DNA-Protein Crosslink Proteolysis Repair

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Abstract

Proteins that are covalently bound to DNA constitute a specific type of DNA lesions known as DNA-protein crosslinks (DPCs). DPCs represent physical obstacles to the progression of DNA replication. If not repaired, DPCs cause stalling of DNA replication forks that consequently leads to DNA double strand breaks, the most cytotoxic DNA lesion. Although DPCs are abundant DNA lesions, the mechanism of DPC repair was unclear until now. Recent work unveiled that DPC repair is orchestrated by proteolysis performed by two distinct metalloproteases, SPARTAN in metazoans and Wss1 in yeast. This review summarises recent discoveries on two proteases in DNA replication-coupled DPC repair and establishes DPC proteolysis repair as a separate DNA repair pathway for genome stability and protection from accelerated ageing and cancer.
Overview of DNA-protein crosslinks

DNA contains all genetic information a cell and organism requires to grow, differentiate, survive and divide. DNA is an extremely fragile molecule embedded in highly reactive environment such as H2O. DNA is constantly damaged by various endogenous and exogenous agents, such as reactive oxygen species and UV light, respectively. Loss of DNA integrity leads to various defects in cellular physiology and consequently to diseases such as cancer, diabetes, accelerated aging, neurodegeneration or cell death. To preserve DNA integrity all organisms possess an elaborate genome maintenance apparatus, consisting of multiple DNA damage repair (DDR) and DNA damage tolerance (DDT) pathways (see Glossary) (Figure 1). Different DDR pathways are involved in the recognition and repair of specific types of DNA lesions [1, 2]. Currently, most of the known DDR pathways are well-characterised. However, it is still not well understood how DNA-protein crosslinks are repaired [3]. Although DPCs are one of the most abundant DNA lesions and their presence, if not removed, is cytotoxic, the existence of a specialised DPC repair pathway remained elusive. Recently, several research groups identified a unique DNA-protein crosslink repair pathway based on proteolysis, which we have termed here as DNA-protein crosslink proteolysis repair (DPC-PR) [4-8]. DPC-PR pathway is conserved from yeast to humans and is orchestrated by DNA-dependent proteases Wss1 in yeast and SPARTAN (SPRTN), also known as DVC1, in metazoans. The aim of this review is to establish DPC-PR as a unique DNA repair pathway based on DNA replication-coupled proteolysis orchestrated by SPRTN protease in metazoans and Wss1 in yeast.
**Origins and chemistry of DNA-protein crosslinks**

DPCs are created when proteins covalently and irreversibly bind to DNA. Virtually any protein in close proximity to DNA can be crosslinked to DNA upon exposure to various endogenous or exogenous crosslinking agents. Mass-spectrometry analyses identified numerous DNA binding proteins including histones, transcription factors, DNA repair and replication proteins, as well as non-DNA binding proteins as DPCs [4, 9-11]. DPC classification is explained in Box 1. Given the high concentration of histones in close proximity to DNA, it is not surprising that histones are among the most abundant DPCs [12]. Aldehydes, reactive oxygen (ROS), nitrogen species (NOS) and DNA helical alterations are one of the most common endogenous DPC-inducing sources. Aldehydes are generated by normal cellular metabolic pathways including histone demethylation [13], AlkB-type repair [14] amino acid metabolism [15] and lipid peroxidation [16]. It is estimated that endogenous formaldehyde concentration in human plasma is 100 µM, and in some tissues can reach up to 400 µM [17, 18]. Similarly, ROS and NOS are metabolic side products of cellular respiration, immune responses and inflammation [19]. While aldehydes, ROS and NOS crosslink any protein in the vicinity of DNA, DNA helical alterations such as DNA abasic sites and oxanines (nitric oxide induced guanine lesions) are so far shown to crosslink histones [20], DNA glycosylases, DNA polymerase β [21] and high mobility group (HMG) proteins [22].

Cancer treatment including Ionizing Radiation (IR) and anticancer drugs as well as exposure to UV-light are the main exogenous DPC-inducing factors. IR causes DPCs directly, through the irradiation of the DNA backbone which creates unstable DNA cation radicals or indirectly, through the transfer of radiation energy to water molecules which in turn enhances local concentrations of ROS. Anticancer drugs like
mitomycin C, platinum compounds, nitrogen mustard, DNA methyltransferase inhibitors [23] and Topoisomerase poisons (camptothecin and etoposide) [24, 25] cause DPCs through agent-specific mechanisms. UV-light excites DNA bases, most commonly thymidines, which in turn covalently bind to amino acids, specifically cysteine, lysine, phenylalanine, tryptophan or tyrosine with highest efficiency [26]. For the detailed chemistry of DPCs see Box 2. Altogether, DPCs are constantly formed in our genome and if not repaired cause severe threat to genome integrity.

