

REVIEW

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DNA folds threaten genetic stability and can be leveraged for chemotherapy

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Damaging DNA is a current and efficient strategy to fight against cancer cell proliferation. Numerous mechanisms exist to counteract DNA damage, collectively referred to as the DNA damage response (DDR) and which are commonly dysregulated in cancer cells. Precise knowledge of these mechanisms is necessary to optimise chemotherapeutic DNA targeting. New research on DDR has uncovered a series of promising therapeutic targets, proteins and nucleic acids, with application notably *via* an approach referred to as combination therapy or combinatorial synthetic lethality. In this review, we summarise the cornerstone discoveries which gave way to the DNA being considered as an anticancer target, and the manipulation of DDR pathways as a valuable anticancer strategy. We describe in detail the DDR signalling and repair pathways activated in response to DNA damage. We then summarise the current understanding of non-B DNA folds, such as G-quadruplexes and DNA junctions, when they are formed and why they can offer a more specific therapeutic target compared to that of canonical B-DNA. Finally, we merge these subjects to depict the new and highly promising chemotherapeutic strategy which combines enhanced-specificity DNA damaging and DDR targeting agents. This review thus highlights how chemical biology has given rise to significant scientific advances thanks to resolutely multidisciplinary research efforts combining molecular and cell biology, chemistry and biophysics. We aim to provide the non-specialist reader a gateway into this exciting field and the specialist reader with a new perspective on the latest results achieved and strategies devised.

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Introduction

High levels of genetic instability and mutations is a general enabling hallmark of cancer, as defined by Douglas Hanahan and Robert A. Weinberg in their cornerstone reviews.^{1,2} This is likely a result of the need for tumour progression to acquire multiple mutations, *e.g.* in oncogenes and tumour-suppressor genes, for successful tumour initiation, and potentially metastasis. The genome maintenance systems limit such genetic defects in healthy cells but this is often impaired in cancer cells, making the accumulation of DNA lesions a common feature in cancer. Cancer cells' reduced ability to repair DNA damage, driving their genomic instability, provides an important vulnerability that is exploited therapeutically.^{3–5}

Recent advances in genetics, genomics and proteomics have provided a better understanding of the complex interplay between DNA lesions and repair mechanisms, and what is therapeutically relevant to halt cancer cell proliferation.⁶ These advances have generated momentum for designing more selective and efficient anticancer strategies, often relying on combinations of DNA damaging agents and DNA repair inhibitors.^{5,7,8} Recent advances in our understanding of the complex aspects of DNA/RNA secondary structure^{9–12} provide brand new opportunities to exploit DNA damage in a more precise manner. Alternative nucleic acid structures are defined as structures that deviate from the canonical Watson–Crick DNA double helix, B-DNA or duplex-DNA. From a molecular viewpoint, the selective small-molecule targeting of such DNA structures holds potential over gene sequence targeting, given that they present well-defined 3D-structures,^{13,14} in a manner reminiscent of modern pharmaceutical targeting of proteins and enzymes.¹⁵ Furthermore, alternative DNA structure formation is coupled with DNA transactions (replication, transcription) due to transient strand separation and local DNA deformation.^{16–18} Such structures thus offer the added advantage of creating replication/transcription-associated DNA damage whose repercussions will be felt primarily in highly

proliferating cells. The present review aims at gathering the most recent results obtained in this new and promising approach, to demonstrate the relevance of targeting higher-order DNA structures to treat cancer.

1. The discovery of DNA as a chemotherapeutic target

1a. Discovering the vulnerability of DNA. It was known that organisms could be mutated or damaged through radiative and chemical attack before what we now call the blueprint of life, the DNA, was discovered.¹⁹ Ionising (IR) and ultraviolet (UV) radiation were known to induce lethal and non-lethal mutation in cells from the 1930s,^{20–22} in a U.S. American push to understand radiation biology for military applications, proceeding the fabrication of atomic weapons. Repair mechanisms in irradiated cells were first described in bacteria independently by two United States laboratories in 1949.^{23,24} The mechanistic basis of this repair process, referred to as enzymatic photoreactivation,²⁵ was elucidated only several years later in light of the discovery of DNA structure in 1953,^{26–28} as well as the description of irradiation-promoted formation of cyclobutane pyrimidine dimers (CPD) *in vitro*²⁹ and the discovery of the DNA photolyase that directly repairs CPD formed by UV-light.^{30,31}

DNA is constantly exposed to damaging agents – “*tens of thousands of DNA lesions per day*”³ – of both endogenous and exogenous origins.^{32,33} Beyond irradiation and cross-linking chemicals, different types of lesion have now been unravelled, which include chemical modifications (*e.g.* oxidations, deaminations), DNA replication errors (*e.g.* mismatched Watson–Crick base pairs) and DNA transaction impediments (*e.g.* non-B helix structured DNA, sometimes called ‘difficult-to-replicate’ sequences^{34,35}) which can stall replication and transcription (Fig. 1). The plurality of DNA damage explains why cells have evolved such a highly sophisticated and efficient network of surveillance, signalling and repair pathways, collectively referred to as the DNA damage response (DDR).^{3,4,33,36,37} The



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and in 2018 he was awarded the CNRS Bronze Medal for his discoveries. Starting in 2021, he will lead the “Deciphering and Drugging DNA Repair” team at IPBS (Toulouse, France).



David Monchaud

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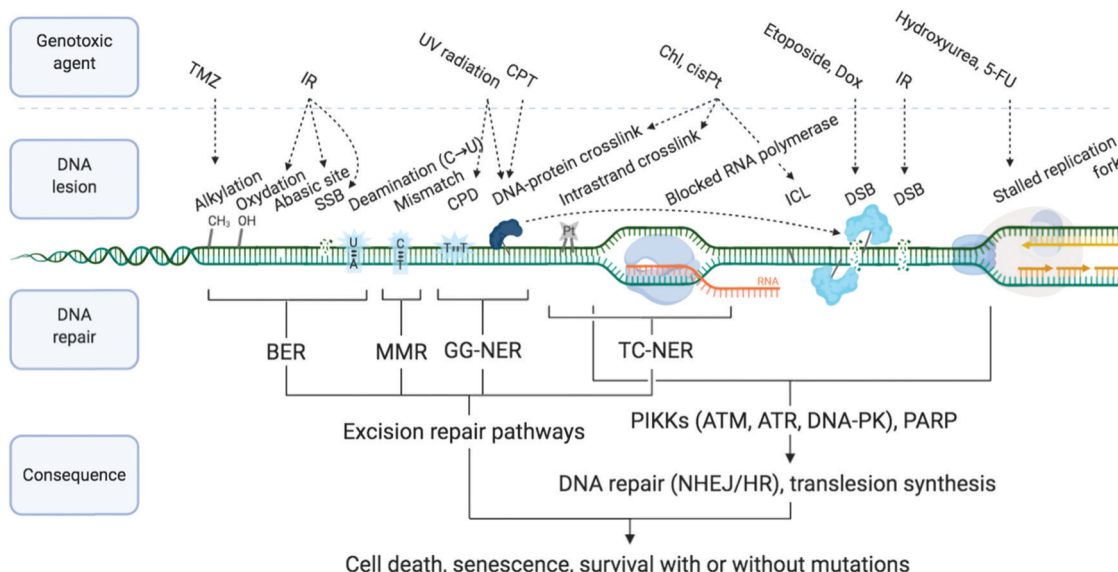


Fig. 1 Examples of DNA-damaging agents, the lesion which they induce, and the downstream consequence. Temozolomide (TMZ), ionising radiation (IR), camptothecin (CPT), chlorambucil (Chl), cisplatin (cisPt), doxorubicin (Dox), single-strand break (SSB), double strand break (DSB), cyclobutane pyrimidine dimer (CPD), interstrand cross-link (ICL), base excision repair (BER), mismatch repair (MMR) and global genome nucleotide excision repair (GG-NER), transcription-coupled NER (TC-NER). Adapted from ref. 70, created with BioRender.

molecular complexity of the DDR is just starting to be understood, because of the multiplicity of pathways involved, including excision (BER, NER, MMR), recombination (HR) and joining (NHEJ) pathways (see Sections 2a and b) and their constant and intricate cross-talk. DDR ultimately aims at controlling genomic instability in normal cells, whilst from a therapeutic viewpoint, the DDR presents a key strategic Achille's heel of cancer cells, which can be targeted to impede their anarchic proliferation.

1b. Damaging DNA to treat cancer. Cancer has likely existed since the beginning of mankind but ageing and the constant modification of our lifestyles has dramatically increased its occurrence. Treatment consisted solely of surgery up until radiotherapy treatments were introduced, with the first uses of X-rays in 1897 and of radium in 1904, following the discoveries of physicists Wilhelm C. Rontgen and Marie Skłodowska Curie respectively.^{38,39} Unknown at the time, radiotherapy exploits the principle of inducing DNA damage, including DNA double strand breaks (DSBs) that show moderate selectivity in killing rapidly dividing cells. The first chemotherapy approaches again targeted DNA, yet still in a somewhat empirical manner due to the lack of mechanistic details on how DNA damage is created and repaired. The chemical arms-race headed by the Germans from the beginning of WWI (1914) led to the military use of mustard gas (bis(2-chloroethyl)sulfide),^{40,41} along with a wide insight into DNA-targeting agents.^{42,43} DNA cross-linking molecules such as mustard gas exert cytotoxic effects that resembled those triggered by irradiations.⁴⁴ Shrouded in secrecy, Alfred Gilman and Louis S. Goodman at Yale University made the link between the leukopenic effects (drop in white blood cell count) of low doses of mustard gas and the potential for lymphosarcoma treatment, which was transposed to a single undocumented clinical trial in 1942.⁴⁵ These crude results from 'personal recollection' of a 'misplaced chart' provided the first chemotherapy proof-of-concept,⁴⁶ giving way

to an extended study that was communicated after the end of WWII.^{41,47,48} Elucidation of the cross-linking mechanism,⁴⁹ again driven by military interests to conceive possible antidotes to chemical weapons, fuelled numerous studies that built the foundations for modern chemotherapy of new DNA repair mechanisms.

These results gave new impetus to the quest for small molecule therapy. Similarly to Gilman and Goodman, Sidney Farber made the link between the high lymphocyte levels observed in leukaemia patients' blood and potentially leukopenic vitamins antifolates, who began pioneering trials on last-chance child leukaemia patients with synthetic antifolates in 1947.^{48,50,51} These research efforts led to the first example of targeted therapy with the use of 5-fluorouracil (5-FU) in 1957,⁵² which specifically targets the abnormal uracil dependence of hepatomas.⁵³ The benchmark example of targeted therapy, imatinib (Gleevec) was reported in 1996,⁵⁴ being the first approved anticancer drug molecularly designed for its specific protein target, the constitutively active BCR-ABL tyrosine kinase produced by the Philadelphia chromosome in chronic myelogenous leukaemia.⁵⁵

The most defining characteristic of cancer cells is their uncontrolled growth, and this is exploited by antiproliferative agents. This class of chemicals encompasses the oldest and still most popular strategy of chemotherapy. Such agents now include several major subgroups (Fig. 1): chemicals which directly alkylate the DNA such as bifunctional cross-linking agents (nitrogen mustards including chlorambucil (Chl) and platinum drugs including cisplatin (cisPt), carboplatin and oxaliplatin) which cause intrastrand cross-links (and less frequently interstrand and DNA–protein cross-links)^{56,57} and monofunctional alkylators such as temozolomide (TMZ); topoisomerase poisons such as etoposide, which intercalates at the



breakage site formed as DNA topoisomerase 2 (TOP2) induces a DSB to disentangle DNA, thus trapping the DNA-TOP2 cleavage complex,^{58–60} and camptothecin (CPT) and derivatives (topotecan, irinotecan), which similarly trap the DNA-topoisomerase 1 (TOP1) complex;^{61,62} antimetabolites (inhibitors of nucleotide metabolism such as folate antagonist methotrexate, and 5-FU);⁶³ and antimitotic agents such paclitaxel that impedes progression through mitosis by targeting tubulin.

It is widely accepted that cytotoxic agents can attack normal cells as well. In order to treat patients whilst reducing extreme side-effects, drug combinations are often used, accompanied by surgery and radiotherapy of solid tumours. The first combination of DNA targeting agents associated several cytotoxic drugs (e.g. etoposide and cisPt for small-cell lung carcinoma or FOLFIRINOX protocol for metastatic pancreatic cancers).

Another hallmark of cancer cells is their genetic instability, notably caused by an imbalance in DDR mechanisms.^{2,64} Among the most documented examples of oncogenic dysregulations, BRCA1/2 and TP53 mutations rank highly. Mutations in *Breast cancer susceptibility genes* BRCA1 and BRCA2 are associated with breast and ovarian cancers.^{65,66} BRCA1/2 play a critical role in DSB repair by homologous recombination (HR), a key DDR pathway (see Section 2a2). TP53, the gene encoding the apoptosis- and DNA-damage-checkpoint-regulating protein p53, is the most frequently mutated gene in cancer, at approximately half of all cancers.^{67,68} It is classically proposed that these genetic dysregulations are advantageous to tumour progression,² and in a recent study, most cancers analysed from The Cancer Genome Atlas were enriched in mutations coding for DDR proteins.⁶⁹ For this reason, modern therapeutic strategies that target DDR, and combinations of DNA-damaging agents and DDR inhibitors, are evolving rapidly (discussed in Section 2d).

2. The DNA damage response (DDR)

The DNA damage response (DDR) is a broad term that designates the mechanisms involved in the response to all DNA lesions. However it is frequently used in a restrictive manner to designate the mechanisms responding to DNA double-strand breaks (DSB). Indeed, owing to their high cytotoxicity, DSBs

induce a robust response that is initiated by the detection of the lesion by PIKKs and/or PARPs (see Section 2b) and that leads to (1) activation of cell cycle checkpoints that blocks or slow down cell cycle progression at specific boundaries; (2) DSB repair that is principally mediated by HR and NHEJ, and (3) activation of a gene expression program that dictates long term response. The DDR can result in cell survival when the amount and type of DSBs is manageable, or in senescence or cell death when damage is too severe (Fig. 2).^{3,71,72} DNA repair mechanisms can be faithful or instead fix mutations in the genome. While DSB-induced genetic variations can be deleterious, with for example the formation of translocations that can be responsible for secondary cancers, they can also be a desired outcome of several physiological processes. This is the case for antibody diversification by variable (diversity) joining (V(D)J) recombination and class-switch recombination (CSR) in which recombination-activating gene (RAG) nuclease for V(D)J, and activation-induced cytidine deaminase (AID) for CSR promote sites-specific DSBs, and for meiotic recombination which relies on SPO11 for genome-wide DSBs formation to promote recombination between homologous chromosomes.^{73–75} The DDR can be activated by *bona fide* DSBs (leading to ATM and DNA-PKcs activation), by accumulation of single-stranded DNA (leading to ATR activation), by conversion of a DNA lesion into one these structures as a result of defective DNA repair, or by conversion of stalled DNA replication or transcription complexes.

2a. DNA repair mechanisms

2a1. Single-strand breaks and damaged/mismatched nucleotides repair. Tomas Lindahl, Paul L. Modrich and Aziz Sancar were awarded the Nobel prize in chemistry in 2015^{76–78} for their pioneering work describing DNA repair mechanisms, including base excision repair (BER), mismatch repair (MMR) and nucleotide excision repair (NER). Multiple repair mechanisms exist for all types of DNA damage (Fig. 1),^{3,33,79,80} and a large degree of overlap exists within the multiple DNA repair pathways and with double-strand break (DSB) repair pathways.⁶⁹ Single-strand breaks (SSBs) are the most common DNA damage occurring throughout a cell's lifetime, with a large amount being generated endogenously by reactive oxygen species (ROS)

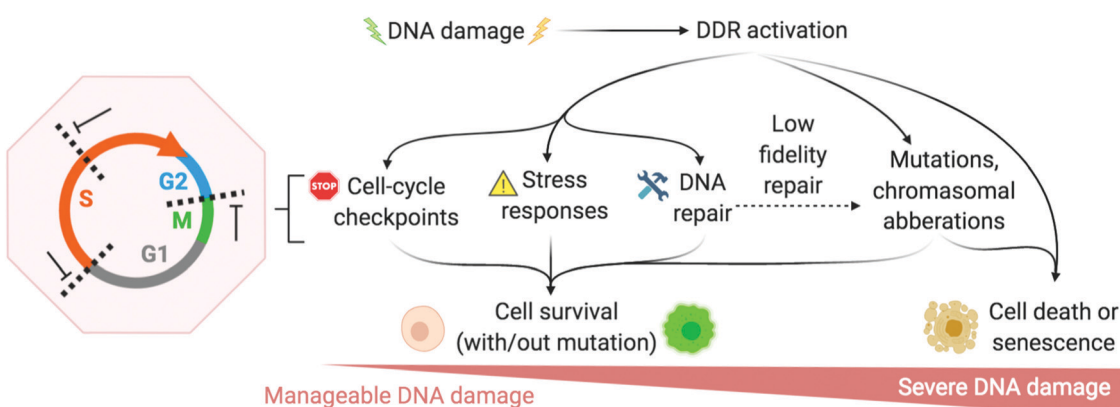


Fig. 2 DNA damage response (DDR) is activated to varying degrees depending on the extremity of DNA damage. Adapted from ref. 72, created with BioRender.



generating SSB directly or resulting in DNA lesions processed by DNA glycosylases and apurinic/apyrimidinic endonucleases (APEs). We provide below a brief description of the SSB repair pathways:

BER: small nucleobase lesions such as alkylations, oxidations and deaminations are repaired by BER, in which a single nucleobase is removed in a concerted dual excision by a DNA glycosylase (e.g. 8-oxoguanine DNA glycosylase 1, OGG1) that specifically recognises the base adduct, and an endonuclease recruited to the abasic site (e.g. APE1). The SSB thereby generated is recognised by the crucial mediator protein PARP-1 that promotes the assembly of several proteins, including SSB repair enzymes, through the formation of poly-ADP ribose chains on local DNA-associated proteins (see Section 2b). Depending on the DNA lesion, short- or long-patch BER will process the damage. In short-patch BER, in which a single nucleotide is repaired, the abasic site is filled by DNA polymerase β and sealed by DNA ligase 3 (Lig3) in complex with the X-ray repair cross-complementing protein 1 (XRCC1).³² In long-patch BER, where 2–12 nucleotides must be replaced due to a bulkier DNA lesion, DNA polymerases (DNA Pol δ , ϵ or β) act to fill the gap in concert with the action of flap endonuclease 1 (FEN1) that processes the resulting flap intermediate and DNA ligase I (Lig1) that seals the nick.⁸⁰

MMR: the mismatch recognition protein complex Mutator S alpha (MutS α), heterodimer of mutS homologues 2 and 6 (MSH2/MSH6), senses Watson–Crick base mismatch protrusions formed from imperfect DNA replication. It moves with DNA replication machinery including the proliferating cell nuclear antigen (PCNA) and DNA Pol δ , although it may or may not be coupled directly, scanning the newly synthesised DNA for mismatches in leading and lagging strands.^{81,82} At a point of damage, the endonuclease complex MutL α , heterodimer of MutL homologue 1 (MLH1) and the post-meiotic segregation increased 2 (PMS2) proteins are recruited and forms a nick on either the 3' or the 5' side of the damage. From this nick, exonucleases, primarily Exo1, remove a stretch of 1 to 2 kilobases (kb) from the damaged strand. The exposed single-stranded DNA (ssDNA) is stabilised by Replication Protein A (RPA) and the gap is filled by DNA replication machinery (PCNA-DNA Pol δ) and ligated by Lig1. New mediators and damage substrates of this pathway are still being discovered, and this pathway partly overlaps with NER, for which reason Sanear once warned against the rigid, over-simplified one-damage-one-pathway classification.⁸³

NER: bulkier, helix-distorting lesions such as CPD, intrastrand cross-links (covalent linkages between two bases of the same strand) and small DNA–protein cross-links (<8–10 kDa)^{61,84,85} are repaired by NER, which is further categorised into transcription-coupled (TC-NER) and global genome (GG-NER) repair pathways, differing in when and where in the cell cycle they occur and in the initial sensor proteins they involve. TC-NER occurs in rhythm with the advancing RNA polymerase (RNAP) and is initiated when the transcription complex is blocked by other lesions.⁸⁶ Unlike TC-NER, GG-NER operates throughout the genome after recognition by the protein sensors *xeroderma pigmentosum* group C (XPC) or

group E (XPE) coupled with the damage DNA binding protein 2 (DDB2). These distinct sensors recruit the same downstream NER mediators: first, a ternary complex that includes the transcription factor IIH (TFIIC) and two helicases, *xeroderma pigmentosum* group B and group D (XPB and XPD) that unwind the damaged DNA and create a ~30 nucleotide bubble; then, an endonuclease complex comprising the *xeroderma pigmentosum* group F (XPF) and excision repair cross-complementation group 1 (ERCC1) proteins incises on the 5' end of the bubble while the *xeroderma pigmentosum* group G (XPG) endonuclease incises at the 3' end, releasing an oligonucleotide carrying the DNA lesion. Like in MMR, the gap is then filled by DNA Pol ϵ and Pol δ , the leading and lagging strand polymerases respectively⁸⁷ and ligated (Lig1).^{78,80,88} GG-NER is not constitutive like TC-NER, but is induced during DNA-damage repair signalling, and is less sensitive to certain DNA adducts than TC-NER.⁷⁰

Direct repair: certain common methylation adducts are reversed in a single step, such as methyl group transfer (O^6 -methylguanine) or oxidative demethylation (1-methyladenine and 3-methylcytosine), by O^6 -meG-DNA methyltransferase (MGMT) and 1meA/3meC-DNA dioxygenase respectively (AlkB homologues).⁸⁹

2a2. Double-strand breaks (DSBs) repair. As compared to SSBs, DSBs pose a more serious threat to genetic integrity since they result in the loss of chromosomal integrity which can lead to the loss of chromosome arms, translocations and cell death. DSBs are highly toxic and a single DSB can result in cell cycle arrest⁹⁰ and cell death.³ DSBs can be generated directly, such as those generated by TOP2 poisoning, ionising radiation, or as a result of the conversion of a primary lesion or blocking structure by transcription or DNA replication.^{91,92} DSBs are repaired by two principal mechanisms (Fig. 3): the non-homologous end joining (NHEJ) pathway, which joins two DNA ends without a DNA template; and the homologous recombination (HR), in which nucleases strip away a stretch of nucleotides at broken ends, exposing a 3' overhang of ssDNA in a process called DNA end resection, then use the corresponding sister chromatid as a template to repair the damaged strand.^{6,37,93}

NHEJ: NHEJ is initiated when the DNA ends are recognised by the Ku complex, the heterodimer of Ku70 and Ku80.³³ Ku shields the DNA ends from exonuclease activities and is a hub for recruiting the other NHEJ factors, including the catalytic subunit of the DNA-dependent protein kinase (DNA-PKcs). Multiple DNA processing enzymes are involved in cleaning the DNA ends for repair by NHEJ, including Artemis (DNA cross-link repair 1C, DCLRE1C), the X polymerases Pol μ and Pol λ , and the polynucleotide kinase/phosphatase (PNKP). Finally, the DNA ends are sealed by the action of the ligation complex composed of XLF-XRCC4-DNA ligase 4 (Lig4). Since NHEJ does not rely on nucleotide sequence matching and it allows short deletions or insertions upon repair, it is considered an error-prone repair mechanism. And yet, sequence fidelity is mostly maintained with little loss in genetic material or chromatin rearrangement, likely thanks to fast recognition of blunt ends of DSB by highly abundant sensor proteins such as Ku, which keeps breaks in close proximity to be repaired.⁹⁴



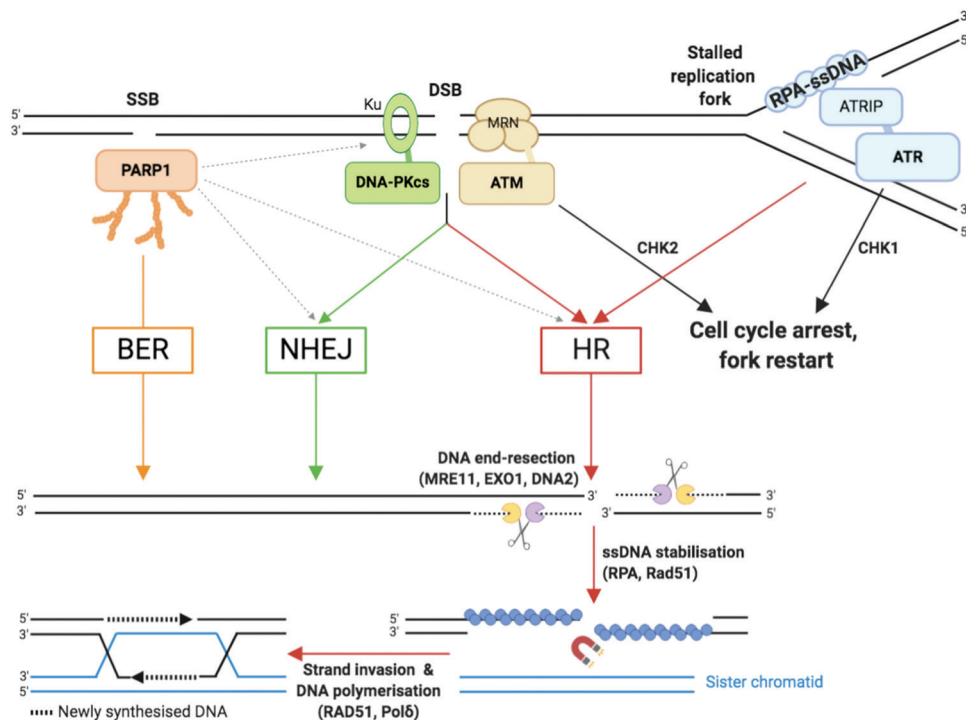


Fig. 3 Single-strand breaks (SSB) and double-strand breaks (DSB) repair pathways. Dotted arrows indicate that PARP1 can be activated by DSB and stimulate DSB repair by both HR and NHEJ.^{33,109} Adapted from ref. 37, created with BioRender.

