tr>
<tr>
<td>ERCC2</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 2</td>
</tr>
<tr>
<td>ERCC3</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 3</td>
</tr>
<tr>
<td>ERCC4</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 4</td>
</tr>
<tr>
<td>ERCC5</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 5</td>
</tr>
<tr>
<td>ERCC6</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 6</td>
</tr>
<tr>
<td>ERCC8</td>
<td>Excision repair cross-complementing rodent repair deficiency, complementation group 8</td>
</tr>
<tr>
<td>EXO1</td>
<td>Exonuclease 1</td>
</tr>
<tr>
<td>FEN1</td>
<td>Flap structure-specific endonuclease 1</td>
</tr>
<tr>
<td>LIG1</td>
<td>Ligase I, DNA, ATP-dependent</td>
</tr>
<tr>
<td>LIG3</td>
<td>Ligase III, DNA, ATP-dependent</td>
</tr>
<tr>
<td>LIG4</td>
<td>Ligase IV, DNA, ATP-dependent</td>
</tr>
<tr>
<td>MGMT</td>
<td>O-6-methylguanine-DNA methyltransferase</td>
</tr>
<tr>
<td>MLH1</td>
<td>MutL homolog 1, colon cancer, nonpolyposis type 2 (E. coli)</td>
</tr>
<tr>
<td>MLH3</td>
<td>MutL homolog 3 (E. coli)</td>
</tr>
<tr>
<td>MMS19</td>
<td>MMS19 nucleotide excision repair homolog (S. cerevisiae)</td>
</tr>
<tr>
<td>MPG</td>
<td>N-methylpurine-DNA glycosylase</td>
</tr>
<tr>
<td>MRE11A</td>
<td>MRE11 meiotic recombination 11 homolog A (S. cerevisiae)</td>
</tr>
<tr>
<td>MSH2</td>
<td>MutS homolog 2, colon cancer, nonpolyposis type 1 (E. coli)</td>
</tr>
<tr>
<td>MSH3</td>
<td>MutS homolog 3 (E. coli)</td>
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<tr>
<td>MSH4</td>
<td>MutS homolog 4 (E. coli)</td>
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<td>MSH5</td>
<td>MutS homolog 5 (E. coli)</td>
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<td>MSH6</td>
<td>MutS homolog 6 (E. coli)</td>
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<tr>
<td>MUTYH</td>
<td>MutY homolog (E. coli)</td>
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<tr>
<td>NEIL1</td>
<td>Nei endonuclease VIII-like 1 (E. coli)</td>
</tr>
<tr>
<td>Gene</td>
<td>Description</td>
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<td>NEIL2</td>
<td>Nei endonuclease VIII-like 2 (E. coli)</td>
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<td>NEIL3</td>
<td>Nei endonuclease VIII-like 3 (E. coli)</td>
</tr>
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<td>NTHL1</td>
<td>Nth endonuclease III-like 1 (E. coli)</td>
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<td>OGG1</td>
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<tr>
<td>PARP1</td>
<td>Poly (ADP-ribose) polymerase 1</td>
</tr>
<tr>
<td>PARP2</td>
<td>Poly (ADP-ribose) polymerase 2</td>
</tr>
<tr>
<td>PARP3</td>
<td>Poly (ADP-ribose) polymerase family, member 3</td>
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<tr>
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<td>PMS1 postmeiotic segregation increased 1 (S. cerevisiae)</td>
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<tr>
<td>PMS2</td>
<td>PMS2 postmeiotic segregation increased 2 (S. cerevisiae)</td>
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<tr>
<td>PNKP</td>
<td>Polynucleotide kinase 3'-phosphatase</td>
</tr>
<tr>
<td>POLB</td>
<td>Polymerase (DNA directed), beta</td>
</tr>
<tr>
<td>POLD3</td>
<td>Polymerase (DNA-directed), delta 3, accessory subunit</td>
</tr>
<tr>
<td>POLL</td>
<td>Polymerase (DNA directed), lambda</td>
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<tr>
<td>DNA-PKcs</td>
<td>Protein kinase, DNA-activated, catalytic polypeptide</td>
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<td>RPA1</td>
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<tr>
<td>RPA3</td>
<td>Replication protein A3, 14kDa</td>
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<td>STE20-like kinase</td>
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<td>SMUG1</td>
<td>Single-strand-selective monofunctional uracil-DNA glycosylase 1</td>
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<tr>
<td>TDG</td>
<td>Thymine-DNA glycosylase</td>
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<td>TOP3A</td>
<td>Topoisomerase (DNA) III alpha</td>
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<tr>
<td>TOP3B</td>
<td>Topoisomerase (DNA) III beta</td>
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<td>TREX1</td>
<td>Three prime repair exonuclease 1</td>
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<td>XPA binding protein 2</td>
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<tr>
<td>XPC</td>
<td>Xeroderma pigmentosum, complementation group C</td>
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<td>XRCC1</td>
<td>X-ray repair complementing defective repair in Chinese hamster cells 1</td>
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<td>XRCC2</td>
<td>X-ray repair complementing defective repair in Chinese hamster cells 2</td>
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<td>XRCC3</td>
<td>X-ray repair complementing defective repair in Chinese hamster cells 3</td>
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<td>XRCC4</td>
<td>X-ray repair complementing defective repair in Chinese hamster cells 4</td>
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<td>XRCC5</td>
<td>X-ray repair complementing defective repair in Chinese hamster cells 5 (double-strand-break rejoining)</td>
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APPENDIX C. Other related DNA repair genes under-expressed in BRCA1 deficient cell lines compared to BRCA1 proficient cell lines.