NHEJ occurs throughout the cell cycle and is the most prevalent DSB repair mechanism in non-dividing cells.

HR: HR is initiated by the generation of a 3' ended ssDNA extension produced by exonucleolytic processing of the DNA ends. This process, called DNA end resection, is controlled by the nuclease activity of MRE11 (Fig. 3). MRE11, as part of the MRE11-RAD50-NBS1 (MRN complex) associates to the chromatin flanking each DSB, through ATP-dependent RAD50 DNA clamping activity. CtIP (C-terminal binding protein 1 (CtBP)-interacting protein, also known as retinoblastoma-binding protein 8, RBBP8), an interaction partner of MRN, controls MRE11 endonuclease activity which is stimulated when CtIP is phosphorylated by CDKs and ATM/ATR (see Section 2b).⁹⁵ This triggers the formation of a nick which is the initiation site for bidirectional resection performed in the 3'-5' direction by the exonuclease activity of MRE11 and in the 5'-3' direction by the exonuclease activity of EXO1 and/or DNA2 in complex with the Bloom syndrome helicase (BLM). Following end resection, the ssDNA is stabilised by RPA, which comprises the heterotrimer RPA1/2/3 known to rapidly bind to and stabilise ssDNA via its oligonucleotide/oligosaccharide-binding fold (OB-fold) domains during DNA replication and repair⁹⁶ (Fig. 3). RPA is replaced by Rad51 (homologue of bacterial recombination protein A, recA), responsible for scanning the sister chromatid for the homologous strand, aided by BRCA2 and BRCA1 in association with BRCA1-associated RING domain 1 (BARD1).^{97,98} Homologous strand invasion requires stabilisation of the non-template sister strand by Rad51, whilst DNA Pol δ fills in the invading strand using the template strand (Fig. 3). HR occurs during and after synthesis

(S) phase, when two sister chromatids are present, and enables faithful DNA replication (Section 2b).

2a3. Interstrand cross-links (ICLs) repair. ICLs are covalent linkages between two different DNA strands, as opposed to intrastrand cross-links that occur on the same strand and which are primarily repaired by NER. ICLs are detected and repaired during DNA replication by the Fanconi Anemia (FA) pathway, named after the disease resulting from genetic mutations of key components of this pathway. The FA pathway requires activation by ATR kinase, a master mediator of DDR signal transduction (see Section 2b), which simultaneously reduces replication fork speed.⁹⁹⁻¹⁰¹ Currently, 22 genes have been described as FA genes, some of them being implicated in other DNA repair mechanisms: FANCA, -B, -C, -D1 (BRCA2), -D2, -E, -F, -G (XRCC9), -I, -J (BRIP1), -L (PHF9), -M, -N (PALB2), -O (RAD51C), -P (SLX4), -Q (ERCC4/XPF), -R (RAD51), -S (BRCA1), -T (UBE2T), -U (XRCC2), -V (REV7/MAD2L2) and -W (RFWD3).¹⁰² The FANCA, -D, -C, -E, -F, -G, -L and -M proteins form the core complex which monoubiquitinates FANCD2/FANCI upon ICL recognition, while the other proteins mediate the lesion incision or “unhooking” forming a DSB whose repair involves components of translesion synthesis (TLS, DNA polymerase κ) and HR pathways. The FA complex recognises the ICL, often at the collision site of one or two stalled replication forks, leading to incision by endonucleases (the structure-specific endonucleases SLX1, MUS81 (also known as SLX3), XPF and ERCC1) in one strand on either side of the ICL thus “unhooking” the ICL lesion and forming a DSB. Replicative



polymerases act on the intact strand to fill the template whilst retaining the incised strand in proximity. Finally the HR machinery repairs the DSB in the incised strand from the template double strand.¹⁰³ ICLs can also be repaired by a replication-independent mechanism involving incisions on the ICL flanks by the NER machinery and lesion bypass by Polk.^{99,104,105}

2b. The DDR mediators. The key mediators in the repair of DNA lesions resulting from SSB, DSB and arrested transcription and replication forks are the poly(ADP-ribose) polymerases (PARPs) and the phosphatidylinositol 3-kinase-related kinases (PIKKs, *i.e.* DNA-PK, ATM and ATR, Fig. 3).

PARP1 and PARP2 (the latter accounting for 5–10% of total PARP activity)¹⁰⁶ are considered a major first line of defence in the DDR response.^{4,33,37} PARP activation, PARylation (synthesis of branched and linear chains of poly(ADP-ribose) (PAR) on proteins) and auto-PARYlation (Fig. 4a) occurs rapidly at sites of SSB, DSB and stalled replication forks.^{80,107} PAR attachments on PARP and on nearby histones act as a platform promoting the recruitment of DDR factors including XRCC1/Lig3 and inducing chromatin remodelling.¹⁰⁸ Importantly, PARP1 is also activated by DSBs where it promotes the recruitment of NHEJ and HR factors.^{33,109} Repair of SSB can either be processed in a short- or long-patch DNA gap filling mechanism similar to that of BER,¹⁰⁷ mediated by nucleases such as FEN1 and DNA Polδ, the lagging strand polymerase capable of strand displacement.⁸⁷

The role of PARP3 in DNA repair is more poorly understood, although has been shown to have more protein PARylation targets than PARP1 and 2 with minor overlap.¹¹⁰ PARP3 has a different DNA-binding domain, it is described to recognise DSB only¹¹¹ and it has been shown to promote the recruitment of the NHEJ ligation machinery, the XRCC4/Lig4 complex.¹¹² Of interest, PARP3 was found to promote chromosome rearrangements and its genetic knockout in lung adenocarcinoma cells showed extreme sensitivity to small molecules which induce DSB¹¹³ or stabilise the secondary DNA structure G-quadruplexes (G4).¹¹⁴

The PIKKs coordinate the central alarm system to DSB and stalled forks, often during NHEJ/HR (see Section 2a2), through activation by (auto-)phosphorylation and recruitment of a vast range of signalling molecules.^{33,115} These signal transducers have significant structure and function similarity, however are activated by distinct sensor proteins which recognise the DNA damage. PIKKs comprise two families of effectors, DNA dependent protein kinase (DNA-PK) on one hand, and ataxiatelangiectasia mutated (ATM) and ataxiatelangiectasia and Rad3-related (ATR) on the other hand, which are recruited to DNA damage sites through molecular interactions with analogous C-terminal motifs of Ku80, Nibrin (or Nijmegen) breakage syndrome gene (NBS1) and ATR-interacting protein (ATRIP) respectively (Fig. 3).^{37,116}

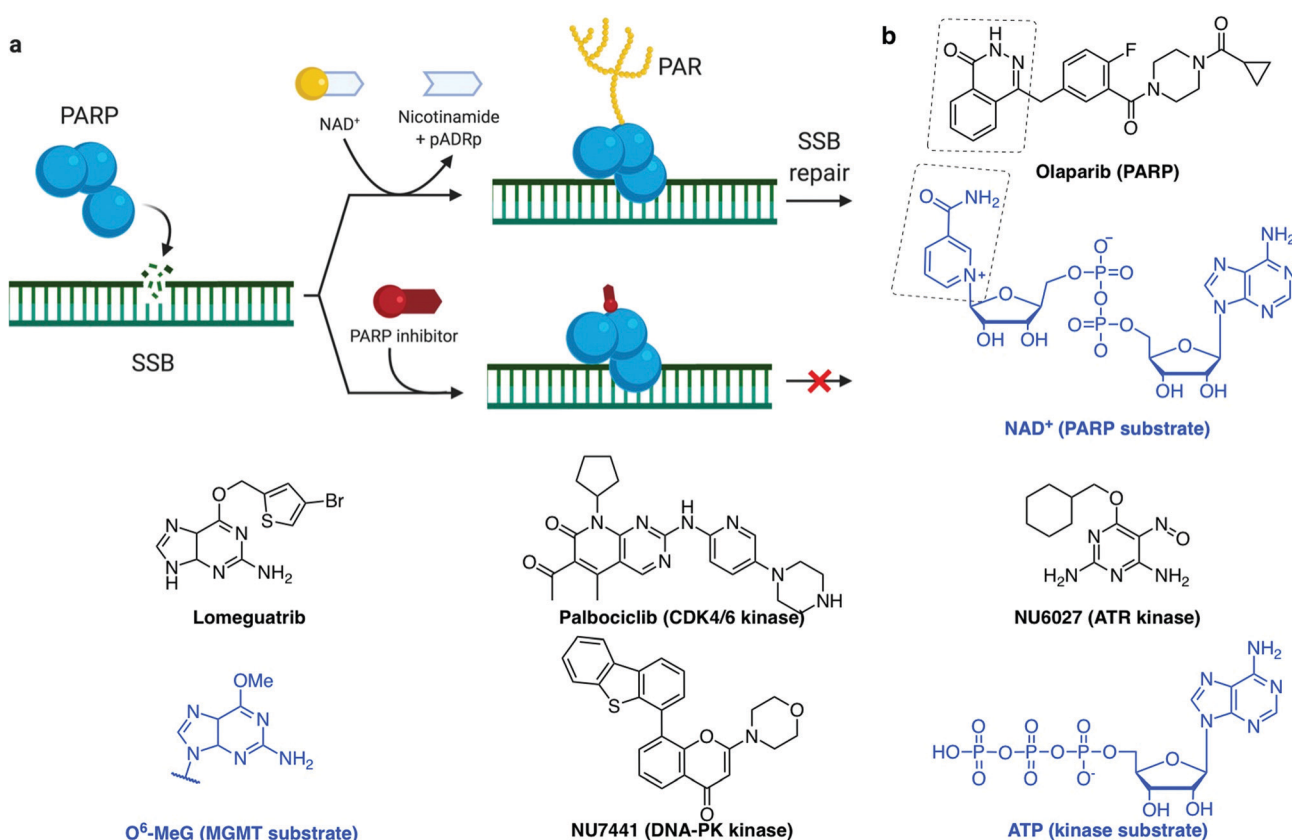


Fig. 4 (a) PARP recruitment to SSB and synthesis of poly(ADP-ribose) (PAR) to activate SSB repair. Inhibition of repair by PARP-trapping at the break site. (b) Examples of inhibitors (with their protein target) and the substrate they mimic (blue). (a) Created with BioRender.



– DNA-PKcs (also known as DNA dependent protein kinase catalytic subunit) is involved in DSB repair by NHEJ. The Ku heterodimer comprising Ku70/Ku80 (also known as X-ray repair cross-complementing protein 6 and 5, XRCC6 and XRCC5, respectively) binds to free DSB ends (amongst other DNA structures) and recruits DNA-PKcs. The Ku-DNA-PKcs-DNA complex forms the active DNA-PK holoenzyme which is a serine/threonine kinase able to phosphorylate several substrates including itself. Autophosphorylation of DNA-PKcs regulates DNA end processing.

– ATM, through a specific amplification loop, plays an apical role in DSB signalling and cell cycle checkpoint activation, in addition to its function in promoting DSB repair *via* HR.^{6,71} The MRN complex associates to the flank of each DSB through the ATP-dependent RAD50 clamp which recognises the DNA damage. The MRN complex mediates the recruitment of ATM by direct interaction between ATM and the NBS1 C-terminus.¹¹⁷ However, in contrast to Ku whose recruitment is limited to one or two Ku proteins per DSB end,¹¹⁸ multiple MRN-ATM complexes are recruited to chromatin thanks to an amplification loop relying on ATM kinase activity. Indeed, ATM is responsible for the phosphorylation of a large number of proteins including the C-terminus of the histone variant H2AX, generating γH2AX, the S139-phosphorylated form of H2AX. This initial phosphorylation recruits MDC1 (mediator of DNA damage checkpoint protein 1), *via* its BRCA1 C terminus (BRCT) repeat domains,^{119–122} which is itself phosphorylated by casein kinase 2 (CK2). The phosphorylated form of MDC1 is recognised by the forkhead-associated (FHA) and BRCT domains of NBS1 which drives the secondary recruitment of the MRN-ATM complex that subsequently phosphorylates other more distant substrates including H2AX.^{123–126} This amplification loop leads the γH2AX signal to spread from kilo- to megabases from the DSB, making the detection of the γH2AX signal a sensitive and widespread approach to monitor the induction of DNA damage in cells (examples in Section 3c). The rapid spreading of γH2AX from DSB sites is controlled by chromatin domain boundaries delimited by DNA-binding cohesin protein complexes,^{127,128} which also repress transcription of damaged genes.¹²⁹ In addition to NBS1, MDC1 recruits other proteins that govern cell cycle checkpoints, such as checkpoint kinase 2 (CHK2), and mechanisms of DNA repair pathway choice including 53BP1 (promoting NHEJ) and BRCA1 (promoting HR).^{130,131} As well as recruiting DNA repair proteins (MRN/ATM), γH2AX recruits chromatin remodelling machinery to aid damage repair.^{122,132} It is noteworthy that ATM can also be directly activated by oxidative stress *via* oxidation of some of its cysteines.⁷¹ In parallel to its signaling functions, ATM also stimulates HR repair (Fig. 3) by phosphorylating several proteins including the MRN interacting protein CtIP. CtIP phosphorylations promotes MRE11 endonuclease activity, responsible for nicking the side(s) of the DSB to initiate end resection. The ATM-dependent CtIP phosphorylations promote Ku eviction from the DSB end, thereby directing DSB repair towards HR.¹³³ In addition, ATM stimulates HR by phosphorylating several HR factors, such as BRCA2, EXO1 and BLM, and NHEJ factors such

as XRCC4, XRCC4-like factor (XLF) and DNA-PKcs.^{134–137} ATM also plays a crucial role in enforcing the cell cycle checkpoints through activation by phosphorylation of CHK2, which in turn activates p53, a key apoptosis regulator.⁶⁸ 53BP1 is recruited to nucleosomes with specific histone modifications: Ring Finger Protein 168 (RNF168)-ubiquitylated H2AK15 and dimethylated H4K20;¹³⁸ where it functions as a central determinant in the repair pathway choice made at DSB by promoting the recruitment of the shieldin complex,¹³⁹ thereby limiting DNA end resection by BRCA1/BARD1, which exposes ssDNA to be repaired by HR. As such 53BP1 inhibits HR and supports NHEJ repair in the G1-phase of the cell cycle.^{6,140} PARP3, in interaction with Ku, similarly is described to favour DSB repair to NHEJ by limiting end resection.¹¹³

– ATR is the key damage signalling mediator at replication forks and in dividing cells. ATR binds through ATRIP to the RPA:ssDNA complex,^{141,142} which forms rapidly at DNA replication fork blockages during the S phase of the cell cycle (Fig. 2 and 3) and stabilises ssDNA during DNA replication. Upon DNA damage, ATR induces a signalling cascade, phosphorylating a long list of cell cycle-related substrates, notably checkpoint kinase 1 (CHK1) and apoptosis and replication stress regulator p53.^{4,143} ATR activation stabilises the stalled fork (preventing fork collapse which can otherwise lead to DSB formation)¹⁴⁴ and prevents proximal replication origin restart.^{142,145,146} Stalled replication forks can be restarted by an HR-dependent mechanism. Unlike ATM and DNA-PKcs, ATR is necessary for cell growth and embryonic development,^{147,148} meaning mechanistic studies of ATR were greatly facilitated by the recent development of ATR-inhibiting chemicals. Cross-talk exists between the ATM and ATR kinases since ATR can activate ATM in some instances,¹⁴⁹ the two kinases having many substrates in common and their cellular roles strongly overlapping. However, most of the details regarding the intricate genetic regulations in which ATM and/or ATR remain to be discovered.

2c. DNA damage checkpoints. Healthy cell cycle progression is driven by the cyclin-dependent kinases (CDKs) and CDK-cyclin complexes¹⁵⁰ and exit is controlled by p53 and p21 signalling.^{68,151} In addition to regulation by cell mass and proliferative signals, cell cycle progression can be interrupted upon DNA damage at specific cell cycle boundaries, called the cell cycle checkpoints.¹⁵² Three DNA damage checkpoints exist: the G1/S, G2/M and intra-S phase checkpoints (Fig. 2). The G1/S checkpoint blocks S-phase entry of cells carrying DNA damage, allowing repair and preventing the conversion of simple DNA lesions into more complex ones during the replication process. It is enforced by activation of the ATM-CHK2-p53 axis, which results in the accumulation of the CDK-cyclin inhibitor p21.¹⁵³ As a result of its dependence on p53, this checkpoint is frequently lost in cancer cells, providing an explanation for the high level of genomic instability in cancer cells. The intra-S checkpoint relies on both ATM and ATR-CHK1¹⁵⁴ and consists of blocking the initiation of new replication origins once DNA damage is detected. Activation of the intra-S phase checkpoint delays the progression through S-phase although it does not



inhibit it completely. Finally, the G2/M checkpoint relies on the activation of ATM-CHK2 and ATR-CHK1 and prevents damaged cells from initiating mitosis.¹⁵⁵ Progression through mitosis could result in generating two daughter cells with incomplete or abnormal genomes or in failure to partition DNA leading to mitotic catastrophe, a type of cell death that is frequently observed when treating cancer cells with DNA damaging agents. Key enforcers of the DNA damage checkpoints are the CHK2 and CHK1 serine/threonine kinases that are activated by ATM and ATR respectively, and can phosphorylate various substrates,¹⁴¹ including the cell division cycle 25 (CDC25) phosphatases A, B and C. CDC25 dual-specificity phosphatases normally remove inhibitory phosphorylations of the CDKs,^{156,157} a rate-limiting step in CDK activation which allows cell cycle progression.¹⁵⁸ Once phosphorylated by CHK1/CHK2, CDC25 phosphatases are degraded, inactivated or sequestered into the cytoplasm by association with the 14-3-3 proteins, thereby blocking cell cycle progression. Considering that cancer cells display high genomic instability, due to increased levels of replication, mitosis and metabolism and/or dysregulation of the DDR machinery, it has been envisioned that they should be more dependent on DNA damage checkpoints than normal cells. This idea led to a race to develop DNA damage checkpoint inhibitors, which could induce selective cell death of cancer cells as a standalone treatment, or in combination could increase the selectivity and activity of current DNA damaging anticancer treatments (see Sections 1 and 2d).^{4,159}

2d. Small molecule targeting of DDR. Impairments in the mechanisms responsible for the surveillance and repair of genetic material throughout cell division are unsurprisingly causative of oncogenetic dysregulations. Because cancer cells are often DDR-deficient, current strategies aim at fostering the genetic instability of cancers by further deactivating these surveillance and repair systems.^{5,8,160} There are in fact several rationales justifying the use of DDR-disruptive agents in cancer treatment. Small molecule inhibitors of DNA repair proteins are able to increase the level of DNA damage in cancer cells (see below), whereas inhibitors of checkpoint regulators, such as WEE1,¹⁵⁵ a crucial regulator of the G2/M checkpoint which stops cells entering into mitosis on activation by CHK1, function by metaphorically speeding up the cell cycle to an uncontrollable speed, causing it to 'derail' under the pressure of unsustainable genetic aberrations.¹⁶¹ DDR inhibition can be used in combination with a DNA damaging agent, or crucially can be used in cancers which are already deficient in a DDR pathway, a strategy known as synthetic lethality.

Synthetic lethality describes the relationship between two gene deactivations which cause cell lethality when both genes are silenced (through endogenous mutation, exogenous inhibition or knockdown) but cells are viable when only one is deactivated.^{7,162–165} As discussed above, DDR pathways function with significant redundancy, yet with one pathway missing, cancer cells become more dependent on the remaining pathways. A myriad of clinical trials are in place to assess the benefit of inhibiting multiple DDR pathways concomitantly, with the hope of being able to widen the therapeutic window by

increasing the combined tolerated dose and decreasing the effective dose.⁸ The first-in-class example of a synthetic lethality strategy applied to cancer is the PARP inhibitor olaparib (Fig. 4b),^{8,108} which was approved for treatment of several BRCA-deficient cancer types by the US Food and Drug Administration (FDA) and European Medicines Agency (EMA) in 2014. More recently (in 2018), another PARP inhibitor talazoparib was approved as it displays a higher cytotoxicity in BRCA-deficient tumours, likely due to its stronger PARP–DNA trapping affinities (Fig. 4a).¹⁶⁶ PARP inhibitors have proved efficacious in multiple cancers with deficient DDR pathways, as well as in maintenance therapy for cancers which have previously responded to antiproliferative agents.^{5,160} However, various resistance mechanisms to PARP inhibitors have been described, including *via* the restoration of HR, for example *via* inactivation of 53BP1, or by increased replication fork stability, for example *via* inactivation of the junction endonuclease MUS81 or the *N*-methyltransferase enhancer of zeste homolog 2 (EZH2).^{109,166,167}

Other DDR inhibitors (see Section 2c) have shown potential in the clinic, including inhibitors of specific DNA repair pathways such as DNA-PK inhibitors (M3814 and CC-115, currently in phase I trials^{168,169}) which block NHEJ-dependent DSB repair, and MGMT inhibitors (lomeguatrib, Fig. 4b) which block the removal of alkylated bases. Also in clinical testing are the inhibitors of cell cycle checkpoint mediators ATR, ATM, WEE1 and CHK1, such as M6620, M3541, AZD1775, LY2606368 respectively, for both single-agent and combination therapies.^{8,170} Despite promising, the therapeutic indexes of these molecules (toxic dose/therapeutic dose) are often limiting, as demonstrated in the cases of lomeguatrib¹⁶⁰ and UCN-01 (first specific CHK1 inhibitor developed from the broad-spectrum kinase inhibitor staurosporine^{155,171}). PARP inhibitors had initially been trialled in combination with DNA damaging agent temozolomide and failed due to excessive toxicity, but following these studies, its synthetic lethality properties in BRCA-deficient cancers were uncovered.¹⁶⁰ This exemplifies the advantage of synthetic lethality strategies, in which a single agent can be applied without incurring unacceptable collateral damage, by taking advantage of a known weakness in the patient's cancer. With the aid of modern high-throughput and genome-wide sequencing techniques,¹⁷² the body of knowledge on genetic biomarkers is being established for each cancer type, opening up the development of specific small molecules suited to personalised cancer therapy.^{4,162} Thus, our understanding of the cancer genotypes that are likely to be sensitive to selected treatment is growing.

A common trend in the design of small molecule inhibitors is to structurally mimic the biological substrate of the target protein: *O*⁶-benzylguanine and lomeguatrib (*O*⁶-(4-bromophenyl)guanine) both mimic *O*⁶-methylated guanine (*O*⁶-MeG), the DNA base lesion repaired by MGMT (*O*⁶-methylguanine–DNA methyltransferase, Fig. 4b); olaparib mimics NAD⁺, the building block used by PARP for PARylation signalling (Fig. 4a); and palbociclib fits into the ATP-pocket of checkpoint kinase CDK4/6¹⁷³ (Fig. 4b), whereas other kinase inhibitors can mimic the target protein. Some DDR inhibitors, such as NU6027¹⁷⁴ and UCN-01¹⁷⁵ (ATR and CHK1 inhibitors respectively, Fig. 4b), were initially designed to target CDKs, and the



marginal specificity for DDR checkpoint kinases over another kinase was developed by often lengthy medicinal chemistry methods: trial-and-error synthesis, structure–activity relationship (SAR) studies and enzyme assays, since many active site crystal structures of these targets have not yet been resolved.

In the next chapter, we discuss another approach, which could bypass certain toxic and resistance-incurring pitfalls of DDR, or prove synergistic. Inspired by a strategy at the very foundation of chemotherapy, we look again to triggering the DDR by drugs targeting the DNA itself. Accumulating evidence now supports the existence of non-B DNA structures in cells that fold as a result of cellular activity. The resulting higher-order structures are closer to protein than B DNA in terms of 3D structure and offer more structurally defined binding sites for small molecules (e.g. the accessible G-quartets of G-quadruplexes, the central cavity of DNA junctions, see Sections 3a and 4a) than major/minor groove interaction and/or intercalation in between two successive base pairs of B DNA. This offers the possibility of targeting higher-order nucleic acid structures with a better degree of selectivity, thus re-establishing DNA in its many forms as a promising chemotherapeutic target with unique and structurally defined small molecule binding sites. In the following sections, we will describe the new trend that is emerging based on the stabilisation of non-B DNA structures with specifically designed small molecules (ligands), as a way to damage cancer cells' DNA in a more specific manner.

3. G-Quadruplexes (G4s) as targets to induce DNA damage

3a. The prevalence of G4 in the human genome. The non-B DNA structure which has been most studied for triggering DNA damage is undoubtedly the four-stranded structure named G-quadruplex-DNA, or G4-DNA. If a transiently formed

guanine-rich ssDNA displays the motif $G_{\geq 3}N_xG_{\geq 3}N_xG_{\geq 3}N_xG_{\geq 3}$ (where G is guanine, and N any intervening nucleobase, and x ranges from 1 to > 20), the sequence can give rise to a G4 (Fig. 5).^{179,180} The basic building block of G4s is a G-quartet¹⁸¹ resulting from the self-association of 4 Gs in a square planar arrangement, and was first reported *in vitro* in 1962.¹⁸² Contiguous G-stretches within a strand of ssDNA come together into contiguous planar G-quartets, which stack on top of each other through π -system interactions, with each G-quartet being stabilised by a central physiological cation (K^+ , Na^+) to form a G4 structure that can be classified as antiparallel-, hybrid- or parallel-type G4, depending on the polarity of the strands (Fig. 5). Of note, this topological diversity leads to a variety of intervening loops, which can be diagonal, lateral or reversal loops (Fig. 5). Initially only thought of as an *in vitro* oddity, G4s are now considered key players in cellular processes: recent sequencing-based methods have demonstrated that thousands of quadruplex-forming sequences (QFS) are present in our genome, $> 700\,000$ by G4-seq.¹⁸³ These sequences are predominantly maintained in an unfolded state, as exemplified by the detection in live cells of only $> 10\,000$ G4s by G4 ChIP-seq¹⁸⁴ and *ca.* 3000 G4s by the fluorophore SiR-PyPDS (see Section 3c),¹⁸⁵ with a strong correlation between individual G4 formation and the transcriptional activity of the gene they fold from. This transiency originates from the various mechanisms the cell has evolved to regulate G4 formation, among which the G4 helicases are being actively studied (see Section 3b). Interestingly, the QFS distribution is not random as they are significantly enriched in key regulatory regions including gene promoters, replication origins and telomeres.^{10,186}

G4s in gene promoters: the occurrence of QFS in gene promoters (defined as 1 kb upstream of the transcription start

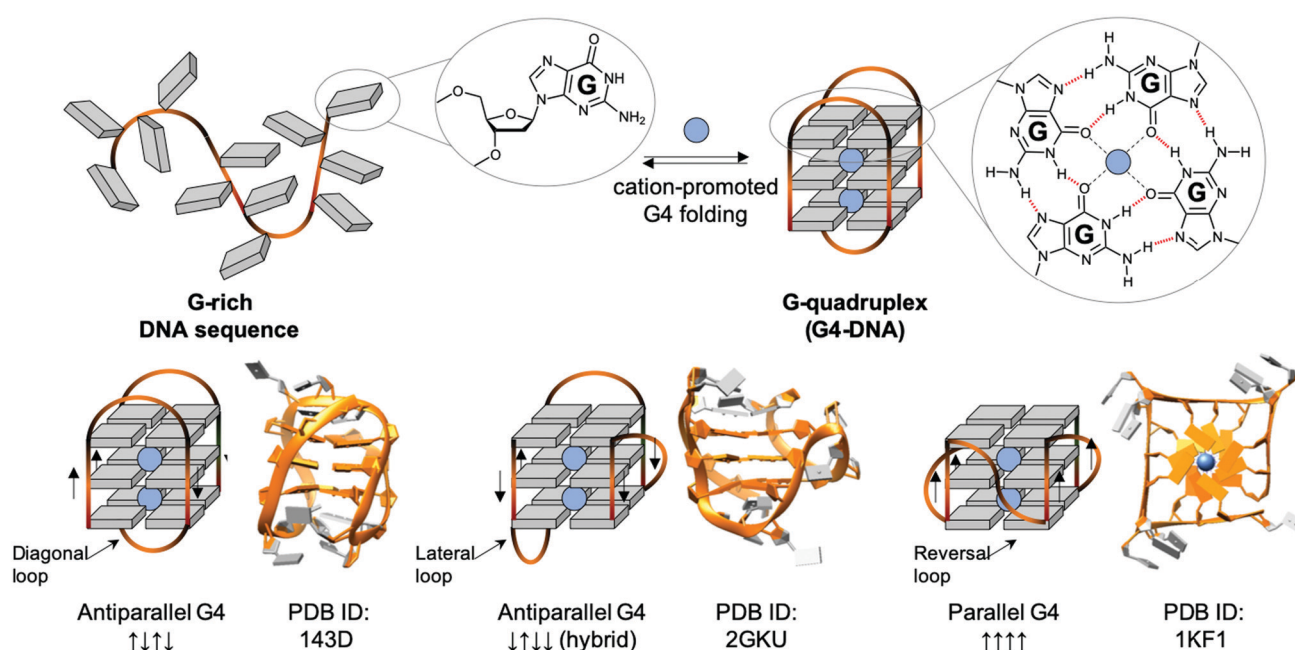


Fig. 5 Schematic representation of a G-rich sequence that folds into a G4 structure (upper panel), highlighting the structure of a guanine (G, upper panel, left) and a G-quartet (right). Topological diversity of G4s that can adopt parallel, hybrid and antiparallel conformation (lower panel, arrows indicate the polarity of the DNA strands), as elucidated by either NMR (PDB IDs 143D¹⁷⁶ and 2GKU¹⁷⁷) or X-ray structure analysis (PDB ID 1KF1¹⁷⁸).



site, TSS) is significantly high in mammals (>2000),^{187–189} inferring a regulatory role of G4s in gene expression.¹⁹⁰ The genesis of G4 structures is linked to a high transcriptional activity, likely due to physical requirements for the DNA duplex to open and for chromatin structure to be relaxed (euchromatin, as opposed to tightly-bound heterochromatin) thus allowing G4s to fold, but this link also indicates a possible causative role of G4s in the recruitment of transcription factors to euchromatin.¹⁸⁴ The observation that G4 sequences are significantly enriched in oncogenes and regions predisposed to amplification in cancers explains and warrants the active search for chemicals (small molecule ligands) that can interact specifically with promoter G4s,¹⁹¹ in order to gain control of processes underlying cancer onset and progression. The textbook example is c-MYC, a transcription factor and proto-oncogene whose protein was considered undruggable owing to its lack of catalytic activity, but for which gene expression can be down-regulated by ligand-mediated stabilisation of its G4-containing promoter region.⁴³

G4s in replication origins: in homeostatic conditions, DNA replication origins are regulated by two triggers, transcription and G4 formation.¹⁹² Genome mapping showed a correlation between replication origins and G4s in metazoans: $>70\%$ of initiation sequences are followed by a QFS, also known as Origin G-rich Repeated Element (ORE).¹⁹³ Furthermore, a genome-wide CRISPR-mediated gene editing study showed that by inserting or deleting QFSs, replication origin activity could be increased or decreased respectively,¹⁹⁴ demonstrating one of the multiple regulatory roles of quadruplexes in healthy dividing cells. However, the detailed mechanism by which G4s act at replication origins is not yet fully understood.

G4s in telomeres: mammalian telomeres are formed by thousands of non-coding repeats ($5'$ TTAGGG $3'$ / $5'$ CCCTAA $3'$) and a single-stranded G-rich 3'-tail, known as the 3' overhang.^{195,196} The telomeric G4 that folds from the G-rich ($5'$ TTAGGG $3'$) 3'-overhang is undoubtedly the most intensively studied G4 structure, given that it was the first discovered, and that its 3 TTA loops are flexible enough to give rise to different G4 topologies, as exemplified by the 3 structures seen in Fig. 5 (lower panel), formed in either Na^+ - (PDB ID 143D¹⁷⁶) or K^+ -rich conditions (PDB IDs 2GKU¹⁷⁷ and 1KF1¹⁷⁸). Telomeres act as a cap at the end of chromosomes, protecting them from enzymatic degradation. Repeated units are removed after each round of DNA replication and cell division (the so-called 'end replication problem'),^{197,198} making telomeric regions the 'mitotic clock' of the cell, limiting the number of cell cycle divisions before the onset of senescence.¹⁹⁶ Telomeres are thus implicated in both cancer and the ageing process.^{199,200} It is largely accepted that G4s form in telomere ends,^{201,202} and the earliest evidence of biological G4 formation was obtained at telomeres. Even if telomeric G4s can either protect telomeres against exonucleases²⁰³ or jeopardise telomere organisation by preventing telomeric loop (T-loop) formation and telomerase recognition,^{204–207} they have been mostly studied as targets for fostering chromosomal fragility in cancers.²⁰⁸

3b. G4 unwinding enzymes – G4 helicases. Given the widespread distribution of QFSs in our genome, the formation of G4s

represents a constant impediment to the advancing transcription/replication complexes along the DNA. Cells have evolved an enzymatic machinery to tackle this threat, the helicases, some of which have now been shown to interact with and resolve G4s.^{209,210} Helicases are increasingly being studied for their role in diseases,^{211,212} notably because their genetic silencing is involved in severe dysregulations that confer premature ageing and cancer susceptibility. To date, 95 different helicases are known, 31 for DNA and 64 for RNA.²¹¹ The two most studied families are (1) the RecQ helicases, including Bloom (BLM) and Werner (WRN) helicases, along with RecQ-like helicases 1, 4 and 5 (RECQL1, RECQL4 and RECQL5, respectively), and (2) the iron–sulfur (Fe–S) helicases, including XPD, Fanconi anaemia complementation group J (FANCIJ, or BRIP1), regulator of telomere length helicase 1 (RTEL1) and DEAD/H-Box helicase 11 (DDX11).^{211,213}

These enzymes are deeply involved in genome surveillance and DNA damage repair (DDR). They are known to resolve higher-order DNA structures formed during DNA transactions (e.g. the 5' flap intermediate resulting from the processing of Okazaki's fragments during replication, or the DNA:RNA hybrids referred to as R-loops formed during transcription) and structures formed during DDR (e.g. the four-way junction known as the Holliday junction that is the central intermediate of HR). Some examples have already been discussed above: BLM is involved in HR-mediated DSB repair (see Section 2b); XPD mediates NER when embedded in the ternary complex TFIIH (see Section 2a1). Of particular interest here is the prevalence of reports that helicases are inhibited by G4 ligands, likely indirectly through DNA blockages (Table 1).

More recently, certain helicases have been shown to unwind G4s *in vitro* and evidence is now accumulating that they do so *in vivo* (Table 1).^{212,214} For instance, BLM localises at telomeres where it allows for proper replication to occur, resolving telomeric G4, an action that is retarded by G4 ligands (PhenDC3).²⁴⁰ Similarly, BLM-deficiency leads to genomic instability caused by elevated levels of recombination by sister chromatid exchange (SCE) events at G4 sites in transcribed regions of the genome.²⁴¹ WRN is also found at telomeres where it resolves telomeric G4 to repress chromosomal instability, and interact with the shelterin complex to regulate T-loop formation.²⁴² A pioneering study conducted with Dog-1, the *Caenorhabditis elegans* homologue of FANCIJ, showed that deletions in Dog-1 helicase promotes genetic instability by allowing G4s to act as replication barriers.²⁴³ In eukaryotic cells, FANCIJ-knockdown (FANCIJ^{KD}) cell growth is strongly inhibited upon treatment with G4 ligand telomestatin (TMS).²³⁵ Similarly, FANCIJ-null cells show replication fork slowing upon TMS treatment.²⁴⁴ Genome-wide analysis performed with *S. cerevisiae* by Pif1-ChIP-qPCR showed an enrichment of G4 motifs at Pif1-binding sites, which are responsible for stalling replication forks and are prone to DNA breakage in Pif1-deficient cells.²⁴⁵ This was further substantiated by the demonstration that Pif1-deficient *S. cerevisiae* displayed enhanced genetic instability triggered by G4 motifs of different (and controllable) stability, and chemical G4 stabilisation *via* G4 ligand treatment (PhenDC3).²⁴⁶

Beyond their biological roles, G4-helicases have been implemented as biomolecular tools to study and/or modulate G4



Table 1 G4-helicases^{209,212,214,215}

Name	Family ^{216,217}	Direct- ionality Species	Activity (for ligand interaction details, see Section 3c)	Associated human disease	Ref.
BLM	RecQ-like SF2	3' → 5' <i>H. sapiens</i>	No preferences for a given subclass of G4 (intra- vs. inter-molecular G4, parallel vs. antiparallel G4s); requires 3'-ss overhang for loading; involved in DNA repair via DSB resection and HR; inhibited by G4-ligands (e.g. BRACO19)	Bloom syndrome	218–220
Sgs1	RecQ-like SF2	3' → 5' <i>S. cerevisiae</i>	Orthologue of <i>H. sapiens</i> BLM helicase; inhibited by G4-ligands PIPER	—	221 and 222
WRN	RecQ-like SF2	3' → 5' <i>H. sapiens</i>	Wide G4 spectrum; requires 3'-ss overhang for loading; manages replication stress, fork arrest and collapse; inhibited by G4-ligand BRACO19	Werner syndrome	219, 220 and 223
RecQ	RecQ-like SF2	3' → 5' <i>E. coli</i>	Unwinds both intra- and intermolecular G4s; orthologue in <i>H. sapiens</i> is RecQL4; inhibited by G4-ligand NMM	—	224
DHX36	DEAH, SF2	3' → 5' <i>H. sapiens</i>	Also known as RHAU (RNA helicase associated with AU-rich element) or G4R1 (G4-resolvase 1); unfolds both DNA and RNA G4s; preference for parallel G4s; inhibited by G4 ligands PDS and PhenDC	—	225–227
Pif1	Pif1-like SF1	5' → 3' <i>S. cerevisiae</i>	Found from yeast to human (hPif1); no preference for a given subclass of G4; requires 5'-ss overhang for loading; inhibited by G4-ligand PhenDC3	—	228–231
RTETL1	DEAH, SF2	5' → 3' <i>H. sapiens</i>	Involved in genome stability and telomere integrity; unwinds intra- and inter-molecular G4s; inhibited by G4 ligand TMPyP4	Hoyerlaa-Hreibardsson syndrome	232 and 233
FANCJ	Fe-S, SF2	5' → 3' <i>H. sapiens</i>	Orthologue of nematode DOG-1 helicase (<i>C. elegans</i>); promotes DNA repair via HR on interaction with BRCA1; resolves telomeric G4; inhibited by G4-ligand telomestatin (TMS)	Fanconi anemia	234–236
DDX1	Fe-S (DEAD- box), SF2	5' → 3' <i>H. sapiens</i>	Unlike FANCJ, DDX1 does not resolve intra- and tetra-molecular G4s (preference for two-stranded G4s); insensitive to TMS treatment	Warsaw breakage syndrome	236 and 237
XPD	Fe-S (Rad3/ XPD), SF2	5' → 3' <i>S. acidocaldarius</i>	Unlike FANCJ, XPD does not resolve intra- and tetra-molecular G4s; orthologue in <i>H. sapiens</i> is ERCC2; insensitive to TMS treatment	Cockayne syndrome (for ERCC2)	236, 238 and 239

landscapes in cells. For instance, Pif1 has been labelled (Pif1-GFP) to track G4 formation in cells through co-localisation with the clickable G4 ligand pyridostatin- α (PDS- α , visualised following *in situ* click chemistry, see Section 3c).⁹² Pif1 has also been overexpressed in cells to reverse G4-mediated genetic instability in *S. cerevisiae*,²⁴⁷ repair dysfunctions in cancer cells²⁴⁸ and cognitive dysregulations in neurons.²⁴⁹ In the latter two examples, Pif1 overexpression was found to remedy phenotypes of G4-mediated functional impairment induced by G4-stabilising PDS treatments. Similarly, BLM was overexpressed to reverse reactive oxygen species (ROS)-induced G4 (and R-loop) formation at transcriptionally active sites, thus restoring the repair process.²⁵⁰

3c. G4 stabilising ligands and chemical biology. As indicated above, the use of small molecule ligands (PhenDC3, TMS, PDS) to perturb cellular equilibria is often used to provide readouts amenable to mechanistic interpretations. This approach abides by the initial definition of chemical biology by Linus Pauling (1954)²⁵¹ and complies better with more modern interpretations of Karl-Heinz Altmann (chemicals are used “*to interrogate, modify, and manipulate biological systems at the cellular level in a highly controlled manner*”)²⁵² or of Stuart L. Schreiber (chemicals are used as “*reversible modulators*” of biological systems “*to gain a deep understanding of cell circuitry and diseases biology*”).²⁵³

Since the very first prototypes at the end of the 1990s,^{254–256} hundreds of G4 ligands¹³ have been developed to act as both reversible modulators of G4 cellular functions^{257,258} and as probes to visualise G4s intracellularly (thus confirming their *bona fide* existence).²⁵⁹ Selectivity for G4s over dsDNA was imperative in the development of G4 ligands. Guidelines for selective G4 ligand design include structural prerequisites of a

wide π -surface to enable optimal π - π stacking interaction with the accessible G-quartet of a G4, cationic appendages to interact with the negatively charged DNA backbones, and water-solubilising sidechains.^{260,261} Structural studies *in vitro* reveal that most G4 ligands stack above the top G-quartet of the G4.^{262–264} Chemical investigation and design optimisation are always accompanied by the development of *ad hoc* biophysical techniques to assess in an ever more precise manner the intrinsic quality (affinity, selectivity) of hundreds of candidates that have yet been evaluated. Such techniques include FRET-melting, fluorescence indicator displacement (FID) assays, ESI-MS and surface plasmon resonance (SPR) investigations.^{265,266} Rational chemical design is now also complemented by high-throughput screening methods (small molecule microarrays,²⁶⁷ and DNA microarrays²⁶⁸) that assess the interaction of hundreds of ligands with hundreds of G4 targets. Such methods offer great potential for the discovery of new G4-targeting chemical scaffolds, as well as a new methodology to compare G4 specificity, where the intra-G4-selectivity is currently a challenging barrier to application.

The ability of G4 ligands to alter cancer cell proliferation is now firmly established.^{14,257} However, the precise mechanisms by which they exert their anticancer activity must still be clarified through impartial experimentation and in multicellular conditions. In light of what has been described above, it is tempting, but not necessarily justified, to link the observed global antiproliferative activity of a given ligand to a change on the gene that contains the G4 sequence being targeted, and for which the ligand shows *in vitro* binding to the truncated oligonucleotide (*i.e.* oncogene expression modulation for promoter G4s; telomere stability for telomeric G4s).^{190,192,269}



And yet, G4 formation is intimately linked to all DNA transactions, favoured by the transient formation of ssDNA during DNA-replication and DNA-to-RNA transcription, when the B-helix is split apart and the single strands are subjected to negative supercoiling, in which helical tension is theorised to cause the DNA to curl like a twisted string. Whilst transactions induce their formation, G4s can physically impede these two processes, stopping or stalling the advancing polymerase, triggering DNA damage and activating DDR mechanisms. This way to consider G4 as a bulky steric hinderance creating DNA damage was – and still is – not well accepted in the field since it removes G4 from its position as a unique genetic lever able to control complicated cellular processes with exquisite spatio-temporal specificity (Fig. 6). However, this approach offers a model that accounts for most if not all results collected in cellular proliferation studies reported so far. Amongst the myriad of candidates synthesised to date, only a handful of ligands have been thoroughly characterised for their ability to trigger DNA damage, and for which the therapeutic potential through DDR modulation has been exploited.

Telomestatin (TMS): one of the first G4 ligands to be studied in cells was the *Streptomyces anulatus*-derived natural product TMS, isolated from wide range screening of telomerase inhibitors (chemical structures in Fig. 7a).²⁷⁰ A structural analysis (NMR) of a TMS/G4 complex confirmed that TMS interacts with G4 mostly *via* π -stacking atop the accessible G-quartet of the telomeric G4.²⁶² When treated with TMS, leukaemia cells proliferate for the first 20 days and apoptosis occurs after this lag (over ten more days), accounting for the delay of telomerase-overexpressing

immortal cancer cells to suffer TMS-induced telomerase inhibition and remortalisation (progressive shortening of the telomere leading to chromatin damage and apoptosis, involving activation of p53 and CHK2).²⁷¹ TMS exerts its activity beyond telomerase inhibition, either disrupting telomere organisation (uncapping telomere-associated proteins),²⁷² or inducing DNA damage (quantified by immunodetection of γ H2AX) in both telomeric (telomere dysfunction-induced foci, or TIFs) and non-telomeric regions, which triggers cell cycle arrest in G1 phase and apoptosis.^{273,274} TMS was also found able to inhibit the expression of genes containing QFS in their promoters (e.g. c-Myb in glioma cells),²⁷⁵ presumably by blocking polymerase processivity. The only sour note of TMS is its complicated accessibility; to tackle this issue, TMS derivative 6OTD was designed (Fig. 7a), as it is more readily synthesised than TMS.^{270,276} 6OTD equally shows interesting G4-interactions *in vitro*²⁷⁷ and inhibits the proliferation of glioma stem cells *via* both telomeric and non-telomeric DNA damage, G1-cell cycle arrest and apoptosis.

RHPS4: the pentacyclic acridinium salt RHPS4, whose NMR structure in complex with a G4 has been established,²⁷⁸ has been instrumental in the discovery of G4 ligand-mediated DNA damage. Six human melanoma lines treated with RHPS4 showed an increase in telomere dysfunction and cell cycle inhibition,²⁷⁹ and telomeric DNA damage (TIFs),²⁸⁰ demonstrated by foci of γ H2AX, 53BP1 and RAD17 colocalised with shelterin component telomeric repeat factor 1 (TRF1). Analysis of ATR^{KD} and ATM^{KD} cells and with the use of the ATR inhibitor caffeine revealed that RHPS4-induced DDR signalling is dependent on ATR, but not ATM, despite both being activated in WT

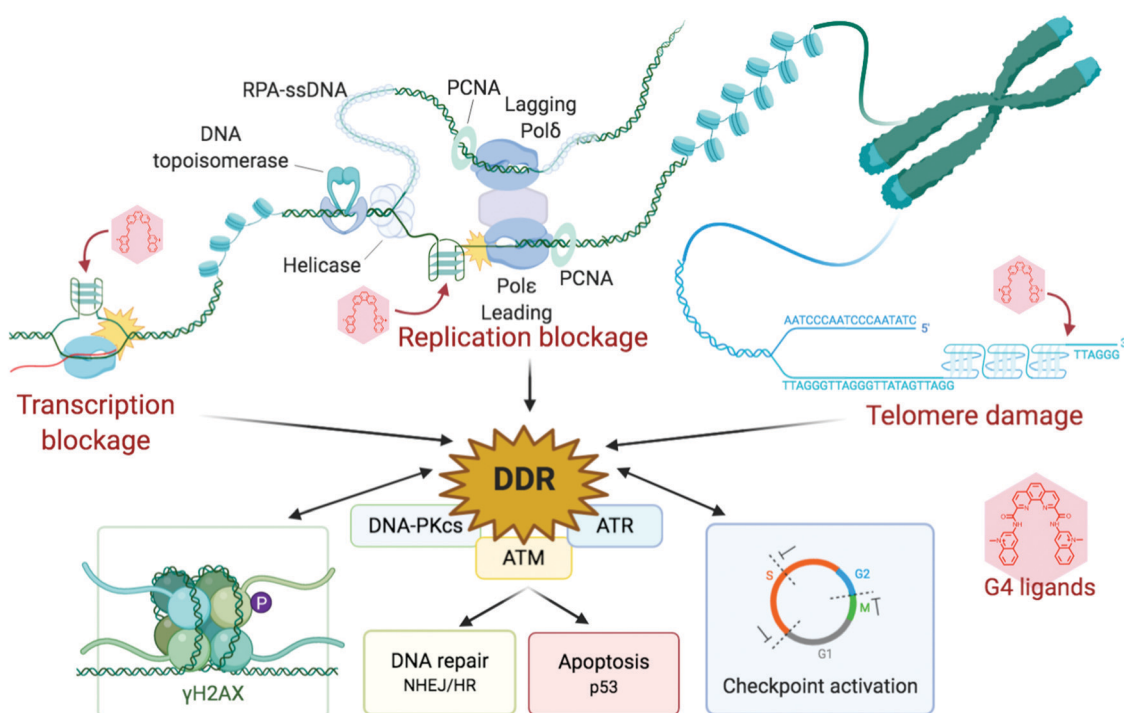


Fig. 6 Schematic representation of the stabilisation of G4s, inducing replication and transcription blockages and telomere damage, producing a DDR response, which feeds back into damage signalling (γ H2AX) and checkpoint inhibition, and eventually leads to recombination repair and/or cellular shutdown through apoptosis. Created with BioRender.