Statistical significance determined using the Bonferroni correction for multiple testing, with p values less than 0.05 (threshold $p < 0.000595 = 0.05/84$).

<table>
<thead>
<tr>
<th>Other related DNA repair genes</th>
<th>Fold Change</th>
<th>P Value</th>
<th>Fold Change</th>
<th>P Value</th>
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<tr>
<td><strong>BRCA1 knockdown cell</strong></td>
<td></td>
<td></td>
<td><strong>MDA-MB-436 cells compared to MCF7 cells</strong></td>
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<td>ATR</td>
<td>0.8129</td>
<td>0.622017</td>
<td>0.1521</td>
<td>0.00402</td>
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<td>EXO1</td>
<td>0.7888</td>
<td>0.572304</td>
<td>0.143</td>
<td>0.002984</td>
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<td>MGMT</td>
<td>0.3817</td>
<td>0.022209</td>
<td>0.2671</td>
<td>0.043123</td>
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<tr>
<td>RAD18</td>
<td>0.7589</td>
<td>0.511457</td>
<td>0.2366</td>
<td>0.02729</td>
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<td>RFC1</td>
<td>0.7687</td>
<td>0.531243</td>
<td>0.4667</td>
<td>0.242016</td>
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<td>TOP3A</td>
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<td>0.450213</td>
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<td>TOP3B</td>
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<td>0.312452</td>
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<td>MRE11A</td>
<td>0.7363</td>
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<td>0.422161</td>
<td>0.3424</td>
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<td>TDG</td>
<td>1.1639</td>
<td>0.717939</td>
<td>0.147</td>
<td>0.003411</td>
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</tbody>
</table>

APPENDIX D. Evaluation of drug interaction

Isobologram, combination index and curve shift analysis are useful methods for analyzing drug-drug interaction, and provide complimentary information.

**Isobologram analysis:** The isobologram analysis provides a graphical presentation of the nature of interaction of two drugs (e.g. drug A and drug B). Firstly, in a two-coordinate plot with one coordinate representing concentration of drug A ($IC_{50(A)}$) and the other representing concentration of drug B ($IC_{50(B)}$), the concentrations of drugs A and B required to produce a defined effect x,
when used as single agents, are placed on the X and Y axes, respectively. The line of additivity is constructed by connecting these two points. Secondly, the concentrations of the two drugs used in combination to provide the same effect x, denoted by point \((C_{(A, x)}, C_{(B, x)})\), are placed in the same plot. Synergy, additivity, or antagonism is indicated when this point is located below, on, or above the line, respectively.

**Combination index analysis:** Combination index (CI) provides a quantitative measure of the extent of drug interaction at a given effect level. That is, the combination concentrations of first drug (A) and second drug (B) to produce an effect (D), combination drugs are normalized by their corresponding concentrations that produces the same effect as a single agent (Chou and Talalay, 1984). This is a useful method for analysing drug-drug interactions (Zhao et al., 2010). The effect of the combined treatment is analysed for the combination of drugs by applying the following CI equation: 

\[(Ac/Ae) + (Bc/Be) = D,\]

where Ac and Bc correspond to the concentrations of drugs used in the combination treatment, and Ae and Be corresponds to the concentrations of drugs able to produce the same effect by themselves. CI is considered synergistic if D (combination index) is < 1, whereas considered additive if D = 1 or antagonistic if D is > 1.

**Curve shift analysis:** Curve shift analysis allows simultaneous presentation of the studied concentration-effect curves of single agent and combination treatments in a single plot. The effects of combination therapy can be analysed by applying the following equation:

\[
\text{Combination Therapy Effect} = \frac{E_{\text{max}} \left( \frac{C_{A,x}}{IC_{50,A}} + \frac{C_{B,x}}{IC_{50,B}} \right)^{n_{\text{combo}}}}{\left( \frac{C_{A,x}}{IC_{50,A}} + \frac{C_{B,x}}{IC_{50,B}} \right)^{n_{\text{combo}}} + (IC_{50,\text{combo}})^{n_{\text{combo}}}}
\]

Where \(E_{\text{max}}\) is the full range of drug effect; \(C_{A,x}\) or \(C_{B,x}\) is the IC\(_{50}\)-equivalent concentration of drug A or B in combination; IC\(_{50}\) is the drug concentration producing the median effect of 50%; and n is the curve shape parameter describing the steepness of the concentration-effect relationship; IC\(_{50, X}\) is the IC\(_{50}\) value of drug X; IC\(_{50,\text{combo}}\) and \(n_{\text{combo}}\) are the values for the combination therapy.

Plotting the effects of both single agents and combinations against (IC\(_{50}\)-equivalent drug concentrations) enables the simultaneous presentation of these concentration effect curves in a single plot. Due to the normalization, the curves for the single agents will have an IC\(_{50}\) value of one, while synergistic combinations will have a lower IC\(_{50}\) value resulting in a leftward shift, and antagonistic combinations will show a rightward shift (Zhao et al., 2010).