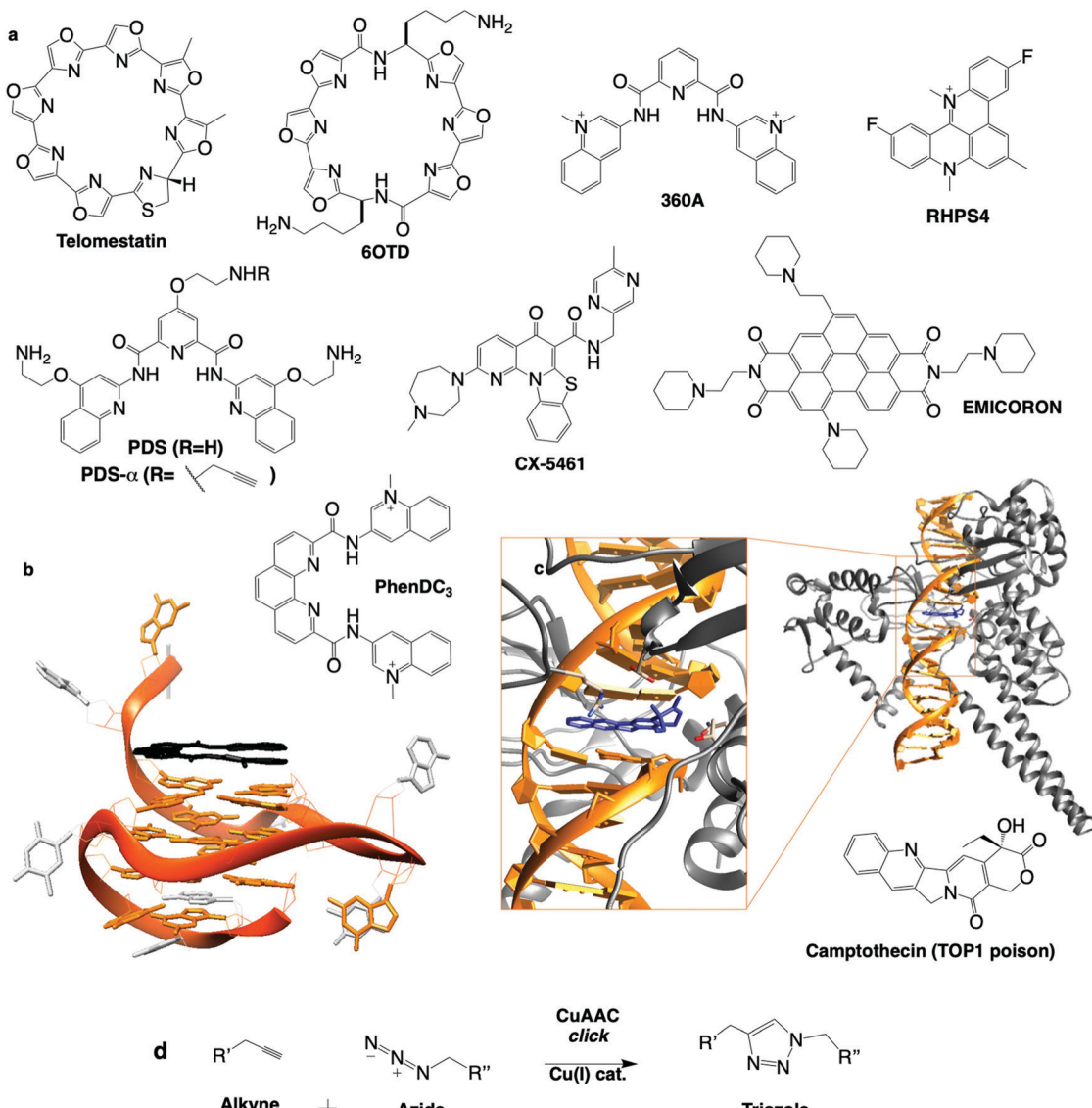


Fig. 7 (a) Chemical structures of G4 ligands. (b) NMR structure of G4 with PhenDC ligand (PDB ID: 2MGN).²⁶³ (c) Crystal structure of TOP1cc with camptothecin (PDB ID: 1T8I).³¹¹ (d) Bioorthogonal copper catalysed click ligation performed in cells between PDS α and an azide-labelled fluorophore.

cells, positioning ATR upstream of ATM in the telomeric DNA damage signalling cascade (Table 2).²⁸⁰ CHK2 was also activated upon RHSP4 treatment and p53/p21 found to be the principal transducers inducing cell death in the same cell line.²⁸¹ In the same study, RHSP4 was shown to induce damage preferentially in replicating cells, where damage signalling was largely ATR-mediated, indicating its mechanism is carried out by replication fork stalling through stabilisation of G4 structures. This was evidenced by a significant colocalisation of DNA damage marker γ H2AX with replication marker PCNA, whereas ionising radiation (IR) induced γ H2AX foci independent of PCNA.²⁸¹ Additionally, helicase activity (WRN and BLM) increased upon RHSP4 treatment, and BLM^{KD} cells showed an increased DDR response to RHSP4-induced damage, further confirming that RHSP4 elicits its damage-inducing activity through the stabilisation of G4 structures.

360A: the pyridodicarboxamide 360A displays a high affinity for G4 and exhibits strong telomerase inhibition *in vitro*.²⁸² The tritiated 360A (^3H -360A) was found to localise at telomeres in T-cell lymphoblastic peripheral blood lymphocytes and human glioma cells by autoradiography.²⁸³ 360A is a telomere-specific DDR-inducing agent for which ATM was first described as an important DDR mediator, since 360A treatment triggers an increase of chromatin rearrangements and telomere damage (TIFs, telomeric γ H2AX and 53BP1 foci) in ATM deficient patient-derived cells and in ATM^{KD} HeLa cells compared to ATM-proficient cells.²⁸⁴ Later, it was shown that 360A treatment induced higher levels of telomere aberrations in ATR^{KD} cells compared to wild type, implicating ATR, as well as ATM, in the response to 360A.²⁸⁵ To further investigate the nature of the repair mechanism triggered on 360A-mediated damage induction, both RAD51- and DNA-PK-depleted cells were treated with

Table 2 Synthetic lethal interactions between G4 stabilising drugs and genetic DDR defects

Ligand	Sensitising genetic change (knockout/knockdown)	Mechanism implicated	Marker	Cell type/model	Ref.
TMS	Helicase FANCJ ^{KD}	Apoptosis	γH2AX	HeLa adenocarcinoma cells, telomerase-negative U2OS osteosarcoma cells	235 and 312
PDS, CX-5461 RHP54	Helicase FANCJ ^{KD}	Replication stress		HeLa cells	313
ATR ^{KD}		Telomere damage	γH2AX, Rad17 and 53BP1 colocalisation with TRF1	Breast and colon carcinoma-derived mice xenograft	280
PhenDC	Pif1 ^{KO} ; KO of Rad51, Rad52 and Rad54 desensitises cells	Chromosomal rearrangements		Yeast <i>Saccharomyces cerevisiae</i>	229, 246 and 288
TMS 360A	ATRX ^{KO}	DNA damage, apoptosis	γH2AX	Murine neuroprogenitor cells	314
	ATM ^{KD} , ATR ^{KD} , RAD51 ^{KD} , DNA-PKcs ^{KD}	Telomere damage and dysfunction,	γH2AX/TRF1 colocalisation, CHK2, p53, RAD51, DNA-PK	HeLa, HCT116 colorectal carcinoma, SV40-immortalised As3wt2 human fibroblasts	284–286
PhenDC	Helicase BLM ^{KD}	Induction of subtelomeric origins, replication fork slowing, increased G4 in telomeres	BG4 antibody staining colocalised with TRF1	SV40-immortalised mouse ear fibroblasts	240
PDS, PhenDC CX-5461, CX-3543 (PDS)	RAD51 ^{KD} , BRCA1 ^{−/−} , BRCA2 ^{−/−} (PDS only)	PARP1 cleavage, apoptosis, cell death	γH2AX	MEF fibroblasts	289
	BRCA2 ^{KO} , BRCA1 ^{KD} , (BRCA2 ^{KD} in U2OS), DNA-PK ^{KO} , LIG4 ^{KO}	DNA damage, apoptosis	γH2AX, 53BP1, DNA damage (Comet assay), CHK1/2, RAD51	HCT116 colon cancer cells, BRCA2 ^{+,−} ovarian, BRCA2 ^{+,−} breast, U2OS, HCT116-derived and patient-derived xenografts	304
CX-5461, PDS	TDP2 ^{KO}	Inhibition of TOP2cc repair		RPE-1 fibroblasts	79
PDS	DNA-PKcs ^{KO}	Inhibition of NHEJ-dependent DNA repair		DNA-PK ^{−/−} MO59J vs. DNA-PK ^{+/+} MO59K glioblastoma cells	92 and 315
PDS	XPF ^{KO/KD} and FANCM ^{KO/KD} (HR/DSB repair mediators)	Stalled replication forks (DNA fibre analysis)	γH2AX, CHK1 and RPA2	U2OS	316
PDS	Polq ^{KO/KD} (Polq protein is involved in alternative end joining DSB repair)	Chromosomal rearrangements, cell death		MEF	317

360A,²⁸⁶ RAD51 being associated with HR, while DNA-PK with NHEJ (see Section 2b). Both Rad51^{KD} and DNA-PK^{KD} cells incubated with 360A showed a progressive decline and eventual stop in cell growth. DNA-PK^{KD} was found to inhibit the 360A-induced telomere fusions observed in WT cells, while Rad51^{KD} increased telomere losses and formation of telomere doublets. These results highlight the central role of both HR and NHEJ in 360A-mediated damage repair.

PhenDC: the design of PhenDC3 and PhenDC6 resulted from the structural optimisation of 360A, changing V-shape for U-shape ligands to increase the overlap with the external G-quartet of a G4,²⁸⁷ as demonstrated by NMR (Fig. 7b).²⁶³ PhenDC displays tremendous G4-interaction and telomerase inhibition *in vitro*, sequestering telomere DNA into its G4 fold. PhenDC has been found to protect telomeres against nucleolytic resection by Exo1²⁰³ and to trigger genetic instability in budding yeast as a result of Pif1 helicase inhibition.²²⁹ A later study by the same group indicated that genetic instability resulted from disrupted HR repair mechanisms (misaligned synthesis-dependent strand annealing reactions), and G4-mediated large-scale genomic rearrangements (in a recombination-dependent manner involving RAD51, RAD52 and RAD54) occurring when G4s fold on the leading strand.²⁸⁸ Similar observations were reported in human cells where PhenDC counteracts helicase processivity (BLM and WRN) and decreases replication

fork speed, particularly at telomeres,²⁴⁰ as well as triggering DSBs upon loss of PARP3, which acts as a promoter of chromosomal rearrangements primarily occurring *via* NHEJ.¹¹⁴ PhenDC treatment of BRCA1-, BRCA2- and RAD51-depleted cells leads to apoptosis, further confirming its capacity to jeopardise HR-mediated DSB repair.²⁸⁹ *Via* an unbiased genome-wide short hairpin RNA (shRNA) genetic silencing screen, a series of PhenDC sensitiser genes was identified (many were in common with PDS treatment, see Table 2), chief among them BRCA1 and TOP1.²⁹⁰

Pyridostatin (PDS): the pyridodicarboxamide PDS displays exceptional *in vitro* binding properties.^{92,291} Initially designed as a telomere-targeting agent, it was rapidly shown that PDS targets both genomic G4s (2–5 μM) and telomeric G4s (> 5 μM).⁹² PDS treatment triggers DNA damage, evidenced by phosphorylation of H2AX, and ATM/ATR pathway activation.^{92,289} PDS stabilises G4 during both transcription and replication, causing DNA and RNA polymerase stalling.^{92,292} DNA damage induced by PDS is repaired in both a NHEJ- and HR-dependent manner.^{92,289,293} PDS treatment of BRCA2^{KD} or RAD51^{KD} cells showed robust activation of CHK1 and RPA thus implicating ATM/ATR signalling. More recently, independent non-biased genetic approaches showed that the cytotoxic effects of PDS (and CX-5461, see Table 2) are dependent on TOP2 activity suggesting that PDS could trap TOP2 at G4.^{79,91} Beyond its therapeutic properties,



PDS is also an exquisite molecular tool to study G4 both *in vitro* and *in vivo*. For instance, PDS is widely used to favour G4 formation in cell lysate for G4-seq and rG4-seq sequencing-based techniques.²⁵⁸ Several derivatives of PDS have been designed and used to assess the relevance of G4 in cells. For instance, a biotinylated PDS was used to pull down G4s from cell extracts,²⁹⁴ and an alkynylated PDS (PDS- α) was used to localise G4s directly in the nucleus by fluorescence microscopy after copper-catalysed alkyne azide cycloaddition (CuAAC) click ligation with a fluorophore in fixed cells⁹² (Fig. 7d). This bioorthogonal technique to visualise a small molecule's cellular localisation *in situ* is particularly precise due to its non-perturbing nature;²⁹⁵ the small molecule of interest with only a minuscule chemical modification binds to its biological target, and can be observed following specific conjugation with the clickable fluorophore, thus introducing the least experimental bias possible. This was a cornerstone study as PDS- α -binding sites were found to significantly co-localise with γ H2AX and only minorly with telomeric TRF1, indicating that PDS induces DNA damage primarily at genomic G4 sites. Beyond building the link between G4s and DNA damage, this approach also provided the very first description of the G4 landscape within human cells (*via* an accurate mapping of the distribution of G4s in the endogenous chromatin context) and highlighted the druggability of the SRC proto-oncogene (involved in multiple pathways that regulate tumour progression),²⁹⁶ thus uncovering a novel G4-mediated anticancer strategy. Recently, a PDS analogue conjugated to a fluorophore (SiR-PyPDS) was used in single-molecule live-cell fluorescence imaging to track dynamic G4 formation,¹⁸⁵ demonstrating the density and lifetime of G4s, and showing a significant increase in G4s in cells undergoing transcription and replication compared to steady state (G0/G1) cells. An alkynylated PDS derivative was chemically ligated to a ICL-inducing nitrogen mustard chlorambucil (Chl) (by CuAAC *in vitro*) to form a G4 ligand-cross-linking conjugate²⁹³ (Fig. 7d). This PDS-Chl drug conjugate showed a greater specificity for NER-deficient cancer cells compared to each of the two drugs alone. Such G4-targeted drug conjugates could be further exploited in NER-related diseases such as skin, testicular and drug resistant cancers. Finally, PDS has been used in non-cancer studies, to demonstrate a G4-mediated down-regulation of BRCA1²⁹⁷ and Autophagy related Gene (ATG7) in neurons,²⁴⁹ thus linking G4s to ageing and neurodegenerative diseases.¹⁸⁸

CX-5461: fluoroquinolones are a group of molecules described to have topoisomerase II (TOP2) and G4 binding affinities.^{191,298} Structural optimisation of fluoroquinolones to improve G4 *versus* TOP2 affinity first gave rise to CX-3543 (also known as quarfloxin), the first G4-interacting ligand to reach clinical trials. The described mechanism of action involves the disruption of nucleolin binding to G4-containing ribosomal DNA resulting in the inhibition of RNA polymerase 1 (Pol1) (overexpressed in tumours due to their increased need for protein production) leading to cancer cell death.²⁹⁹ CX-5461 was designed subsequently and described again as a Pol1 inhibitor although with no mention of G4 interaction in publications from Cylene Pharma.^{300,301} In an independent study, CX-5461 was later shown to induce a DDR response (ATR/ATM activation) and G2 phase cell cycle arrest in leukaemia cells.³⁰² This

unexpected behaviour was later correlated with its ability to stabilise G4s and stall replication forks.^{303,304} In this study, the ability of CX-5461 to interact with G4s was established both *in vitro* (*via* FRET-melting assay) and *in vivo* (with a significant increase in the G4 nuclear foci labelled with the G4-specific antibody BG4).³⁰⁵ BRCA2-depleted cells were found far more sensitive to CX-5461 treatment compared to WT cells, supporting an impact of G4 ligands on replication forks and implicating HR in the repair of the resulting structures. Chromatin immunoprecipitation (ChIP) of RAD51 and ChIP-seq studies performed in U2OS cells supported that some of the DNA damage caused by CX-5461 is repaired by RAD51.³⁰⁴ NHEJ is also involved in the response to CX-5461 as a hypersensitivity to CX-5461 was also observed in DNA-PK^{KD} cells. However, recent unbiased genetic approaches supported that both CX-5461 and PDS trigger DNA damage and mediate their cytotoxic effect through TOP2 interaction more specifically with TOP2A.^{79,91,306} These studies provide mechanistic insights into how CX-5461, PDS and potentially other G4 ligands produce DNA damage through structure specific interference with DNA topoisomerases.

To conclude, these six ligands are illustrative examples of G4 ligands that trigger DNA damage with varying ability, leading to multiple and varying cellular responses, all of them with clinical potential. A more comprehensive view of the genetic modifications described to sensitise cells to G4 ligands is given below, and illustrates the extreme correlation with DDR pathways (Table 2). New ligands are regularly reported (*e.g.* SYUIQ, 20A and FG) with yet again slightly different DNA damaging specificities.^{307–309} All of these chemicals target G4 *in vitro* and *in vivo*, and only with the help of non-biased analyses,^{79,91,290} will we decipher how stabilisation of these structures translates into phenotypes that could be exploited to cure diseases. Surprisingly or not, topoisomerase inhibition (TOP1/2) is often evoked, leading one to consider that G4 ligands could trap DNA-bound TOP1 or TOP2 complexes in a structure-specific manner. TOP1/2 are responsible for untwisting DNA under torsional strain *via* formation of a topoisomerase–DNA cleavage complex.⁵⁸ Topoisomerase poisons are commonly used anti-cancer agents (see Section 1b). They behave as interfacial inhibitors inserting into the topoisomerase–DNA complex and thereby blocking the religation step of the topoisomerase catalytic cycle and stabilising the cleavage complex (cc, TOP1cc or TOP2cc) (Fig. 7c). This is exemplified by prototype anticancer agents camptothecin (TOP1 poison, active site crystal structure in Fig. 7c),³¹⁰ doxorubicin and etoposide (TOP2 poisons).⁶⁰

The advantage that G4 ligands would then have over classical topoisomerase inhibitors is a double specificity, for a structure of nucleic acids and for regulatory sequences in which G4 forming sequences are more prevalent. It is to be noted that topoisomerases accumulate at sites of DNA aberrations,⁶² and maybe even at G4s. Only further studies will help gain insights into the actual targets of G4 ligands in cells, with the hope of gaining “*a deep understanding of cell circuitry and diseases biology*” that the G4s are involved in.

3d. Drug combination with G4-ligands. Radiotherapy induces DSBs which are repaired by HR/NHEJ, whilst antiproliferative



Table 3 Synthetic lethal interactions between G4 stabilising drugs and pharmacological DDR inhibition

Ligand	Synergistic DDR drug/treatment	DDR mechanism implicated	Marker	Cell type/model	Ref.
TMS	Imatinib, daunorubicin, mitoxantrone, vincristine	Telomere damage	CHK2, ATM, p21, p27	K562, OM9;22 leukaemia cells	271
TMS	NSC19630 (WRN helicase inhibitor)	Stalled replication, DNA damage, cell death	γ H2AX, PCNA	U2OS	318
RHSP4	Taxol	Telomere damage, chromosome fusions	γ H2AX/TRF1	UXF1138L cells and xenograft	319
RHPS4	Camptothecin (TOP1 inhibitor) then RHPS4 (CI < 0.2), bleomycin (CI \approx 0.7)	Apoptosis, tumour growth inhibition	γ H2AX	M14 melanoma cells, HT29 colon-derived xenograft	320
RHPS4	GPI 15427 (PARP inhibitor) followed by RHPS4, camptothecin SN-38 or ST1484 (TOP1 inhibitors) followed by RHPS4 (CI \approx 0.1)	Telomere damage, chromosome aberrations (or PARP1 ^{KD} sensitisation)	PARP/telomere colocalisation	BJ fibroblasts, HT29-derived xenografts	321
RHPS4	No synergy with adriamycin and doxorubicin (TOP2a inhibitors) SN-38 (CI < 0.5)	Telomere damage, cell death	γ H2AX, TOP1 accumulation at telomeres	BJ fibroblasts, HT29	322
RHSP4 derivates		Telomere damage, stalled replication	γ H2AX/TRF1, PCNA	HT29	323
RHPS4	Ionising radiation (IR)	Telomere damage, G2-phase block, chromosomal rearrangements, tumour growth inhibition	γ H2AX and 53BP1/TRF1 colocalisation	U251MG glioblastoma cells and xenograft	324
PDS	NU7441 (DNA-PK inhibitor)	Induction of DNA:RNA hybrid structures (R-loops), DNA damage, DSBs and DDR signalling	γ H2AX, 53BP1, ATM, antibody colocalisation: BG4 (G4) and S9.6 (R-loop)	DNA-PK ^{-/-} MO59J vs. DNA-PK ^{+/+} MO59K glioblastoma cells	325
PDS, FG	NU7441 (DNA-PK inhibitor)	DSBs, ATM-dependent DDR	γ H2AX, ATM	SV40-transformed MRC-5 fibroblasts	92
PDS, PDSI	NU7441 (DNA-PK inhibitor)	DSBs, DDR, apoptosis	53BP1, RAD51 (or BRCA2 ^{-/-})	HCT116 colon cancer cells	315
PDS	MK1775 (WEE1 cell cycle kinase inhibitor) and pimozide (FA repair USP1 inhibitor)	G4 stabilisation, cell death	Sensitised to KD of BRCA1&2, POLQ, USP1, TOP1, WEE1 etc.	A375 melanoma cells, HT1080 fibrosarcoma cells	290
PhenDC		G4 stabilisation, cell death	Sensitised to KD of BRCA1 and TOP1		
PDS, PhenDC	ME0328 (PARP3 inhibitor), KU0058948 (PARP1 inhibitor)	Chromosome rearrangements, DSBs, G2-phase block	γ H2AX, 53BP1, G4-binding 1H6 antibody staining, PARP3 ^{KO} sensitised	A549 lung adenocarcinoma cells	114
CX-5461, CX-3543.	IR, hydroxyurea	DNA damage	γ H2AX, 53BP1, CHK1/2, RAD51, ATRX ^{KD} sensitised	Murine normal human astrocytes, glioma stem cells	326
PDS					
CX-5461	BMN-673 (PARP inhibitor)	Pol1 transcription inhibition, replication stress, DNA damage, G2-phase block, no G4-stabilisation effect	ATM/ATR, RPA, γ H2AX, HR-deficient, BRCA1 ^{-/-} and MYC ^{-/-} cells sensitised	Ovarian cancer cells and xenografts	303
EMICORON	SN-38 then EMICORON; EMICORON then oxaliplatin or 5-FU (all CI < 0.5)	Telomeric DNA damage		HT29 colon cancer cells and xenografts	327

chemotherapeutic agents such as alkylating agents (TMZ) and cross-linking agents (nitrogen mustards and cisPt) induce lesions that are repaired by BER and NER/FA pathways respectively (see Section 1b). TOP1/2c stabilised by TOP1/2 poisons (Fig. 7c) are bulky DNA-protein lesions which are processed into SSBs or DSBs following collision with either the transcription or replication machinery.⁶¹ The cancer cell selectivity of all of these damage-inducing treatments relies on the principle that cancers are fast-replicating and DDR-deficient. Treatment regimens combining antiproliferative treatments with DDR inhibitors are proving efficient enough to reach clinical trials.⁸ G4 ligands are beginning to be viewed as DNA-damaging agents, and on the shoulders of intensive mechanistic studies of G4 ligand interactions, this chemical toolbox is now being trialled in combination therapies.

Mechanistic studies frequently describe a (hyper)sensitisation (increased growth inhibition) of cancer models to G4 ligands, when comparing the DDR-deficient cells/tissue with

the wild type (Table 2) or when cells/tissue are cotreated with antiproliferative agents or DDR inhibitors (Table 3). Exploiting DDR-deficient cancer cells has been shown for BRCA1/2-deficient cells, which are sensitised to G4 ligands (PDS,³¹⁵ PDS and PhenDC²⁸⁹) compared to BRCA1/2-proficient cells and one molecule, CX-5461, is in clinical trials for BRCA1/2-deficient tumours.³⁰⁴ PDS and CX-5461 exacerbate the damage occurring in DDR-deficient cells, similar to PARP inhibitors. In the same system, the damage-inducing effect was amplified by cotreatment with a DNA-PK inhibitor NU7441.³¹⁵

One of the earliest studies on TMS investigated the combinatorial effects with multiple antiproliferative agents.²⁷¹ Tauchi *et al.* described an additive effect of TMS with imatinib (inhibitor of anti-apoptotic tyrosine kinases), daunorubicin (TOP2 inhibitor), mitoxantrone (intercalator and TOP1 inhibitor) and vincristine (microtubule inhibitor), analysed by cell growth inhibition in single cell culture (Table 3). Several studies have



exploited the telomere selectivity of RHPS4 to induce anti-tumoural activity. Such interactions are often analysed by colocalisation of γ H2AX damage signalling with telomere marker TRF1, particularly in the case of RHPS4 and 360A, which exhibit various synergistic and sensitising interactions. WRN inhibitor NSC-19630 was found to induce replication-dependent DNA damage (γ H2AX/PCNA colocalisation), cell cycle inhibition and apoptosis as a single agent.³¹⁸ Since G4s are a known substrate of WRN helicase, NSC-19630 was combined with a G4-ligand to enhance these DDR-inducing effects. A non-toxic concentration of NSC-19630 (1 μ M, where $IC_{50} = 3$ μ M) in combination with a non-toxic concentration of TMS (0.6 μ M) was shown to induce 70% cell death in 3 days. Still, more data points are required to draw a strict conclusion on synergism.

Synergistic interactions can be confused with additive interactions. To be sure that a combinatorial effect is synergistic, a matrix of concentrations of the two chemical components should be analysed. One popular method pioneered by Chou and Talalay takes into account the shape of each IC_{50} curve (S-shaped or linear), in order to calculate a combination index (CI), where $CI < 1$ indicates synergism, $CI \approx 1$ indicates additivity and $CI > 1$ indicates antagonism.^{328,329} This method was used by Biroccio *et al.* to show strong synergistic effects between TOP1 inhibitors SN-38 or ST1484 and RHPS4 with extremely low combination indexes ($CI < 0.1$).³²² Another mathematical method to distinguish additive from synergistic effects is the Bliss independence model, in which the calculated additive effects are subtracted from the observed effects to produce a surface plot for a matrix of concentrations.³³⁰ In a field in which combinatory drug regimens are standard in the clinic, and clinical trials rarely test every combinatorial option (for a 2 drug combination of A and B, four regimens are possible: $\pm A/\pm B$), it is crucial to understand the interactions of each drug in fine detail in pre-clinical studies.

Many studies show that G4-induced damage occurs in both the HR and NHEJ pathways,²⁸⁶ and replication- and transcription-dependent mechanisms.⁹² Due to the vast range of interactions observed for G4 ligands, it is crucial to undertake unbiased experiments, despite the frequent and disappointing discovery of small molecules with poor specificity. CX-5461 was first described as a RNA Pol1 transcription inhibitor *via* a phenotypic screen, whilst its principal mechanism was more recently shown to be through G4-mediated interference with TOP2.³⁰⁴ Bruno *et al.* recently showed that CX-5461 incurs resistance to doxorubicin in a colorectal cancer model,³⁰⁶ an important finding because the TOP2 poison doxorubicin is a common treatment for colorectal cancer. CX-5461 is currently in clinical trials for relapse therapy based on the idea that it targets RNA Pol1-dependent transcription.³⁰³ However, the recent discovery that it acts through G4-dependent interference with TOP2 should lead to the patient/cancer selection process for this drug being revisited.^{79,91,306}

Advances in genome-wide sequencing techniques such as ChIP-seq and Chem-seq have facilitated the characterisation of genetic and epigenetic modulators.³³¹ A combination of specific DNA targeting agents with chromatin-interacting protein modulation (a field also currently undergoing important drug

development innovations)^{332–334} could prove to be another promising strategy in synthetic lethality chemotherapy.

In light of the wealth of preclinical data acquired with G4 ligands in the past recent years (Table 3), it is now tempting to develop drugs targeting other DNA folds that offer more defined anchoring sites than the external G-quartet. This opens new avenues to develop drugs with higher selectivity and new ways to interfere with DNA transactions in cancer cells. Their relevance to DDR and their therapeutic potential thus merit further investigation, which are highlighted in the following sections.

4. Non-B DNA structures

4a. Other higher-order DNA structures in the genome.

Repetitive DNA sequences make up 50% of the genome^{335–337} and are often found in regulatory non-coding regions. The transient and local deformation of the genomic duplex to form ssDNA during DNA transactions favours these repeated DNA sequences to fold up on themselves, forming molecular knots of varying structures and stability. In addition to G4s, a handful of non-B DNA structures have been shown to trigger genetic instability. To date, many examples have been described; some of them will be provided below (R-loops and DNA junctions) but this topic is regularly covered by authoritative reviews that interested readers are invited to refer to.^{11,17,18,335,338–340} Over ten types of non-B DNA structures have been described, including Z-DNA,³⁴¹ R-loops,^{342–345} triplex-DNA (or H-DNA)³⁴⁶ (see Fig. 8) with varying levels of biological and pathological relevance.

R-Loop structures were first described in 1976³⁴⁷ and currently are receiving renewed attention. R-Loops are DNA:RNA hybrids formed during transcription when the nascent RNA strand folds back and hybridises to the dsDNA sequence it was transcribed from, displacing the DNA strand of similar sequence to form a three-stranded RNA displacement loop, or R-loop.^{342,348} Initially thought to be formed only accidentally, genome-wide analyses such as DRIP-seq^{349,350} (DNA:RNA immunoprecipitation and sequencing) and DRIVE-seq³⁵⁰ (DNA:RNA *in vitro* enrichment and sequencing) highlighted the prevalence of R-loops in the human genome, whose distribution implies roles in DNA replication initiation, transcription termination and chromatin patterning.^{342,348} In a recent study, R-loops generated by RNA polymerase II localised to non-coding regions in ribosomal DNA and had positive regulatory effects on ribosomal RNA expression by RNA Pol1 in the nucleoli.³⁵¹ Aberrant R-loop formation threatens genomic stability, notably inducing replication and translational stress as they form roadblocks to polymerase processivity that promote DNA breaks and activate both ATM and ATR pathways. However, the relationship between DNA damage and R-loop formation is intricate as R-loops trigger DNA damage and conversely DNA damage favours R-loop formation.^{352,353} From a chemical biology point of view, R-loops thus represent another class of nucleic acid targets to damage DNA, an approach that has thus far only been attempted indirectly as no R-loop ligands are yet reported. For instance, R-loop formation was promoted by impeding canonical RNA maturation and spliceosome assembly^{353,354} using the TOP1 inhibitor diospyrin D1,³⁵⁵ or by stabilising G4 with PDS³⁵⁶ and monohydrazone



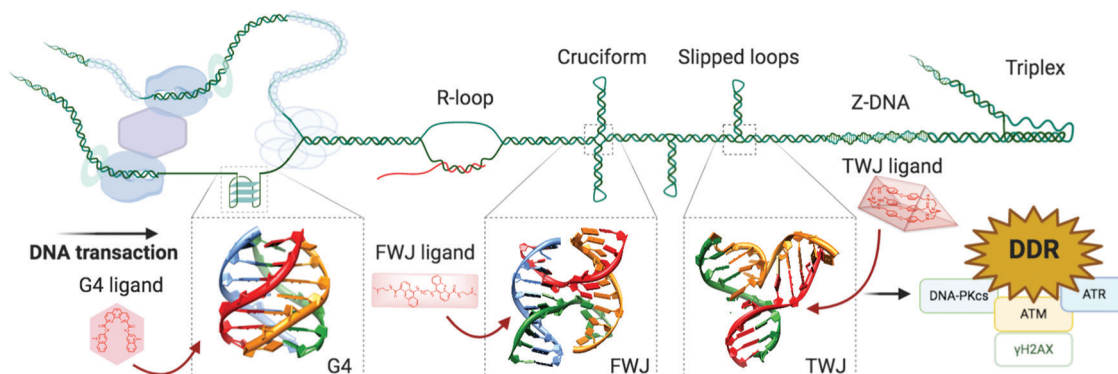


Fig. 8 Alternative DNA structures form in the vicinity of DNA transactions. The junction point of a cruciform structure presents a four-way junction (FWJ, e.g. Holliday junction) and that of a slipped loop presents a three-way junction (TWJ). Stabilisation of these structures with chemicals (ligands) can impede DNA transactions and induce DDR. Adapted from ref. 35, created with BioRender.

derivatives,³⁵⁷ since stabilised G4s promote the formation of R-loops at transcribed genes and the reciprocal G4/R-loop stabilisation can arrest polymerases and trigger DSBs accumulation.

Inverted repeat sequences are also highly prevalent in the human genome, and are also involved in genetic disorders, they are thus interesting targets for small molecules. Inverted repeats fold into hairpins stabilised by Watson–Crick-type hybridisation of an upstream region with the complementary downstream region. Two types of hairpin structures can be formed: if the inverted repeat-containing sequence forms two hairpins directly opposite to each other, the higher-order structure is referred to as cruciform DNA, or a four-way DNA junction (FWJ); alternatively, if the two hairpins are misaligned (shifted relative to each other), the structure is referred to as slippage loops, S-DNA, or three-way DNA junction (TWJ) (Fig. 8).^{11,358} The thermodynamic stability of these DNA junctions is high enough to act as physical obstacles that stall the replication fork/transcription bubble, creating replication- and transcription-coupled DNA damage that trigger DDR. Stabilisation of such blockages by external chemicals equally represents an alternative way to trigger DNA damage, with the strategic advantage of presenting structurally defined binding sites (chiefly the central cavity of the DNA junction, often referred to as the junction point) for small molecule targeting with high specificity (see Section 4b).

The quest for DNA junction-forming sequences in our genome has been far less intensive than that of QFS. However, several bioinformatic tools are being developed, such as the Palindrome analyser³⁵⁹ and the IRFinder,³⁶⁰ and preliminary results indicate that inverted repeat-forming sequences might be even more prevalent than QFS. For instance, comparison of the number of inverted repeat-forming sequences and QFS as identified with IRFinder³⁶¹ and G4-hunter³⁶² algorithms in the genome of *S. cerevisiae* indicates that there are >8000 inverted repeat-forming sequences (of 30 nt length) *versus* >3000 QFSs (of 25 nt length), with a strong enrichment at centromeric regions and replication origins. In the human genome, the advent of next-generation sequencing highlighted the prevalence of repeated sequences, accounting for \approx 50% of our

genome: the most frequent repeats are short and long interspersed nuclear elements (100–300 bp (base pairs) and >300 bp respectively) which account for 36% of the genome (>1 500 000 sequences), while short tandem repeats (referred to as microsatellites (2–10 bp), minisatellites (10–60 bp) and satellites (60–100 bp)) are adjacent repeating sequences and account for 3% of the genome (>400 000 sequences), a percentage comparable with the coding region of the genome.³³⁷ However, the proportion of inverted repeat-forming sequences among repeated sequences is less defined: they can be further classified as short and long inverted repeats (<30 bp and >30 bp respectively). Sergei M. Mirkin pioneered the quantification of short inverted repeats in a pre-sequencing era, developing an algorithm that showed that 50% of 40 kbp segments in the human genome contain at least one inverted repeat-forming sequence (with 2 to 6 nt loops).³⁶³ Subsequently, Karen M. Vasquez showed that short inverted repeats are significantly enriched at translocation breakpoints (an average of 28.5 short inverted repeats within a \pm 100 bp region surrounding the 20 000 breakpoints, *versus* 21.4 in 20 000 control sequences).³⁶⁴ Long inverted repeats were quantified in a more straightforward manner, with >2500 sequences in the human genome.³⁶⁵

While less studied than G4s, inverted repeats are also known to be responsible for genetic instability, which may have advantageous therapeutic payoffs (see Section 4b). Inverted repeats have been particularly studied in the context of the genetic disorders that hairpin-forming sequences are involved in, known as repeat expansion diseases.¹¹ Indeed, the genetic expansion (in which the sequence gets repeated during replication) of inverted repeats is frequently encountered in both cancers³⁶⁶ and severe neurological disorders (>50 human diseases).¹¹ Repeats are prone to expansion as a result of replication slippage, faulty replication and improper repair.^{18,367} Efforts have been invested to provide a unified model for explaining expansion (such as the breakage-fusion-bridge (BFB) model³⁶⁸), but the diversity of cellular mechanisms, events and protagonists that could account for repeat expansion explains why the mechanism behind expansion is still unknown. The most common expansion is the trinucleotide repeat expansion³⁶⁹ and related diseases (causative



of 16 different inheritable conditions), such as $(CAG)_n$ responsible for the Huntington disease, $(CGG)_n$ for the fragile X syndrome, and $(GGA)_n$ for Friedreich ataxia. The ability of these repeats to fold into hairpin structures has been firmly demonstrated.

As indicated in Section 3b, the cell has evolved the helicase machinery to resolve branched DNA structures^{219,370} which might be formed during DNA transactions and repair. Inverted repeats were significantly more present at replication fork stall sites in *E. coli*, *S. cerevisiae* and monkey fibroblasts cells, in a hairpin length-dependent manner.^{371,372} In another study, a non-biased RPA-ChIP-Seq sampling of replication fork stall sites reported that inverted repeats and structure-forming (purine-rich and AT-rich) microsatellite sequences were enriched at fork stall sites.³⁷³ This effect was amplified by chemical ATR inhibition, further indicating that the DDR machinery is used to straighten out these inverted repeat structures in homogenous conditions. The instability triggered herein is also counteracted by fork-stabilising proteins topoisomerase 1-associated factor 1 (Tof1) and Mannose Receptor C-Type 1 (Mrc1), that prevent irreversible fork collapse and DNA breakage.^{371,374} Several DDR mechanisms have been implicated in inverted repeat-induced damage and repair. Short inverted repeats were also shown to trigger genetic instability *via* DSB formation after being cleaved by the MRX (MRE11-RAD50-Xrs2 in *S. cerevisiae*, MRN in human) complex associated with Sae2 (SUMO1 activating enzyme subunit 2, CtIP in human)³⁷⁵ and repaired by HR,³⁷⁶ or cleaved by the ERCC1-XPF complex³⁶⁴ and repaired by the microhomology-mediated end joining (MMEJ) mechanism.³⁷⁷ These results explain why targeting inverted repeat-forming sequences with chemicals fosters lethal DNA damage and trigger DDR (Fig. 8).

4b. Non-B DNA targeting and drug combination. The therapeutic potential of DNA junction targeting has been exploited only marginally, despite an initial impetus provided in the late 1980s with the works of Neville R. Kallenbach and co-workers on the targeting of FWJ with chemicals.^{380–382} Early evidence that small molecules fit specifically into TWJs was provided by the use of non-natural TWJ-forming oligonucleotides (aptamers) that could encapsulate small hydrophobic molecules such as cocaine^{383,384} and steroids.^{385,386} The field of research was rejuvenated later with the resolution of the crystal structure of both FWJ:ligand³⁷⁸ and TWJ:ligand complexes³⁷⁹ (Fig. 9a and c), which provided invaluable indications for designing efficient DNA junction ligands. To date, only a few families of compounds have been studied for their ability to interact specifically with DNA junctions: FWJ ligands include organic dyes³⁸⁷ and porphyrins,³⁸² peptides^{388,389} (Fig. 9e) and macrocyclopeptides,^{390,391} acridines^{392,393} and bisacridine derivatives³⁷⁸ (Fig. 9b); TWJ ligands include helical supramolecular cylinders^{379,394} (Fig. 9d), helical metallopeptides,³⁹⁵ poly-aza-macrocycles,³⁹⁶ porphyrins,³⁹⁷ triptycenes,^{398,399} azacryptands⁴⁰⁰ (Fig. 9f), azacyclophanes^{401,402} (Fig. 9g), tetrahedral metallocages⁴⁰³ and calix[3]carbazoles.⁴⁰⁴ Several of these compounds were shown to inhibit cell growth, however only a few of them have been studied for their ability to trigger DNA damage, or studied in cells for any other mechanism of action.

The biological relevance of targeting inverted repeats was first demonstrated with the studies of peptidic ligands that bind the junction point of FWJs. *In vitro* and in *E. coli*, hexapeptides were capable of selectively binding the FWJ of a Holliday

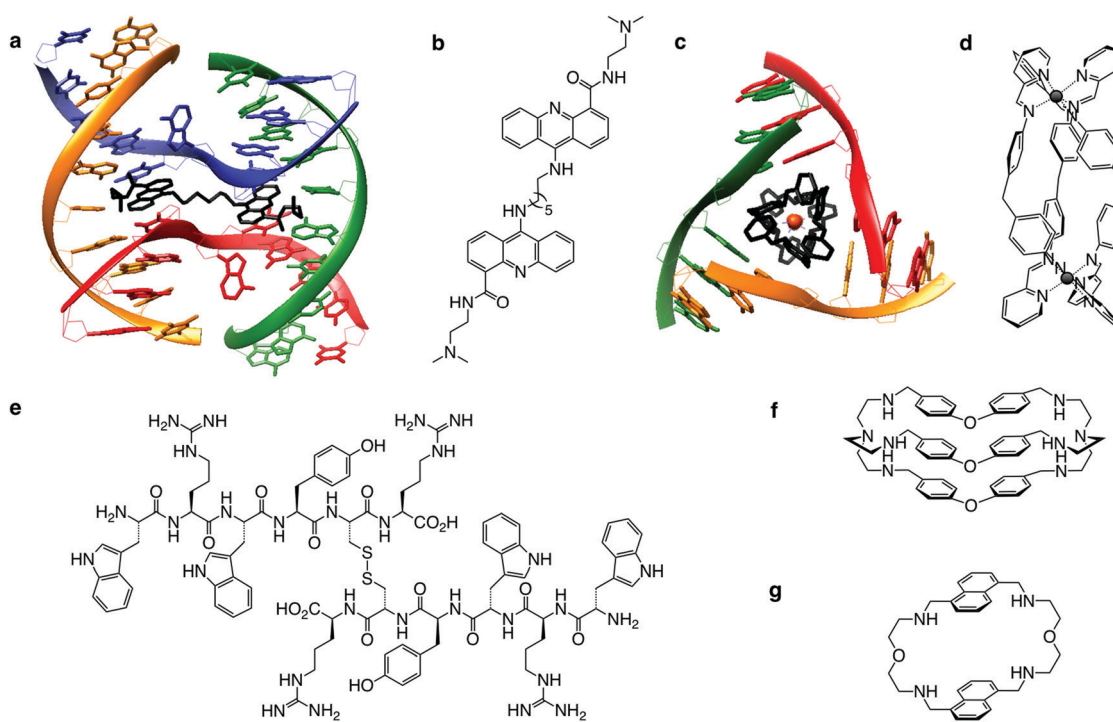


Fig. 9 (a and b) Crystal structure of FWJ with bisacridine ligand (PDB ID: 2GWA).³⁷⁸ (c and d) Crystal structure of TWJ with supramolecular iron cylinder (PDB ID: 2ETO).³⁷⁹ (e) FWJ ligand WRWYCR peptide in active disulfide form, TWJ ligands azacryptand TrisPOB (f) and azacyclophane 1,5-BisNP-O (g).



junction (a crucial intermediate during HR repair of both DSBs and replication-dependent ssDNA gaps, as demonstrated by live cell imaging and HJ-ChIP-seq),^{405,406} thus abrogating the action of Holliday junction-processing enzymes by displacing RecG helicase and inhibiting RuvABC resolvase, in the case of WRWYCR, for which the active form is the disulfide-bridged dimer (Fig. 9e).^{388,389} Furthermore, a similar more selective hexapeptide WKHYNY, triggered the accumulation of DNA breaks by blocking DNA repair through the inhibition of Cre recombinase,⁴⁰⁷ and was shown to act synergistically with other DNA damaging agents such as cross-linking agent mitomycin C.⁴⁰⁸ Later on, these peptides were shown to inhibit proliferation of numerous cancer cell lines (PC3, Du145, LnCAP, DuPro-1, PPC-1, HeLa and A549) *via* DNA damage induction, demonstrated by the accumulation of γ H2AX and 53BP1 foci and activation of CHK1 and CHK2.⁴⁰⁹ Again, the antiproliferative activities of these peptides were amplified by DSB-inducers (TOP2 inhibitors doxorubicin and etoposide) but not by unrelated therapeutics such as the mitotic inhibitor docetaxel, lending further credence to the DNA repair-based therapeutic activity of FWJ-ligands.

In these approaches, FWJ-targeting agents were used to impair DNA repair and their cellular activity is potentiated by DNA damaging agents. Similarly, in a more recent study, TWJ-targeting agents were shown to induce DNA damage, and their cellular activity potentiated by DNA repair inhibitors. The use of TWJ-targeting was greatly aided on obtaining the crystal structure of a supramolecular iron cylinder nestled into the junction point of a TWJ³⁷⁹ (Fig. 9c and d). This structure offers a brand new antiproliferative approach based on the hypothesis that an “*iron cylinder is able to bind to and stall DNA replication forks (a three-way junction structure)*”,⁴¹⁰ a hypothesis not entirely accurate given that replication forks are not TWJ structures *per se*. Despite being able to disrupt DNA transactions *in vitro*,⁴¹¹ these compounds were found to be poorly toxic in both cancer (HBL100, SKOV3, T47D and HL-60) and non-cancer (MRC5) cells, unable to trigger DNA damage, but capable of driving cells towards apoptosis.⁴¹⁰ Alternative investigations questioned both their TWJ-specificity^{412–416} and even the DNA-interacting properties of this class of ligands.⁴¹⁷

These results created momentum for further development, notably *via* the studies of cryptand-type ligands of two types: azacyclophanes, also known as cyclo-bis-intercalators; and azacryptands, also known as cyclo-tris-intercalators (Fig. 9f and g). These ligands, initially designed to target mismatched base pairs^{418,419} and abasic sites,^{420,421} have a 3D structure fully suited to fit within the TWJ junction point.⁴⁰⁰ The TWJ-interacting properties of these ligands were first thoroughly assessed *via* a series of *in vitro* assays (TWJ-screen, FRET-melting assays, etc.)^{401,402} to select candidates displaying high affinity and selectivity for TWJs, in order to provide a reliable *bona fide* model for interpreting cellular studies. The best candidates displayed antitumoral activity in the low micromolar range against MCF7 and MDA-MB-231 cells^{402,422} due to their ability to create DNA damage (γ H2AX accumulation) mostly at the G1/S transition. The TWJ ligands were then used in drug combinations with DDR inhibitors, targeting both HR

and NHEJ: NU7441 (DNA-PK inhibitor which impairs NHEJ, structure in Fig. 4b), KU55933 (ATM inhibitor which impairs DSB signalling and HR) and B02 (RAD51 inhibitor which impairs HR). Combination indexes (CI) as low as 0.39 (B02), 0.72 (KU55933) and 0.80 (NU7441) demonstrated that these azacryptands trigger both transcription- and replication-associated DSBs. The excellent synergy (CI = 0.39) obtained with azacryptand TrisPOB (Fig. 9f) and RAD51 inhibitor B02 supports that TWJ-ligands induce DSBs repaired by HR, potentially by promoting fork stalling in S and G2 phase.^{6,16}

Conclusion

Francis H. C. Crick, one of the founders of modern genetics, the co-discoverer of the double helix of DNA (1953)²⁶ and the father of the central dogma of biology (1957),^{423,424} wrote in 1974: “*we totally missed the possible role of enzymes in repair although (...) I later came to realise that DNA is so precious that probably many distinct repair mechanisms would exist*”.⁴²⁵ Because of this oversight, DNA repair has required a long time to be discovered, studied and understood. This lag also stems from the multiplicity of pathways involved in DNA repair thoroughly described above, along with their intricate cross-talk. As a consequence, a therapeutic approach that aims at damaging DNA is far more challenging than anticipated, as repair takes place under many different guises and innumerate feedback mechanisms to optimise its efficiency.

While it has been attempted to target the DNA double helix directly, we highlight here that DNA is a highly dynamic molecule that can adopt many higher-order structures, which represent a new class of targets to inflict severe injuries on cancer cells’ genome. The targeting of these structures can be performed with good-to-excellent specificity (as compared to the targeting of B DNA) and it has now been firmly demonstrated that the stabilisation of these structures by small molecules triggers extensive DNA damage that ultimately leads to cell death. Here, we focus on how damage is triggered in cells and describe the many different signalling and repair pathways that are activated in response to non-B DNA structure-mediated DNA damage.

Pioneering studies using G4 ligands have paved the way for understanding how the stabilisation of G4s creates damage and the therapeutic utility this presents in cancer. New pre-clinical and clinical data is emerging on the use DNA damaging agents (G4 ligands) with inhibitors of DNA repair, an approach referred to as combination therapy or chemically induced synthetic lethality. It is of unquestionable importance to perform detailed mechanistic investigations, and the recently developed, non-biased approaches such as chemoproteomics, chemogenomics and functional genomics are invaluable. Only an in-depth understanding of the damage caused by targeting alternative DNA structures can help define them as new biomarkers for cancers therapies.

The recent development of ligands which recognise other alternative DNA folds (such as three- and four-way DNA



junctions) are of immense value in determining their prevalence in cells, understanding their functions and assessing their relevance as therapeutic targets. Although in its infancy, this strategy benefits from the momentum provided by G4 studies (in terms of tools and strategies, techniques and assays, *etc.*) and, importantly, opens brand new therapeutic opportunities and chemical space. The recent demonstration that the stabilisation of DNA junctions triggers DNA damage enables the design of new therapeutic cocktails, combining ever more specific DNA damaging agents and DNA repair inhibitors, to halt cancer cell proliferation in a synergistic manner. Efforts must now be invested to validate this strategy at the pre-clinical/clinical levels, which will represent long but worthwhile efforts because, as cancer is a major pressing societal concern, all opportunities are valuable.

The examples depicted above show that alternative nucleic acid structures have gone a long way since their discovery, from being considered *in vitro* artefacts to biologically functional and targetable entities. These advances have been made possible thanks to an ever deeper understanding of the cell's inner workings which results from decades of resolutely multidisciplinary research combining molecular and cell biology, chemistry and biophysics. These results thus are a perfect demonstration of the absolute necessity to work at the interface between different, strongly interdependent scientific disciplines to enable science to meet societal expectations for the future.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 D. Hanahan and R. A. Weinberg, *Cell*, 2000, **107**, 57–70.
- 2 D. Hanahan and R. A. Weinberg, *Cell*, 2011, **144**, 646–674.
- 3 S. P. Jackson and J. Bartek, *Nat. Rev.*, 2009, **461**, 1071–1078.
- 4 M. J. O'Connor, *Mol. Cell*, 2015, **60**, 547–560.
- 5 A. Minchom, C. Aversa and J. Lopez, *Ther. Adv. Med. Oncol.*, 2018, **10**, 1–18.
- 6 R. Scully, A. Panday, R. Elango and N. A. Willis, *Nat. Rev. Mol. Cell Biol.*, 2011, **20**, 698–714.
- 7 N. J. O'Neil, M. L. Bailey and P. Hieter, *Nat. Rev. Genet.*, 2017, **18**, 613–623.
- 8 P. G. Pilié, C. Tang, G. B. Mills and T. A. Yap, *Nat. Rev. Clin. Oncol.*, 2019, **16**, 81–104.
- 9 D. Varshney, J. Spiegel, K. Zyner, D. Tannahill and S. Balasubramanian, *Nat. Rev. Mol. Cell Biol.*, 2020, **21**, 459–474.
- 10 R. Hänsel-Hertsch, M. Di Antonio and S. Balasubramanian, *Nat. Rev. Mol. Cell Biol.*, 2017, **18**, 279–284.
- 11 A. N. Khristich and S. M. Mirkin, *J. Biol. Chem.*, 2020, **295**, 4134–4170.
- 12 C. K. Kwok and C. J. Merrick, *Trends Biotechnol.*, 2017, **35**, 997–1013.
- 13 S. Neidle, *J. Med. Chem.*, 2016, **59**, 5987–6011.
- 14 S. Neidle, *Nat. Rev. Chem.*, 2017, **1**, 0041.
- 15 G. A. Holdgate, T. D. Meek and R. L. Grimley, *Nat. Rev. Drug Discovery*, 2018, **17**, 115–132.
- 16 N. Puget, K. M. Miller and G. Legube, *DNA Repair*, 2019, **81**, 102661.
- 17 J. Zhao, A. Bacolla, G. Wang and K. M. Vasquez, *Cell. Mol. Life Sci.*, 2010, **67**, 43–62.
- 18 B. P. Belotserkovskii, S. M. Mirkin and P. C. Hanawalt, *Chem. Rev.*, 2013, **113**, 8620–8637.
- 19 R. Dahm, *Dev. Biol.*, 2005, **278**, 274–288.
- 20 A. Hollaender and B. M. Duggar, *J. Bacteriol.*, 1938, **36**, 17–37.
- 21 E. C. Friedberg, *DNA Repair*, 2015, **33**, 35–42.
- 22 E. C. Friedberg, *Cell Res.*, 2008, **18**, 3–7.
- 23 A. Kelner, *Proc. Natl. Acad. Sci. U. S. A.*, 1949, **34**, 73–79.
- 24 R. Dulbecco, *Nature*, 1949, **163**, 949–950.
- 25 G. B. Sancar, *Mutat. Res.*, 2000, **451**, 25–37.
- 26 J. D. Watson and F. Crick, *Nature*, 1953, **171**, 737–738.
- 27 C. Dennis and P. Campbell, *Nature*, 1953, **171**, 740–741.
- 28 M. H. F. Wilkins, A. R. Wilson and H. R. Wilson, *Nature*, 1953, **171**, 738–740.
- 29 R. Beukers, J. Ylstra and W. Berends, *Recl. Trav. Chim. Pays-Bas*, 1958, **77**, 729–732.
- 30 S. Minato and H. Werbin, *Photochem. Photobiol.*, 1972, **15**, 97–100.
- 31 J. J. Madden and H. Werbin, *Biochemistry*, 1974, **13**, 2149–2154.
- 32 T. Lindahl and D. E. Barnes, *Cold Spring Harbor Symp. Quant. Biol.*, 2000, **65**, 127–133.
- 33 A. Ciccia and S. J. Elledge, *Mol. Cell*, 2010, **40**, 179–204.
- 34 D. Cortez, *DNA Repair*, 2015, **32**, 149–157.
- 35 H. Técher, S. Koundrioukoff, A. Nicolas and M. Debatisse, *Nat. Rev. Genet.*, 2017, **18**, 535–550.
- 36 T. Helleday, S. Eshtad and S. Nik-zainal, *Nat. Rev. Genet.*, 2014, **15**, 585–598.
- 37 A. N. Blackford and S. P. Jackson, *Mol. Cell*, 2017, **66**, 801–817.
- 38 S. I. Hajdu, *Cancer*, 2011, **117**, 1097–1102.
- 39 S. I. Hajdu, *Cancer*, 2012, **118**, 4914–4928.
- 40 D. Vilches, G. Alburquerque and R. Ramirez-Tagle, *Educ. Quim.*, 2016, **27**, 233–236.
- 41 J. Hirsch, *J. Am. Med. Assoc.*, 2006, **296**, 1518–1520.



42 K. W. Kohn, *Cancer Res.*, 1996, **56**, 5533–5546.

43 L. H. Hurley, *Nat. Rev. Cancer*, 2002, **2**, 188–200.

44 A. Gilman and F. S. Philips, *Science*, 1946, **103**, 409–436.

45 C. P. Rhoads, *Ca-Cancer J. Clin.*, 1978, **28**, 306–312.

46 P. Christakis, *Yale J. Biol. Med.*, 2011, **84**, 169–172.

47 L. Goodman, M. Wintrobe, W. Dameshek, M. Goodman, A. Gilman and M. McLennan, *J. Am. Med. Assoc.*, 1946, **132**, 126–132.

48 V. T. DeVita and E. Chu, *Cancer Res.*, 2008, **68**, 8643–8653.

49 K. W. Kohn, *Cancer Res.*, 1977, **37**, 1450–1454.

50 S. Mukherjee, *The Emperor of all Maladies: A Biography of Cancer*, Scribner, New York, Toronto, 1st edn, 2010.

51 S. Farber, L. K. Diamond, R. D. Mercer, R. F. Sylvester and J. A. Wolff, *N. Engl. J. Med.*, 1948, **238**, 787–793.

52 C. Heidelberger, N. K. Chaudhuri, P. Danneberg, D. Mooren, L. Griesbach, R. Duschinsky, R. J. Schnitzer, E. Pleven and J. Scheiner, *Nature*, 1957, **179**, 663–666.

53 D. B. Longley, D. P. Harkin and P. G. Johnston, *Nat. Rev. Cancer*, 2003, **3**, 330–338.

54 B. Druker, S. Tamura, E. Buchdunger, S. Ohno, G. M. Segal, S. Fanning, J. Zimmermann and N. B. Lydon, *Nat. Med.*, 1996, **2**, 561–566.

55 T. Hunter, *J. Clin. Invest.*, 2007, **117**, 2036–2043.

56 Y. Jung and S. J. Lippard, *Chem. Rev.*, 2007, **107**, 1387–1407.

57 K. Chválová, V. Brabec and J. Kašpáriková, *Nucleic Acids Res.*, 2007, **35**, 1812–1821.

58 J. L. Delgado, C. M. Hsieh, N. L. Chan and H. Hiasa, *Biochem. J.*, 2018, **475**, 373–398.

59 J. Marinello, M. Delcuratolo and G. Capranico, *Int. J. Mol. Sci.*, 2018, **19**, 3480.

60 J. L. Nitiss, *Nat. Rev. Cancer*, 2009, **9**, 338–350.

61 J. Stingele, R. Belletti and S. J. Boulton, *Nat. Rev. Mol. Cell Biol.*, 2017, **18**, 563–573.

62 Y. Pommier, *Nat. Rev. Cancer*, 2006, **6**, 789–802.

63 G. L. Patrick, *An Introduction to Medicinal Chemistry*, Oxford University Press, Oxford, UK, 5th edn, 2013.

64 R. A. Burrell, N. McGranahan, J. Bartek and C. Swanton, *Nature*, 2013, **501**, 338–345.

65 J. Huszno, Z. Kolasza and E. W. A. Grzybowska, *Oncol. Lett.*, 2019, **17**, 1986–1995.

66 I. Gorodetska, I. Kozeretska and A. Dubrovska, *J. Cancer*, 2019, **10**, 2109–2127.

67 S. Negrini, V. G. Gorgoulis and T. D. Halazonetis, *Nat. Rev. Mol. Cell Biol.*, 2010, **11**, 220–228.

68 E. R. Kastenhuber and S. W. Lowe, *Cell*, 2017, **170**, 1062–1078.

69 L. H. Pearl, A. C. Schierz, S. E. Ward, B. Al-Lazikani and F. M. G. Pearl, *Nat. Rev. Cancer*, 2015, **15**, 166–180.

70 P. C. Hanawalt, *DNA Repair*, 2015, **36**, 2–7.

71 Y. Shiloh and Y. Ziv, *Nat. Rev. Mol. Cell Biol.*, 2013, **14**, 197–210.

72 Y. Shiloh, *Nat. Rev. Cancer*, 2003, **3**, 155–168.

73 J. Stavnezer and C. E. Schrader, *J. Immunol.*, 2014, **193**, 5370–5378.

74 K. J. Zanotti and P. J. Gearhart, *DNA Repair*, 2016, **38**, 110–116.

75 I. Lam and S. Keeney, *Cold Spring Harbor Perspect. Biol.*, 2015, **7**, 0016634.

76 T. Lindahl, *Angew. Chem., Int. Ed.*, 2016, **55**, 8528–8534.

77 P. Modrich, *Angew. Chem., Int. Ed.*, 2016, **55**, 8490–8501.

78 A. Sancar, *Angew. Chem., Int. Ed.*, 2016, **55**, 8502–8527.

79 M. Olivieri, T. Cho, A. Álvarez-Quilón, K. Li, M. J. Schellenberg, M. Zimmermann, N. Hustedt, S. E. Rossi, S. Adam, H. Melo, A. M. Heijink, G. Sastre-Moreno, N. Moatti, R. Szilard, A. McEwan, A. K. Ling, A. Serrano-Benitez, T. Ubhi, S. Feng, J. Pawling, I. Delgado-Sainz, M. W. Ferguson, J. W. Ddennis, G. W. Brown, F. Cortés-Ledesma, R. S. Williams, A. Martin, D. Xu and D. Durocher, *Cell*, 2020, **182**, 481–496.

80 R. Abbotts and D. M. Wilson, *Free Radical Biol. Med.*, 2017, **107**, 228–244.

81 J. Jiricny, *Nat. Rev. Mol. Cell Biol.*, 2006, **7**, 335–346.

82 G. X. Reyes, T. T. Schmidt, R. D. Kolodner and H. Hombauer, *Chromosoma*, 2015, **124**, 443–462.

83 A. Sancar, *Nat. Genet.*, 1999, **21**, 247–249.

84 T. Nakano, S. Morishita, A. Katafuchi, M. Matsubara, Y. Horikawa, H. Terato, A. M. H. Salem, S. Izumi, S. P. Pack, K. Makino and H. Ide, *Mol. Cell*, 2007, **28**, 147–158.

85 D. J. Baker, G. Wuenschell, L. Xia, J. Termini, S. E. Bates, A. D. Riggs and T. R. O'Connor, *J. Biol. Chem.*, 2007, **282**, 22592–22604.

86 G. Spivak and A. K. Ganesan, *DNA Repair*, 2014, **19**, 64–70.

87 P. M. J. Burgers and T. A. Kunkel, *Annu. Rev. Biochem.*, 2017, **86**, 417–438.

88 L. J. Martin, *J. Neuropathol. Exp. Neurol.*, 2008, **67**, 377–387.

89 B. Sedgwick, P. A. Bates, J. Paik, S. C. Jacobs and T. Lindahl, *DNA Repair*, 2007, **6**, 429–442.

90 L. C. Huang, K. C. Clarkin and G. M. Wahl, *Proc. Natl. Acad. Sci. U. S. A.*, 1996, **93**, 4827–4832.

91 A. Pipier, M. Bossaert, J. F. Riou, C. Noirot, L.-T. Nguyêñ, R.-F. Serre, O. Bouchez, E. Defrancq, P. Calsou, S. Britton and D. Gomez, *bioRxiv*, 2020, DOI: 10.1101/2020.02.18.953851.

92 R. Rodriguez, K. M. Miller, J. V. Forment, C. R. Bradshaw, M. Nikan, S. Britton, T. Oelschlaegel, B. Xhemalce, S. Balasubramanian and S. P. Jackson, *Nat. Chem. Biol.*, 2012, **8**, 301–310.

93 K. Karanam, R. Kafri, A. Loewer and G. Lahav, *Mol. Cell*, 2012, **47**, 320–329.

94 L. Deriano and D. B. Roth, *Annu. Rev. Genet.*, 2013, **47**, 433–455.

95 A. Shibata, P. Jeggo and M. Löbrich, *DNA Repair*, 2018, **71**, 164–171.

96 R. L. Flynn and L. Zou, *Crit. Rev. Biochem. Mol. Biol.*, 2010, **45**, 266–275.

97 W. D. Wright, S. S. Shah and W. D. Heyer, *J. Biol. Chem.*, 2018, **293**, 10524–10535.

98 W. Zhao, J. B. Steinfeld, F. Liang, X. Chen, D. G. Maranon, C. Jian Ma, Y. Kwon, T. Rao, W. Wang, C. Sheng, X. Song, Y. Deng, J. Jimenez-Sainz, L. Lu, R. B. Jensen, Y. Xiong, G. M. Kupfer, C. Wiese, E. C. Greene and P. Sung, *Nature*, 2017, **550**, 360–365.



99 A. Datta and R. M. Brosh, *Genes*, 2019, **10**, 170.

100 G.-L. Moldovan and A. D. D'Andrea, *Annu. Rev. Genet.*, 2009, **43**, 223–249.

101 A. M. R. Taylor, C. Rothblum-Oviatt, N. A. Ellis, I. D. Hickson, S. Meyer, T. O. Crawford, A. Smogorzewska, B. Pietrucha, C. Weemaes and G. S. Stewart, *Nat. Rev. Dis. Prim.*, 2019, **5**, 64.

102 J. Niraj, A. Färkkilä and A. D. D'Andrea, *Annu. Rev. Cancer Biol.*, 2019, **3**, 457–478.

103 M. C. Kottemann and A. Smogorzewska, *Nature*, 2013, **493**, 356–363.

104 P. Tonzi, Y. Yin, C. W. T. Lee, E. Rothenberg and T. T. Huang, *eLife*, 2018, **7**, e41426.

105 H. L. Williams, M. E. Gottesman and J. Gautier, *Mol. Cell*, 2012, **47**, 140–147.

106 J. Murai, S. Y. N. Huang, B. B. Das, A. Renaud, Y. Zhang, J. H. Doroshow, J. Ji, S. Takeda and Y. Pommier, *Cancer Res.*, 2012, **72**, 5588–5599.

107 K. W. Caldecott, *Nat. Rev. Genet.*, 2008, **9**, 619–631.

108 C. J. Lord and A. Ashworth, *Science*, 2017, **355**, 1152–1158.

109 L. Palazzo and I. Ahel, *Biochem. Soc. Trans.*, 2018, **46**, 1681–1695.

110 B. A. Gibson, Y. Zhang, H. Jiang, K. M. Hussey, J. H. Shrimp, H. Lin, F. Schwede, Y. Yu and W. L. Kraus, *Science*, 2016, **353**, 45–50.

111 C. Beck, I. Robert, B. Reina-San-Martin, V. Schreiber and F. Dantzer, *Exp. Cell Res.*, 2014, **329**, 18–25.

112 S. L. Rulten, A. E. O. Fisher, I. Robert, M. C. Zuma, M. Rouleau, L. Ju, G. Poirier, B. Reina-San-Martin and K. W. Caldecott, *Mol. Cell*, 2011, **41**, 33–45.

113 C. Beck, C. Boehler, J. Guirouilh Barbat, M. E. Bonnet, G. Illuzzi, P. Ronde, L. R. Gauthier, N. Magroun, A. Rajendran, B. S. Lopez, R. Scully, F. D. Boussin, V. Schreiber and F. Dantzer, *Nucleic Acids Res.*, 2014, **42**, 5616–5632.

114 T. A. Day, J. V. Layer, J. Patrick Cleary, S. Guha, K. E. Stevenson, T. Tivey, S. Kim, A. C. Schinzel, F. Izzo, J. Doench, D. E. Root, W. C. Hahn, B. D. Price and D. M. Weinstock, *Nat. Commun.*, 2017, **8**, 15110.

115 N. Hustedt and D. Durocher, *Nat. Cell Biol.*, 2017, **19**, 1–9.

116 J. Falck, J. Coates and S. P. Jackson, *Nature*, 2005, **434**, 605–611.

117 A. Syed and J. A. Tainer, *Annu. Rev. Biochem.*, 2018, **87**, 263–294.

118 S. Britton, J. Coates and S. P. Jackson, *J. Cell Biol.*, 2013, **202**, 579–595.

119 M. Stucki, J. A. Clapperton, D. Mohammad, M. B. Yaffe, S. J. Smerdon and S. P. Jackson, *Cell*, 2005, **123**, 1213–1226.

120 M. S. Lee, R. A. Edwards, G. L. Thede and J. N. M. Glover, *J. Biol. Chem.*, 2005, **280**, 32053–32056.

121 M. Goldberg, M. Stucki, J. Falck, D. D'Amours, D. Rahman, D. Pappin, J. Bartek and S. P. Jackson, *Nature*, 2003, **421**, 952–956.

122 R. Scully and A. Xie, *Mutat. Res.*, 2013, **750**, 5–14.

123 J. R. Chapman and S. P. Jackson, *EMBO Rep.*, 2008, **9**, 795–801.

124 F. J. Hari, C. Spycher, S. Jungmichel, L. Pavic and M. Stucki, *EMBO Rep.*, 2010, **11**, 387–392.

125 S. Burma, B. P. Chen, M. Murphy, A. Kurimasa and D. J. Chen, *J. Biol. Chem.*, 2001, **276**, 42462–42467.

126 J. D. Friesner, B. Liu, K. Culligan and A. B. Britt, *Mol. Biol. Cell*, 2005, **16**, 2566–2576.

127 P. Caron, F. Aymard, J. S. Iacoboni, S. Briois, Y. Canitrot, B. Bugler, L. Massip, A. Losada and G. Legube, *PLoS Genet.*, 2012, **8**, e1002460.

128 I. Litwin, E. Pilarczyk and R. Wysocki, *Genes*, 2018, **9**, 581.

129 C. Meisenberg, S. I. Pinder, S. R. Hopkins, S. K. Wooller, G. Benstead-Hume, F. M. G. Pearl, P. A. Jeggo and J. A. Downs, *Mol. Cell*, 2019, **73**, 212–223.

130 Z. Lou, K. Minter-Dykhouse, X. Wu and J. Chen, *Nature*, 2003, **421**, 957–961.

131 G. S. Stewart, B. Wang, C. R. Bigneli, A. M. R. Taylor and S. J. Elledge, *Nature*, 2003, **421**, 961–966.

132 W. M. Bonner, C. E. Redon, J. S. Dickey, A. J. Nakamura, O. A. Sedelnikova, S. Solier and Y. Pommier, *Nat. Rev. Cancer*, 2008, **8**, 957–967.

133 P. Chanut, S. Britton, J. Coates, S. P. Jackson and P. Calsou, *Nat. Commun.*, 2016, **7**, 12889.

134 B. P. C. Chen, N. Uematsu, J. Kobayashi, Y. Lerenthal, A. Krempler, H. Yajima, M. Löbrich, Y. Shiloh and D. J. Chen, *J. Biol. Chem.*, 2007, **282**, 6582–6587.

135 D. Normanno, A. Négrel, A. J. de Melo, S. Betzi, K. Meek and M. Modesti, *eLife*, 2017, **6**, e22900.

136 M. Hammel, Y. Yu, B. L. Mahaney, B. Cai, R. Ye, B. M. Phipps, R. P. Rambo, G. L. Hura, M. Pelikan, S. So, R. M. Abolfath, D. J. Chen, S. P. Lees-Miller and J. A. Tainer, *J. Biol. Chem.*, 2010, **285**, 1414–1423.

137 S. Britton, P. Chanut, C. Delteil, N. Barboule, P. Frit and P. Calsou, *Nucleic Acids Res.*, 2020, DOI: 10.1093/nar/gkaa723.

138 M. D. Wilson, S. Benlekbir, A. Fradet-Turcotte, A. Sherker, J. P. Julien, A. McEwan, S. M. Noordermeer, F. Sicheri, J. L. Rubinstein and D. Durocher, *Nature*, 2016, **536**, 100–103.

139 S. M. Noordermeer, S. Adam, D. Setiaputra, M. Barazas, S. J. Pettitt, A. K. Ling, M. Olivieri, A. Álvarez-Quilón, N. Moatti, M. Zimmermann, S. Annunziato, D. B. Krastev, F. Song, I. Brandsma, J. Frankum, R. Brough, A. Sherker, S. Landry, R. K. Szilard, M. M. Munro, A. McEwan, T. G. de Rugy, Z. Y. Lin, T. Hart, J. Moffat, A. C. Gingras, A. Martin, H. van Attikum, J. Jonkers, C. J. Lord, S. Rottenberg and D. Durocher, *Nature*, 2018, **560**, 117–121.

140 S. Panier and S. J. Boulton, *Nat. Rev. Mol. Cell Biol.*, 2014, **15**, 7–18.

141 K. A. Cimprich and D. Cortez, *Nat. Rev. Mol. Cell Biol.*, 2008, **9**, 616–627.

142 V. Costanzo, D. Shechter, P. J. Lupardus, K. A. Cimprich, M. Gottesman and J. Gautier, *Mol. Cell*, 2003, **11**, 203–213.

143 F. A. Derheimer, H. M. O'Hagan, H. M. Krueger, S. Hanasoge, M. T. Paulsen and M. Ljungman, *Proc. Natl. Acad. Sci. U. S. A.*, 2007, **104**, 12778–12783.

144 K. Trenz, E. Smith, S. Smith and V. Costanzo, *EMBO J.*, 2006, **25**, 1764–1774.

145 J. M. Merchut-Maya, J. Bartek and A. Maya-Mendoza, *DNA Repair*, 2019, **81**, 102654.



146 L. I. Toledo, M. Altmeyer, M. B. Rask, C. Lukas, D. H. Larsen, L. K. Povlsen, S. Bekker-Jensen, N. Mailand, J. Bartek and J. Lukas, *Cell*, 2013, **155**, 1088.

147 E. J. Brown and D. Baltimore, *Genes Dev.*, 2000, **14**, 397–402.

148 A. De Klein, M. Muijtjens, R. Van Os, Y. Verhoeven, B. Smit, A. M. Carr, A. R. Lehmann and J. H. J. Hoeijmakers, *Curr. Biol.*, 2000, **10**, 479–482.

149 A. Jazayeri, J. Falck, C. Lukas, J. Bartek, G. C. M. Smith, J. Lukas and S. P. Jackson, *Nat. Cell Biol.*, 2006, **8**, 37–45.

150 M. Malumbres and M. Barbacid, *Nat. Rev. Cancer*, 2009, **9**, 153–166.

151 A. B. Williams and B. Schumacher, *Cold Spring Harbor Perspect. Med.*, 2016, **6**, a026070.

152 K. J. Barnum and M. J. O'Connell, *Methods Mol. Biol.*, 2014, **1170**, 29–40.

153 A. M. Abukhdeir and B. H. Park, *Expert Rev. Mol. Med.*, 2008, **10**, e19.

154 P. Nurse, *Nature*, 2005, **432**, 557.

155 C. J. Matheson, D. S. Backos and P. Reigan, *Trends Pharmacol. Sci.*, 2016, **37**, 872–881.

156 M. Chen, C. E. Ryan and H. Piwnica-Worms, *Mol. Cell. Biol.*, 2003, **23**, 7488–7497.

157 H. Löffler, B. Rebacz, A. D. Ho, J. Lukas, J. Bartek and A. Krämer, *Cell Cycle*, 2006, **5**, 2543–2547.

158 M. Donzelli and G. F. Draetta, *EMBO Rep.*, 2003, **4**, 671–677.

159 H. Wang, X. Zhang, L. Teng and R. J. Legerski, *Exp. Cell Res.*, 2015, **334**, 350–358.

160 T. A. Yap, R. Plummer, N. S. Azad and T. Helleday, *Am. Soc. Clin. Oncol. Educ. B*, 2019, 185–195.

161 M. Dobbelstein and C. S. Sørensen, *Nat. Rev. Drug Discovery*, 2015, **14**, 405–423.

162 S. Murata, C. Zhang, N. Finch, K. Zhang, L. Campo and E. K. Breuer, *BioMed Res. Int.*, 2016, 2346585.

163 W. G. Kaelin, *Nat. Rev. Cancer*, 2005, **5**, 689–698.

164 M. Shaheen, C. Allen, J. A. Nickoloff and R. Hromas, *Blood*, 2011, **117**, 6074–6082.

165 T. Dobzhansky, *Genetics*, 1946, **31**, 269–290.

166 M. Keung, Y. Wu and J. Vadgama, *J. Clin. Med.*, 2019, **8**, 435.

167 J. S. Brown, B. O'Carrigan, S. P. Jackson and T. A. Yap, *Cancer Discovery*, 2017, **7**, 20–37.

168 P. Munster, M. Mita, A. Mahipal, J. Nemunaitis, C. Massard, T. Mikkelsen, C. Cruz, L. Paz-Ares, M. Hidalgo, D. Rathkopf, G. Blumenschein, D. C. Smith, B. Eichhorst, T. Cloughesy, E. H. Filvaroff, S. Li, H. Raymon, H. de Haan, K. Hege and J. C. Bendell, *Cancer Manage. Res.*, 2019, **11**, 10463–10476.

169 B. Van Triest, L. Damstrup, J. Falkenius, V. Budach, E. Troost, M. Samuels, T. Goddemeier and P. F. Geertsen, *J. Clin. Oncol.*, 2017, **35**, e14048.

170 B. Haynes, J. Murai and J. M. Lee, *Cancer Treat. Rev.*, 2018, **71**, 1–7.

171 T. P. Matthews, A. M. Jones and I. Collins, *Expert Opin. Drug Discovery*, 2013, **8**, 621–640.

172 J. Shendure, S. Balasubramanian, G. M. Church, W. Gilbert, J. Rogers, J. A. Schloss and R. H. Waterston, *Nature*, 2017, **550**, 345–353.

173 C. Sánchez-Martínez, M. J. Lallena, S. G. Sanfeliciano and A. de Dios, *Bioorg. Med. Chem. Lett.*, 2019, **29**, 126637.

174 A. Bradbury, S. Hall, N. Curtin and Y. Drew, *Pharmacol. Ther.*, 2020, **207**, 107450.

175 S. Rundle, A. Bradbury, Y. Drew and N. J. Curtin, *Cancers*, 2017, **9**, 41.

176 Y. Wang and D. J. Patel, *Structure*, 1993, **1**, 263–282.

177 K. N. Luu, A. T. Phan, V. Kuryavyi, L. Lacroix and D. J. Patel, *J. Am. Chem. Soc.*, 2006, **128**, 9963–9970.

178 G. N. Parkinson, M. P. H. Lee and S. Neidle, *Nature*, 2002, **417**, 876–880.

179 S. Burge, G. N. Parkinson, P. Hazel, A. K. Todd and S. Neidle, *Nucleic Acids Res.*, 2006, **34**, 5402–5415.

180 D. Rhodes and H. J. Lipps, *Nucleic Acids Res.*, 2015, **43**, 8627–8637.

181 L. Stefan and D. Monchaud, *Nat. Rev. Chem.*, 2019, **3**, 650–668.

182 M. Gellert, M. N. Lipsett and D. R. Davies, *Proc. Natl. Acad. Sci. U. S. A.*, 1962, **48**, 2013–2018.

183 V. S. Chambers, G. Marsico, J. M. Boutell, M. Di Antonio, G. P. Smith and S. Balasubramanian, *Nat. Biotechnol.*, 2015, **33**, 877–881.

184 R. Hänsel-Hertsch, D. Beraldi, S. V. Lensing, G. Marsico, K. Zyner, A. Parry, M. Di Antonio, J. Pike, H. Kimura, M. Narita, D. Tannahill and S. Balasubramanian, *Nat. Genet.*, 2016, **48**, 1267–1272.

185 M. Di Antonio, A. Ponjavic, A. Radzevičius, R. T. Ranasinghe, M. Catalano, X. Zhang, J. Shen, L. M. Needham, S. F. Lee, D. Klenerman and S. Balasubramanian, *Nat. Chem.*, 2020, **12**, 832–837.

186 J. Spiegel, S. Adhikari and S. Balasubramanian, *Trends Chem.*, 2020, **2**, 123–136.

187 A. K. Todd, M. Johnston and S. Neidle, *Nucleic Acids Res.*, 2005, **33**, 2901–2907.

188 N. Maizels, *EMBO Rep.*, 2015, **16**, 910–922.

189 G. Marsico, V. S. Chambers, A. B. Sahakyan, P. McCauley, J. M. Boutell, M. Di Antonio and S. Balasubramanian, *Nucleic Acids Res.*, 2019, **47**, 3862–3874.

190 N. Kim, *Curr. Med. Chem.*, 2017, **26**, 2898–2917.

191 S. Balasubramanian, L. H. Hurley and S. Neidle, *Nat. Rev. Drug Discovery*, 2011, **10**, 261–275.

192 T. M. Bryan, *Molecules*, 2019, **24**, 3439.

193 C. Cayrou, B. Ballester, I. Peiffer, R. Fenouil, P. Coulombe, J. C. Andrau, J. Van Helden and M. Méchali, *Genome Res.*, 2015, **25**, 1873–1885.

194 P. Prorok, M. Artufel, A. Aze, P. Coulombe, I. Peiffer, L. Lacroix, A. Guédin, J. L. Mergny, J. Damaschke, A. Schepers, B. Ballester and M. Méchali, *Nat. Commun.*, 2019, **10**, 3274.

195 Y. Xu, *Chem. Soc. Rev.*, 2011, **40**, 2719–2740.

196 K. Turner, V. Vasu and D. Griffin, *Cells*, 2019, **8**, 73.

197 J. D. Watson, *Nature, New Biol.*, 1972, **239**, 197–201.

198 A. M. Olovnikov, *J. Theor. Biol.*, 1973, **41**, 181–190.

199 P. Martínez and M. A. Blasco, *J. Cell Biol.*, 2017, **216**, 875–887.

200 C. López-Otín, M. A. Blasco, L. Partridge, M. Serrano and G. Kroemer, *Cell*, 2013, **153**, 1194.



201 A. De Cian, L. Lacroix, C. Douarre, N. Temime-Smaali, C. Trentesaux, J. F. Riou and J. L. Mergny, *Biochimie*, 2008, **90**, 131–155.

202 H. Q. Yu, D. Miyoshi and N. Sugimoto, *J. Am. Chem. Soc.*, 2006, **128**, 15461–15468.

203 J. S. Smith, Q. Chen, L. A. Yatsunyk, J. M. Nicoludis, M. S. Garcia, R. Kranaster, S. Balasubramanian, D. Monchaud, M. P. Teulade-Fichou, L. Abramowitz, D. C. Schultz and F. B. Johnson, *Nat. Struct. Mol. Biol.*, 2011, **18**, 478–486.

204 E. H. Blackburn, *Angew. Chem., Int. Ed.*, 2010, **49**, 7405–7421.

205 C. W. Greider, *Angew. Chem., Int. Ed.*, 2010, **49**, 7422–7439.

206 J. W. Szostak, *Angew. Chem., Int. Ed.*, 2010, **49**, 7386–7404.

207 E. H. Blackburn, C. W. Greider and J. W. Szostak, *Nat. Med.*, 2006, **12**, 1133–1138.

208 Y. Deng, S. S. Chan and S. Chang, *Nat. Rev. Cancer*, 2008, **8**, 450–458.

209 O. Mendoza, A. Bourdoncle, J. B. Boulé, R. M. Brosh and J. L. Mergny, *Nucleic Acids Res.*, 2016, **44**, 1989–2006.

210 M. Sauer and K. Paeschke, *Biochem. Soc. Trans.*, 2017, **45**, 1173–1182.

211 R. M. Brosh, *Nat. Rev. Cancer*, 2013, **13**, 542–558.

212 R. M. Brosh and S. W. Matson, *Genes*, 2020, **11**, 255.

213 K. N. Estep and R. M. Brosh, *Biochem. Soc. Trans.*, 2017, **46**, 77–95.

214 P. Lansdorp and N. van Wietmarschen, *Genes*, 2019, **10**, 870.

215 L. K. Lerner and J. E. Sale, *Genes*, 2019, **10**, 95.

216 M. E. Fairman-Williams, U. P. Guenther and E. Jankowsky, *Curr. Opin. Struct. Biol.*, 2010, **20**, 313–324.

217 M. R. Singleton, M. S. Dillingham and D. B. Wigley, *Annu. Rev. Biochem.*, 2007, **76**, 23–50.

218 H. Sun, J. K. Karow, I. D. Hickson and N. Maizels, *J. Biol. Chem.*, 1998, **273**, 27587–27592.

219 P. Mohaghegh, J. K. Karow, R. M. Brosh, V. A. Bohr and I. D. Hickson, *Nucleic Acids Res.*, 2001, **29**, 2843–2849.

220 J. L. Li, R. J. Harrison, A. P. Reszka, R. M. Brosh, V. A. Bohr, S. Neidle and I. D. Hickson, *Biochemistry*, 2001, **40**, 15194–15202.

221 H. Han, R. J. Bennett and L. H. Hurley, *Biochemistry*, 2000, **39**, 9311–9316.

222 H. Sun, R. J. Bennett and N. Maizels, *Nucleic Acids Res.*, 1999, **27**, 1978–1984.

223 M. Fry and L. A. Loeb, *J. Biol. Chem.*, 1999, **274**, 12797–12802.

224 X. Wu and N. Maizels, *Nucleic Acids Res.*, 2001, **29**, 1765–1771.

225 S. D. Creacy, E. D. Routh, F. Iwamoto, Y. Nagamine, S. A. Akman and J. P. Vaughn, *J. Biol. Chem.*, 2008, **283**, 34626–34634.

226 J. P. Vaughn, S. D. Creacy, E. D. Routh, C. Joyner-Butt, G. S. Jenkins, S. Pauli, Y. Nagamine and S. A. Akman, *J. Biol. Chem.*, 2005, **280**, 38117–38120.

227 M. C. Chen, P. Murat, K. Abecassis, A. R. Ferre-D'Amare and S. Balasubramanian, *Nucleic Acids Res.*, 2015, **43**, 2223–2231.

228 C. Ribeyre, J. Lopes, J. B. Boulé, A. Piazza, A. Guédin, V. A. Zakian, J. L. Mergny and A. Nicolas, *PLoS Genet.*, 2009, **5**, e1000475.

229 A. Piazza, J. B. Boulé, J. Lopes, K. Mingo, E. Largy, M. P. Teulade-Fichou and A. Nicolas, *Nucleic Acids Res.*, 2010, **38**, 4337–4348.

230 K. Paeschke, M. L. Bochman, P. Daniela Garcia, P. Cejka, K. L. Friedman, S. C. Kowalczykowski and V. A. Zakian, *Nature*, 2013, **497**, 458–462.

231 S. Dehghani-Tafti, V. Levdkov, A. A. Antson, B. Bax and C. M. Sanders, *Nucleic Acids Res.*, 2019, **47**, 3208–3222.

232 J. B. Vannier, S. Sandhu, M. I. R. Petalcorin, X. Wu, Z. Nabi, H. Ding and S. J. Boulton, *Science*, 2013, **342**, 239–242.

233 J. B. Vannier, V. Pavicic-Kaltenbrunner, M. I. R. Petalcorin, H. Ding and S. J. Boulton, *Cell*, 2012, **149**, 795–806.

234 T. B. C. London, L. J. Barber, G. Mosedale, G. P. Kelly, S. Balasubramanian, I. D. Hickson, S. J. Boulton and K. Hiom, *J. Biol. Chem.*, 2008, **283**, 36132–36139.

235 Y. Wu, K. Shin-ya and R. M. Brosh, *Mol. Cell. Biol.*, 2008, **28**, 4116–4128.

236 S. K. Bharti, J. A. Sommers, F. George, J. Kuper, F. Hamon, K. Shin-Ya, M. P. Teulade-Fichou, C. Kisker and R. M. Brosh, *J. Biol. Chem.*, 2013, **288**, 28217–28229.

237 Y. Wu, J. A. Sommers, I. Khan, J. P. De Winter and R. M. Brosh, *J. Biol. Chem.*, 2012, **287**, 1007–1021.

238 L. T. Gray, A. C. Vallur, J. Eddy and N. Maizels, *Nat. Chem. Biol.*, 2014, **10**, 313–318.

239 A. R. Lehmann, *Genes Dev.*, 2001, **15**, 15–23.

240 W. C. Drosopoulos, S. T. Kosiyatrakul and C. L. Schildkraut, *J. Cell Biol.*, 2015, **210**, 191–208.

241 N. Van Wietmarschen, S. Merzouk, N. Halsema, D. C. J. Spierings, V. Guryev and P. Lansdorp, *Nat. Commun.*, 2018, **9**, 271.

242 T. De Lange, *Genes Dev.*, 2005, **19**, 2100–2110.

243 I. Cheung, M. Schertzer, A. Rose and P. Lansdorp, *Nat. Genet.*, 2002, **31**, 405–409.

244 R. A. Schwab, J. Niemuszczky, K. Shin-ya and W. Niedzwiedz, *J. Cell Biol.*, 2013, **201**, 33–48.

245 K. Paeschke, J. A. Capra and V. A. Zakian, *Cell*, 2011, **145**, 678–691.

246 A. Piazza, M. Adrian, F. Samazan, B. Heddi, F. Hamon, A. Serero, J. Lopes, M. Teulade-Fichou, A. T. Phan and A. Nicolas, *EMBO J.*, 2015, **34**, 1718–1734.

247 L. Maestroni, J. Audry, P. Luciano, S. Coulon, V. Géli and Y. Corda, *Cell Stress*, 2020, **4**, 48–63.

248 S. Jimeno, R. Camarillo, F. Mejías-Navarro, M. J. Fernández-Ávila, I. Soria-Bretones, R. Prados-Carvajal and P. Huertas, *Cell Rep.*, 2018, **24**, 3262–3273.

249 J. F. Moruno-Manchon, P. Lejault, Y. Wang, B. McCauley, P. Honarpisheh, D. A. Morales Scheihing, S. Singh, W. Dang, N. Kim, A. Urayama, L. Zhu, D. Monchaud, L. D. McCullough and A. S. Tsvetkov, *eLife*, 2020, **9**, e52283.

250 J. Tan, X. Wang, L. Phoon, H. Yang and L. Lan, *FEBS Lett.*, 2020, **594**, 1359–1367.

251 G. Beadle and L. Pauling, *Eng. Sci.*, 1954, **17**, 9.

252 K. H. Altmann, J. Buchner, H. Kessler, F. Diederich, B. Kräutler, S. J. Lippard, R. Liskamp, K. Müller, E. M. Nolan, B. Samori, G. Schneider, S. L. Schreiber, H. Schwalbe, C. Toniolo,



C. A. A. van Boeckel, H. Waldmann and C. T. Walsh, *ChemBioChem*, 2009, **10**, 16–29.

253 S. L. Schreiber, *Nat. Chem. Biol.*, 2005, **1**, 64–66.

254 Y. Li, C. R. Geyer and D. Sen, *Biochemistry*, 1996, **35**, 6911–6922.

255 Q. Chen, I. D. Kuntz and R. H. Shafer, *Proc. Natl. Acad. Sci. U. S. A.*, 1996, **93**, 2635–2639.

256 D. Sun, B. Thompson, B. E. Cathers, M. Salazar, S. M. Kerwin, J. O. Trent, T. C. Jenkins, S. Neidle and L. H. Hurley, *J. Med. Chem.*, 1997, **40**, 2113–2116.

257 S. Müller and R. Rodriguez, *Expert Rev. Clin. Pharmacol.*, 2014, **7**, 663–679.

258 F. Raguseo, S. Chowdhury, A. Minard and M. Di Antonio, *Chem. Commun.*, 2020, **56**, 1317–1324.

259 D. Monchaud, *Annu. Rep. Med. Chem.*, 2020, **54**, 133–160.

260 D. Monchaud and M. P. Teulade-Fichou, *Org. Biomol. Chem.*, 2008, **6**, 627–636.

261 S. A. Ohnmacht and S. Neidle, *Bioorg. Med. Chem. Lett.*, 2014, **24**, 2602–2612.

262 W. J. Chung, B. Heddi, M. Tera, K. Iida, K. Nagasawa and A. T. Phan, *J. Am. Chem. Soc.*, 2013, **135**, 13495–13501.

263 W. J. Chung, B. Heddi, F. Hamon, M. P. Teulade-Fichou and A. T. Phan, *Angew. Chem., Int. Ed.*, 2014, **53**, 999–1002.

264 A. T. Phan, V. Kuryavyi, H. Y. Gaw and D. J. Patel, *Nat. Chem. Biol.*, 2005, **1**, 167–173.

265 P. Murat, Y. Singh and E. Defrancq, *Chem. Soc. Rev.*, 2011, **40**, 5293–5307.

266 J. Jaumot and R. Gargallo, *Curr. Pharm. Des.*, 2012, **18**, 1900–1916.

267 K. M. Felsenstein, L. B. Saunders, J. K. Simmons, E. Leon, D. R. Calabrese, S. Zhang, A. Michalowski, P. Gareiss, B. A. Mock and J. S. Schneekloth, *ACS Chem. Biol.*, 2016, **11**, 138–148.

268 S. Ray, D. Tillo, R. E. Boer, N. Assad, M. Barshai, G. Wu, Y. Orenstein, D. Yang, J. S. Schneekloth, Jr. and C. Vinson, *ACS Chem. Biol.*, 2020, **15**, 925–935.

269 B. L. Allen-Petersen and R. C. Sears, *BioDrugs*, 2019, **33**, 539–553.

270 K. Shin-ya, K. Wierzba, K. Matsuo, T. Ohtani, Y. Yamada, K. Furihata, Y. Hayakawa and H. Seto, *J. Am. Chem. Soc.*, 2001, **123**, 1262–1263.

271 T. Tauchi, K. Shin-ya, G. Sashida, M. Sumi, A. Nakajima, T. Shimamoto, J. H. Ohyashiki and K. Ohyashiki, *Oncogene*, 2003, **22**, 5338–5347.

272 N. Temime-Smaali, L. Guittat, A. Sidibe, K. Shin-ya, C. Trentesaux and J. F. Riou, *PLoS One*, 2009, **4**, e6919.

273 T. Miyazaki, Y. Pan, K. Joshi, D. Purohit, B. Hu, H. Demir, S. Mazumder, S. Okabe, T. Yamori, M. Viapiano, K. Shin-ya, H. Seimiya and I. Nakano, *Clin. Cancer Res.*, 2012, **18**, 1268–1280.

274 D. Gomez, T. Wenner, B. Brassart, C. Douarre, M. F. O'Donohue, V. El Khoury, K. Shin-ya, H. Morjani, C. Trentesaux and J. F. Riou, *J. Biol. Chem.*, 2006, **281**, 38721–38729.

275 S. L. Palumbo, R. M. Memmott, D. J. Uribe, Y. Krotova-Khan, L. H. Hurley and S. W. Ebbinghaus, *Nucleic Acids Res.*, 2008, **36**, 1755–1769.

276 M. Tera, H. Ishizuka, M. Takagi, M. Suganuma, K. Shin-ya and K. Nagasawa, *Angew. Chem., Int. Ed.*, 2008, **47**, 5557–5560.

277 T. Nakamura, S. Okabe, H. Yoshida, K. Iida, Y. Ma, S. Sasaki, T. Yamori, K. Shin-ya, I. Nakano, K. Nagasawa and H. Seimiya, *Sci. Rep.*, 2017, **7**, 3605.

278 E. Gavathiotis, R. A. Heald, M. F. G. Stevens and M. S. Searle, *Angew. Chem., Int. Ed.*, 2001, **40**, 4749.

279 C. Leonetti, S. Amodei, C. D. Angelo, A. Rizzo, B. Benassi, A. Antonelli, R. Elli, M. F. G. Stevens, M. D. Incalci, G. Zupi and A. Biroccio, *Mol. Pharmacol.*, 2004, **66**, 1138–1146.

280 E. Salvati, C. Leonetti, A. Rizzo, M. Scarsella, M. Mottolese, R. Galati, I. Sperduti, M. F. G. Stevens, M. D'Incalci, M. Blasco, G. Chiorino, S. Bauwens, B. Horard, E. Gilson, A. Stoppacciaro, G. Zupi and A. Biroccio, *J. Clin. Invest.*, 2007, **117**, 3236–3247.

281 A. Rizzo, E. Salvati, M. Porru, C. D'Angelo, M. F. Stevens, M. D'Incalci, C. Leonetti, E. Gilson, G. Zupi and A. Biroccio, *Nucleic Acids Res.*, 2009, **37**, 5353–5364.

282 A. De Cian, G. Cristofari, P. Reichenbach, E. De Lemos, D. Monchaud, M. P. Teulade-Fichou, K. Shin-ya, L. Lacroix, J. Lingner and J. L. Mergny, *Proc. Natl. Acad. Sci. U. S. A.*, 2007, **104**, 17347–17352.

283 C. Granotier, G. Pennarun, L. Riou, F. Hoffschir, L. R. Gauthier, A. De Cian, D. Gomez, E. Mandine, J. F. Riou, J. L. Mergny, P. Mailliet, B. Dutrillaux and F. D. Boussin, *Nucleic Acids Res.*, 2005, **33**, 4182–4190.

284 G. Pennarun, C. Granotier, F. Hoffschir, E. Mandine, D. Biard, L. R. Gauthier and F. D. Boussin, *Nucleic Acids Res.*, 2008, **36**, 1741–1754.

285 G. Pennarun, F. Hoffschir, D. Revaud, C. Granotier, L. R. Gauthier, P. Mailliet, D. S. Biard and F. D. Boussin, *Nucleic Acids Res.*, 2010, **38**, 2955–2963.

286 L. R. Gauthier, C. Granotier, F. Hoffschir, O. Etienne, A. Ayouaz, C. Desmaze, P. Mailliet, D. S. Biard and F. D. Boussin, *Cell. Mol. Life Sci.*, 2012, **69**, 629–640.

287 A. De Cian, E. DeLemos, J. L. Mergny, M. P. Teulade-Fichou and D. Monchaud, *J. Am. Chem. Soc.*, 2007, **129**, 1856–1857.

288 J. Lopes, A. Piazza, R. Bermejo, B. Kriegsman, A. Colosio, M. P. Teulade-Fichou, M. Foiani and A. Nicolas, *EMBO J.*, 2011, **30**, 4033–4046.

289 J. Zimmer, E. M. C. Tacconi, C. Folio, S. Badie, M. Porru, K. Klare, M. Tumiati, E. Markkanen, S. Halder, A. Ryan, S. P. Jackson, K. Ramadan, S. G. Kuznetsov, A. Biroccio, J. E. Sale and M. Tarsounas, *Mol. Cell*, 2016, **61**, 449–460.

290 K. Zyner, D. S. Mulhearn, S. Adhikari, S. M. Cuesta, M. Di Antonio, N. Erard, G. J. Hannon, D. Tannahill and S. Balasubramanian, *eLife*, 2019, **8**, e46793.

291 R. Rodriguez, S. Müller, J. A. Yeoman, C. Trentesaux, J. F. Riou and S. Balasubramanian, *J. Am. Chem. Soc.*, 2008, **130**, 15758–15759.

292 D. Koirala, S. Dhakal, B. Ashbridge, Y. Sannohe, R. Rodriguez, H. Sugiyama, S. Balasubramanian and H. Mao, *Nat. Chem.*, 2011, **3**, 782–787.

293 M. Di Antonio, K. I. E. McLuckie and S. Balasubramanian, *J. Am. Chem. Soc.*, 2014, **136**, 5860–5863.



294 S. Müller, S. Kumari, R. Rodriguez and S. Balasubramanian, *Nat. Chem.*, 2010, **2**, 1095–1098.

295 T. Cañeque, S. Müller and R. Rodriguez, *Nat. Rev. Chem.*, 2018, **2**, 202–215.

296 L. C. Kim, L. Song and E. B. Haura, *Nat. Rev. Clin. Oncol.*, 2009, **6**, 587–595.

297 J. F. Moruno-Manchon, E. C. Koellhoffer, J. Gopakumar, S. Hambarde, N. Kim, L. D. McCullough and A. S. Tsvetkov, *Aging*, 2017, **9**, 1957–1970.

298 W. Duan, A. Rangan, H. Vankayalapati, M. Y. Kim, Q. Zeng, D. Sun, H. Han, O. Y. Fedoroff, D. Nishioka, S. Y. Rha, E. Izbicka, D. D. Von Hoff and L. H. Hurley, *Mol. Cancer Ther.*, 2001, **1**, 103–120.

299 D. Drygin, A. Siddiqui-Jain, S. O'Brien, M. Schwaebe, A. Lin, J. Bliesath, C. B. Ho, C. Proffitt, K. Trent, J. P. Whitten, J. K. C. Lim, D. Von Hoff, K. Anderes and W. G. Rice, *Cancer Res.*, 2009, **69**, 7653–7661.

300 M. Haddach, M. K. Schwaebe, J. Michaux, J. Nagasawa, S. E. O'Brien, J. P. Whitten, F. Pierre, P. Kerdoncuff, L. Darjania, R. Stansfield, D. Drygin, K. Anderes, C. Proffitt, J. Bliesath, A. Siddiqui-Jain, M. Omori, N. Huser, W. G. Rice and D. M. Ryckman, *ACS Med. Chem. Lett.*, 2012, **3**, 602–606.

301 D. Drygin, A. Lin, J. Bliesath, C. B. Ho, S. E. O'Brien, C. Proffitt, M. Omori, M. Haddach, M. K. Schwaebe, A. Siddiqui-Jain, N. Streiner, J. E. Quin, E. Sanij, M. J. Bywater, R. D. Hannan, D. Ryckman, K. Anderes and W. G. Rice, *Cancer Res.*, 2011, **71**, 1418–1430.

302 S. S. Negi and P. Brown, *Oncotarget*, 2015, **6**, 18094–18104.

303 E. Sanij, K. M. Hannan, J. Xuan, S. Yan, J. E. Ahern, A. S. Trigos, N. Brajanovski, J. Son, K. T. Chan, O. Kondrashova, E. Lieschke, M. J. Wakefield, D. Frank, S. Ellis, C. Cullinane, J. Kang, G. Poortinga, P. Nag, A. J. Deans, K. K. Khanna, L. Mileskin, G. A. McArthur, J. Soong, E. M. J. J. Berns, R. D. Hannan, C. L. Scott, K. E. Sheppard and R. B. Pearson, *Nat. Commun.*, 2020, **11**, 2641.

304 H. Xu, M. Di Antonio, S. McKinney, V. Mathew, B. Ho, N. J. O'Neil, N. Dos Santos, J. Silvester, V. Wei, J. Garcia, F. Kabeer, D. Lai, P. Soriano, J. Banáth, D. S. Chiu, D. Yap, D. D. Le, F. B. Ye, A. Zhang, K. Thu, J. Soong, S. C. Lin, A. H. C. Tsai, T. Osako, T. Algara, D. N. Saunders, J. Wong, J. Xian, M. B. Bally, J. D. Brenton, G. W. Brown, S. P. Shah, D. Cescon, T. W. Mak, C. Caldas, P. C. Stirling, P. Hieter, S. Balasubramanian and S. Aparicio, *Nat. Commun.*, 2017, **8**, 14432.

305 G. Biffi, D. Tannahill, J. McCafferty and S. Balasubramanian, *Nat. Chem.*, 2013, **5**, 182–186.

306 P. M. Bruno, M. Lu, K. A. Dennis, H. Inam, C. J. Moore, J. Sheehe, S. J. Elledge, M. T. Hemann and J. R. Pritchard, *Proc. Natl. Acad. Sci. U. S. A.*, 2020, **117**, 4053–4060.

307 J. Beauvarlet, P. Bensadoun, E. Darbo, G. Labrunie, B. Rousseau, E. Richard, I. Draskovic, A. Londono-Vallejo, J. W. Dupuy, R. N. Das, A. Guédin, G. Robert, F. Orange, S. Croce, V. Valesco, P. Soubeyran, K. M. Ryan, J. L. Mergny and M. Djavaheri-Mergny, *Nucleic Acids Res.*, 2019, **47**, 2739–2756.

308 W. J. Zhou, R. Deng, X. Y. Zhang, G. K. Feng, L. Q. Gu and X. F. Zhu, *Mol. Cancer Ther.*, 2009, **8**, 3203–3213.

309 A. De Magis, S. G. Manzo, M. Russo, J. Marinello, R. Morigi, O. Sordet and G. Capranico, *Proc. Natl. Acad. Sci. U. S. A.*, 2019, **116**, 816–825.

310 A. Thomas and Y. Pommier, *Clin. Cancer Res.*, 2019, **25**, 6581–6589.

311 B. L. Staker, M. D. Feese, M. Cushman, Y. Pommier, D. Zembower, L. Stewart and A. B. Burgin, *J. Med. Chem.*, 2005, **48**, 2336–2345.

312 A. Henderson, Y. Wu, Y. C. Huang, E. A. Chavez, J. Platt, F. B. Johnson, R. M. Brosh, D. Sen and P. Lansdorp, *Nucleic Acids Res.*, 2014, **42**, 860–869.

313 D. C. Odermatt, W. Ting, C. L. Id, S. Wild, S. K. J. Id, E. Rothenberg and K. G. Id, *PLoS Genet.*, 2020, **16**, e1008740.

314 L. A. Watson, L. A. Solomon, J. R. Li, Y. Jiang, M. Edwards, K. Shin-Ya, F. Beier and N. G. Bérubé, *J. Clin. Invest.*, 2013, **123**, 2049–2063.

315 K. I. E. McLuckie, M. Di Antonio, H. Zecchini, J. Xian, C. Caldas, B. Krippendorf, D. Tannahill, C. Lowe and S. Balasubramanian, *J. Am. Chem. Soc.*, 2013, **135**, 9640–9643.

316 S. Li, H. Lu, Z. Wang, Q. Hu, H. Wang, R. Xiang, T. Chiba and X. Wu, *iScience*, 2019, **16**, 63–78.

317 W. Feng, D. A. Simpson, J. Carvajal-Garcia, B. A. Price, R. J. Kumar, L. E. Mose, R. D. Wood, N. Rashid, J. E. Purvis, J. S. Parker, D. A. Ramsden and G. P. Gupta, *Nat. Commun.*, 2019, **10**, 4286.

318 M. Aggarwal, J. A. Sommers, R. H. Shoemaker and R. M. Brosh, *Proc. Natl. Acad. Sci. U. S. A.*, 2011, **108**, 1525–1530.

319 P. Phatak, J. C. Cookson, F. Dai, V. Smith, R. B. Gartenhaus, M. F. G. Stevens and A. M. Burger, *Br. J. Cancer*, 2007, **96**, 1223–1233.

320 C. Leonetti, M. Scarsella, G. Riggio, A. Rizzo, E. Salvati, M. D'Incalci, L. Staszewsky, R. Frapolli, M. F. Stevens, A. Stoppacciaro, M. Mottolese, B. Antoniani, E. Gilson, G. Zupi and A. Biroccio, *Clin. Cancer Res.*, 2008, **14**, 7284–7291.

321 E. Salvati, M. Scarsella, M. Porru, A. Rizzo, S. Iachettini, L. Tentori, G. Graziani, M. D'Incalci, M. F. G. Stevens, A. Orlandi, D. Passeri, E. Gilson, G. Zupi, C. Leonetti and A. Biroccio, *Oncogene*, 2010, **29**, 6280–6293.

322 A. Biroccio, M. Porru, A. Rizzo, E. Salvati, C. D'Angelo, A. Orlandi, D. Passeri, M. Franceschin, M. F. G. Stevens, E. Gilson, G. Beretta, G. Zupi, C. Pisano, F. Zunino and C. Leonetti, *Clin. Cancer Res.*, 2011, **17**, 2227–2236.

323 A. Rizzo, S. Iachettini, P. Zizza, C. Cingolani, M. Porru, S. Artuso, M. Stevens, M. Hummersone, A. Biroccio, E. Salvati and C. Leonetti, *J. Exp. Clin. Cancer Res.*, 2015, **33**, 81.

324 F. Berardinelli, M. Tanori, D. Muoio, M. Buccarelli, A. Di Masi, S. Leone, L. Ricci-Vitiani, R. Pallini, M. Mancuso and A. Antoccia, *J. Exp. Clin. Cancer Res.*, 2019, **38**, 311.

325 F. Berardinelli, S. Siteni, C. Tanzarella, M. F. Stevens, A. Sgura and A. Antoccia, *DNA Repair*, 2015, **25**, 104–115.



326 Y. Wang, J. Yang, A. T. Wild, W. H. Wu, R. Shah, C. Danussi, G. J. Riggins, K. Kannan, E. P. Sulman, T. A. Chan and J. T. Huse, *Nat. Commun.*, 2019, **10**, 943.

327 M. Porru, P. Zizza, M. Franceschin, C. Leonetti and A. Biroccio, *Biochim. Biophys. Acta*, 2017, **1861**, 1362–1370.

328 T. C. Chou and P. Talalay, *Adv. Enzyme Regul.*, 1984, **22**, 27–55.

329 T. C. Chou, *Cancer Res.*, 2010, **70**, 440–446.

330 C. I. Bliss, *Ann. Appl. Biol.*, 1939, **26**, 585–615.

331 R. Rodriguez and K. M. Miller, *Nat. Rev. Genet.*, 2014, **15**, 783–796.

332 S. F. Cai, C. W. Chen and S. A. Armstrong, *Mol. Cell*, 2015, **60**, 561–570.

333 Y. Cheng, C. He, M. Wang, X. Ma, F. Mo, S. Yang, J. Han and X. Wei, *Signal Transduction Targeted Ther.*, 2019, **4**, 62.

334 D. S. Dizon, L. Damstrup, N. J. Finkler, U. Lassen, P. Celano, R. Glasspool, E. Crowley, H. S. Lichenstein, P. Knoblach and R. T. Penson, *Int. J. Gynecol. Cancer*, 2012, **22**, 979–986.

335 G. Wang and K. M. Vasquez, *DNA Repair*, 2014, **19**, 143–151.

336 T. Kato, H. Kurahashi and B. S. Emanuel, *Curr. Opin. Genet. Dev.*, 2012, **22**, 221–228.

337 T. J. Treangen and S. L. Salzberg, *Nat. Rev. Genet.*, 2012, **13**, 36–46.

338 J. Choi and T. Majima, *Chem. Soc. Rev.*, 2011, **40**, 5893–5909.

339 S. Kaushal and C. H. Freudenreich, *Genes, Chromosomes Cancer*, 2018, **58**, 270–283.

340 M. L. Bochman, K. Paeschke and V. A. Zakian, *Nat. Rev. Genet.*, 2012, **13**, 770–780.

341 S. Ravichandran, V. K. Subramani and K. K. Kim, *Biophys. Rev.*, 2019, **11**, 383–387.

342 A. Aguilera and B. Gómez-González, *Nat. Struct. Mol. Biol.*, 2017, **24**, 439–443.

343 F. Chedin, C. J. Benham and K. Musier-Forsyth, *J. Biol. Chem.*, 2020, **295**, 4684–4695.

344 B. Gómez-González and A. Aguilera, *Genes Dev.*, 2019, **33**, 1008–1026.

345 C. Niehrs and B. Luke, *Nat. Rev. Mol. Cell Biol.*, 2020, **21**, 167–178.

346 M. Duca, P. Vekhoff, K. Oussedik, L. Halby and P. B. Arimondo, *Nucleic Acids Res.*, 2008, **36**, 5123–5138.

347 M. Thomas, R. L. White and R. W. Davis, *Proc. Natl. Acad. Sci. U. S. A.*, 1976, **73**, 2294–2298.

348 M. P. Crossley, M. Bocek and K. A. Cimprich, *Mol. Cell*, 2019, **73**, 398–411.

349 L. A. Sanz and F. Chédin, *Nat. Protoc.*, 2019, **14**, 1734–1755.

350 P. A. Ginno, P. L. Lott, H. C. Christensen, I. Korf and F. Chédin, *Mol. Cell*, 2012, **45**, 814–825.

351 K. J. Abraham, N. Khosraviani, J. N. Y. Chan, A. Gorthi, A. Samman, D. Y. Zhao, M. Wang, M. Bokros, E. Vidya, L. A. Ostrowski, R. Oshidari, V. Pietrobon, P. S. Patel, A. Algouneh, R. Singhania, Y. Liu, V. T. Yerlici, D. D. De Carvalho, M. Ohh, B. C. Dickson, R. Hakem, J. F. Greenblatt, S. Lee, A. J. R. Bishop and K. Mekhail, *Nature*, 2020, **585**, 298–302.

352 M. R. Lieber, *Annu. Rev. Biochem.*, 2010, **79**, 181–211.

353 S. Britton, E. Dernoncourt, C. Delteil, C. Froment, O. Schiltz, B. Salles, P. Frit and P. Calsou, *Nucleic Acids Res.*, 2014, **42**, 9047–9062.

354 S. Tuduri, L. Crabbé, C. Conti, H. Tourrière, H. Holtgreve-Grez, A. Jauch, V. Pantesco, J. De Vos, A. Thomas, C. Theillet, Y. Pommier, J. Tazi, A. Coquelle and P. Pasero, *Nat. Cell Biol.*, 2009, **11**, 1315–1324.

355 J. Tazi, N. Balkour, J. Soret, L. Zekri, B. Hazra, W. Laine, B. Baldeyrou, A. Lansiaux and C. Bailly, *Mol. Pharmacol.*, 2005, **67**, 1186–1194.

356 A. De Magis, S. G. Manzo, M. Russo, J. Marinello, R. Morigi, O. Sordet and G. Capranico, *Proc. Natl. Acad. Sci. U. S. A.*, 2019, **116**, 816–825.

357 J. Amato, G. Miglietta, R. Morigi, N. Iaccarino, A. Locatelli, A. Leoni, E. Novellino, B. Pagano, G. Capranico and A. Randazzo, *J. Med. Chem.*, 2020, **63**, 3090–3103.

358 D. M. J. Lilley, *Q. Rev. Biophys.*, 2000, **33**, 109–159.

359 V. Brázda, J. Kolomazník, J. Lýsek, L. Hároníková, J. Coufal and J. Štastný, *Biochem. Biophys. Res. Commun.*, 2016, **478**, 1739–1745.

360 E. M. Strawbridge, G. Benson, Y. Gelfand and C. J. Benham, *Curr. Genet.*, 2010, **56**, 321–340.

361 M. Čutová, J. Manta, O. Porubiaková, P. Kaura, J. Štastný, E. B. Jagelská, P. Goswami, M. Bartas and V. Brázda, *Genomics*, 2020, **112**, 1897–1901.

362 A. Bedrat, L. Lacroix and J. L. Mergny, *Nucleic Acids Res.*, 2016, **44**, 1746–1759.

363 R. Cox and S. M. Mirkin, *Proc. Natl. Acad. Sci. U. S. A.*, 1997, **94**, 5237–5242.

364 X. Lu, G. Wang, A. Bacolla, J. Zhao, S. Spitser and K. M. Vasquez, *Cell Rep.*, 2015, **10**, 1674–1680.

365 Y. Wang and F. C. C. Leung, *FEBS J.*, 2009, **276**, 1986–1998.

366 D. G. Albertson, *Trends Genet.*, 2006, **22**, 447–455.

367 A. Aguilera and T. García-Muse, *Annu. Rev. Genet.*, 2013, **47**, 1–32.

368 A. B. Reams and J. R. Roth, *Cold Spring Harbor Perspect. Biol.*, 2015, **7**, a016592.

369 C. T. McMurray, *Nat. Rev. Genet.*, 2010, **11**, 786–799.

370 W. K. Chu and I. D. Hickson, *Nat. Rev. Cancer*, 2009, **9**, 644–654.

371 I. Voineagu, V. Narayanan, K. S. Lobachev and S. M. Mirkin, *Proc. Natl. Acad. Sci. U. S. A.*, 2008, **105**, 9936–9941.

372 I. Voineagu, C. F. Surka, A. A. Shishkin, M. M. Krasilnikova and S. M. Mirkin, *Nat. Struct. Mol. Biol.*, 2009, **16**, 226–228.

373 N. Shastri, Y. C. Tsai, S. Hile, D. Jordan, B. Powell, J. Chen, D. Maloney, M. Dose, Y. Lo, T. Anastassiadis, O. Rivera, T. Kim, S. Shah, P. Borole, K. Asija, X. Wang, K. D. Smith, D. Finn, J. Schug, R. Casellas, L. A. Yatsunyk, K. A. Eckert and E. J. Brown, *Mol. Cell*, 2018, **72**, 222–238.

374 L. Gellon, S. Kaushal, J. Cebrián, M. Lahiri, S. M. Mirkin and C. H. Freudenreich, *Nucleic Acids Res.*, 2019, **47**, 794–805.

375 M. L. Nicolette, K. Lee, Z. Guo, M. Rani, J. M. Chow, S. E. Lee and T. T. Paull, *Nat. Struct. Mol. Biol.*, 2010, **17**, 1478–1485.

376 K. S. Lobachev, D. A. Gordenin and M. A. Resnick, *Cell*, 2002, **108**, 183–193.

377 M. McVey and S. E. Lee, *Trends Genet.*, 2008, **24**, 529–538.

378 A. L. Brogden, N. H. Hopcroft, M. Searcey and C. J. Cardin, *Angew. Chem., Int. Ed.*, 2007, **46**, 3850–3854.



379 A. Oleksi, A. G. Blanco, R. Boer, I. Usón, J. Aymamí, A. Rodger, M. J. Hannon and M. Coll, *Angew. Chem., Int. Ed.*, 2006, **45**, 1227–1231.

380 Q. Guo, N. C. Seeman and N. R. Kallenbach, *Biochemistry*, 1989, **28**, 2355–2359.

381 M. Lu, Q. Guo and N. R. Kallenbach, *Crit. Rev. Biochem. Mol. Biol.*, 1992, **27**, 157–190.

382 M. Lu, Q. Guo, R. F. Pasternack, D. J. Wink, N. C. Seeman and N. R. Kallenbach, *Biochemistry*, 1990, **29**, 1614–1624.

383 M. N. Stojanovic, P. De Prada and D. W. Landry, *J. Am. Chem. Soc.*, 2000, **122**, 11547–11548.

384 M. N. Stojanovic, P. de Prada and D. W. Landry, *J. Am. Chem. Soc.*, 2001, **123**, 4928–4931.

385 M. N. Stojanovic, E. G. Green, S. Semova, D. B. Nikić and D. W. Landry, *J. Am. Chem. Soc.*, 2003, **125**, 6085–6089.

386 M. N. Stojanovic and T. S. Worgall, *Curr. Opin. Chem. Biol.*, 2010, **14**, 751–757.

387 M. Lu, Q. Guo, N. C. Seeman and N. R. Kallenbach, *Biochemistry*, 1990, **29**, 3407–3412.

388 K. V. Kepple, J. L. Boldt and A. M. Segall, *Proc. Natl. Acad. Sci. U. S. A.*, 2005, **102**, 6867–6872.

389 K. V. Kepple, N. Patel, P. Salamon and A. M. Segall, *Nucleic Acids Res.*, 2008, **36**, 5319–5334.

390 M. L. Bolla, E. V. Azevedo, J. M. Smith, R. E. Taylor, D. K. Ranjit, A. M. Segall and S. R. McAlpine, *Org. Lett.*, 2003, **5**, 109–112.

391 P. S. Pan, F. A. Curtis, C. L. Carroll, I. Medina, L. A. Liotta, G. J. Sharples and S. R. McAlpine, *Bioorg. Med. Chem.*, 2006, **14**, 4731–4739.

392 L. A. Howell, R. A. Bowater, M. A. O'Connell, A. P. Reszka, S. Neidle and M. Searcey, *ChemMedChem*, 2012, **7**, 792–804.

393 L. A. Howell, Z. A. E. Waller, R. Bowater, M. A. O'Connell and M. Searcey, *Chem. Commun.*, 2011, **47**, 8262–8264.

394 D. R. Boer, J. M. C. A. Kerckhoffs, Y. Parajo, M. Pascu, I. Usón, P. Lincoln, M. J. Hannon and M. Coll, *Angew. Chem., Int. Ed.*, 2010, **49**, 2336–2339.

395 I. Gamba, G. Rama, E. Ortega-Carrasco, J. D. Maréchal, J. Martínez-Costas, M. Eugenio Vázquez and M. V. López, *Chem. Commun.*, 2014, **50**, 11097–11100.

396 S. Vuong, L. Stefan, P. Lejault, Y. Roussel, F. Denat and D. Monchaud, *Biochimie*, 2012, **94**, 442–450.

397 L. Stefan, B. Bertrand, P. Richard, P. Le Gendre, F. Denat, M. Picquet and D. Monchaud, *ChemBioChem*, 2012, **13**, 1905–1912.

398 S. A. Barros and D. M. Chenoweth, *Angew. Chem., Int. Ed.*, 2014, **53**, 13746–13750.

399 S. A. Barros and D. M. Chenoweth, *Chem. Sci.*, 2015, **6**, 4752–4755.

400 J. Novotna, A. Laguerre, A. Granzhan, M. Pirrotta, M. P. Teulade-Fichou and D. Monchaud, *Org. Biomol. Chem.*, 2015, **13**, 215–222.

401 L. Guyon, M. Pirrotta, K. Duskova, A. Granzhan, M. P. Teulade-Fichou and D. Monchaud, *Nucleic Acids Res.*, 2018, **46**, e16.

402 K. Duskova, J. Lamarche, S. Amor, C. Caron, N. Queyriaux, M. Gaschard, M. J. Penouilh, G. De Robillard, D. Delmas, C. H. Devillers, A. Granzhan, M. P. Teulade-Fichou, M. Chavarot-Kerlidou, B. Therrien, S. Britton and D. Monchaud, *J. Med. Chem.*, 2019, **62**, 4456–4466.

403 J. Zhu, C. J. E. Haynes, M. Kieffer, J. L. Greenfield, R. D. Greenhalgh, J. R. Nitschke and U. F. Keyser, *J. Am. Chem. Soc.*, 2019, **141**, 11358–11362.

404 Z. Yang, Y. Chen, G. Li, Z. Tian, L. Zhao, X. Wu, Q. Ma, M. Liu and P. Yang, *Chem. – Eur. J.*, 2018, **24**, 6087–6093.

405 J. Xia, Q. Mei and S. M. Rosenberg, *Trends Genet.*, 2019, **35**, 383–395.

406 J. Xia, L. T. Chen, Q. Mei, C. H. Ma, J. A. Halliday, H. Y. Lin, D. Magnan, J. P. Pribis, D. M. Fitzgerald, H. M. Hamilton, M. Richters, R. B. Nehring, X. Shen, L. Li, D. Bates, P. J. Hastings, C. Herman, M. Jayaram and S. M. Rosenberg, *Sci. Adv.*, 2016, **2**, e1601605.

407 K. Ghosh, K. L. Chi, F. Guo, A. M. Segall and G. D. Van Duyne, *J. Biol. Chem.*, 2005, **280**, 8290–8299.

408 C. W. Gunderson and A. M. Segall, *Mol. Microbiol.*, 2006, **59**, 1129–1148.

409 M. Dey, S. Patra, L. Y. Su and A. M. Segall, *PLoS One*, 2013, **8**, e78751.

410 A. C. G. Hotze, N. J. Hodges, R. E. Hayden, C. Sanchez-Cano, C. Paines, N. Male, M. K. Tse, C. M. Bunce, J. K. Chipman and M. J. Hannon, *Chem. Biol.*, 2008, **15**, 1258–1267.

411 C. Ducani, A. Leczkowska, N. J. Hodges and M. J. Hannon, *Angew. Chem., Int. Ed.*, 2010, **49**, 8942–8945.

412 G. I. Pascu, A. C. G. Hotze, C. Sanchez-Cano, B. M. Kariuki and M. J. Hannon, *Angew. Chem., Int. Ed.*, 2007, **46**, 4374–4378.

413 U. McDonnell, J. M. C. A. Kerchoffs, R. P. M. Castineiras, M. R. Hicks, A. C. G. Hotze, M. J. Hannon and A. Rodger, *Dalton Trans.*, 2008, 667–675.

414 H. Yu, X. Wang, M. Fu, J. Ren and X. Qu, *Nucleic Acids Res.*, 2008, **36**, 5695–5703.

415 C. Zhao, J. Geng, L. Feng, J. Ren and X. Qu, *Chem. – Eur. J.*, 2011, **17**, 8209–8215.

416 C. Zhao, L. Wu, J. Ren, Y. Xu and X. Qu, *J. Am. Chem. Soc.*, 2013, **135**, 18786–18789.

417 V. Brabec, S. E. Howson, R. A. Kaner, R. M. Lord, J. Malina, R. M. Phillips, Q. M. A. Abdallah, P. C. McGowan, A. Rodger and P. Scott, *Chem. Sci.*, 2013, **4**, 4407–4416.

418 A. Granzhan, N. Kotera and M. P. Teulade-Fichou, *Chem. Soc. Rev.*, 2014, **43**, 3630–3665.

419 N. Kotera, A. Granzhan and M. P. Teulade-Fichou, *Biochimie*, 2016, **128–129**, 133–137.

420 N. Kotera, F. Poyer, A. Granzhan and M. P. Teulade-Fichou, *Chem. Commun.*, 2015, **51**, 15948–15951.

421 C. Caron, X. N. T. Duong, R. Guillot, S. Bombard and A. Granzhan, *Chem. – Eur. J.*, 2019, **25**, 1949–1962.

422 K. Duskova, P. Lejault, É. Benchimol, R. Guillot, S. Britton, A. Granzhan and D. Monchaud, *J. Am. Chem. Soc.*, 2020, **142**, 424–435.

423 F. H. Crick, *Symp. Soc. Exp. Biol.*, 1958, **12**, 138–163.

424 F. Crick, *Nature*, 1970, **227**, 561–563.

425 F. Crick, *Nature*, 1974, **248**, 766–769.