**Involvement of canonical DNA repair pathways in DPC repair**

Although DPCs are abundant DNA lesions, the mechanisms of DPC repair were under-investigated and thus poorly understood [12]. A few biochemical and genetic studies in different organisms suggested that two canonical DNA repair pathways, namely nucleotide excision repair (NER) and homologous recombination (HR), orchestrate DPC repair and protect cells from DPC-induced cytotoxicity (Figure 2A) [27]. This concept came from the initial studies in bacteria where NER was found to remove and repair small DPCs (smaller than 16kDa) [28], while HR and subsequent replication restart repaired bulky DPCs [29] [12]. A genome-wide screen in yeast implicated NER in the repair of DPCs after acute exposure to high formaldehyde doses and HR in the repair of DPCs after chronic exposure to low formaldehyde doses [30]. Sensitivity analysis in mutant yeast strains also showed that NER is dominant in the DPC repair after high formaldehyde doses [5]. The coordination of NER and HR in yeast is probably dependent on cell cycle phase (high formaldehyde doses cause cell cycle arrest and thus favour NER), and the size of the DPC, similar as in bacteria, although this has not been shown so far. The analysis of mammalian NER excision capacity in vitro and in vivo showed that NER is only able to remove DNA-
crosslinked proteins of less than 8 - 10 kDa [28, 29, 31-33]. Correspondingly, cells from *Xeroderma pigmentosum* patients (bearing mutations in different NER factors) are sensitive to different DPC-inducing agents [34, 35].

Unlike NER, the involvement of HR in DPC repair has only been shown indirectly in bacteria and yeast, and more recently in metazoans. Thus, it is still difficult to understand whether HR is directly involved in DPC repair. HR-deficient *E. coli* cells (*ΔrecA ΔrecB*) were sensitive to DPC-inducing agents, formaldehyde and azacytidine [29, 36]. In yeast, deletion of the HR genes *sgs1, xrs2, mre11, rad50* and/or *rad52* sensitised cells to chronic doses of formaldehyde [5, 30]. The role of HR in DPC repair was further supported by the observation that formaldehyde-treated cells showed elevated levels of DSBs and Rad51 foci and an increased rate of sister chromatid exchange (SCE) events [37]. However, involvement of HR is not surprising given that one of the main outcomes of DPC accumulation is the emergence of DSBs [4, 38]. As formaldehyde not only induces DPCs, but also protein-protein crosslinks and, more importantly DNA inter- and intra-strand crosslinks (ICL), it activates the Fanconi anemia pathway as well as HR. Thus, all aforementioned experimental endpoints such as cell survival, SCE, and DSB formation measure the total cellular response to the various types of formaldehyde-induced DNA damage, and not DPCs alone. Therefore, it is hard to conclude to what extent HR is involved, if at all, in DPC repair.

Interestingly, recent work demonstrates that depletion of Mre11, a crucial nuclease in HR, does not impair general DPC removal in human cells [4], thus supporting a hypothesis that HR cannot deal with the majority of diverse DPCs. However, two recent studies in human cells have demonstrated that MRE11, independently of its
function in HR and together with another nuclease, CtIP, removes TOP2-ccs. MRE11 does not directly act on TOP2-ccs, but endonucleolytically cleaves DNA 15-20 bp downstream of TOP2-cc [39, 40]. Thus, by removing TOP2-ccs from the 5’ end of DSBs, MRE11 removes DPCs after DSB formation. Similarly, biochemical data in *Xenopus* egg extract demonstrated that TOP2-cc complex is processed by cooperation of MRN (MRE11-RAD50-NBS1) complex, BRCA1 and CtIP [41]. Altogether, these data suggest that MRN complex removes covalently attached TOP2 from DSBs in both HR-dependent and HR-independent manner. However, further studies are needed to clarify a direct role of HR in DPC repair.

Considering that NER can act only on small DPCs, and HR removes only DPCs after DSB formation, it was speculated that the Fanconi Anemia DNA repair pathway might be involved in DPC repair. Fanconi anemia pathway-deficient cells were found to be hypersensitive to formaldehyde and azacytidine but not to the specific TOP1-cc-inducing agent camptothecin [4, 42, 43]. However, similar to HR- and NER-deficient cells, cells deficient of FANCD2, one of the main component of the Fanconi Anemia pathway, do not accumulate DPCs and exhibit normal DPC repair kinetics following treatment with formaldehyde [4, 6]. Furthermore, immunodepletion of FANCD2 from *Xenopus* egg extracts did not affect DNA replication fork progression past DPCs [44]. Therefore, the role of the Fanconi Anemia pathway in DPC repair seems to be strictly associated with the repair of DNA interstrand crosslinks (ICLs), a lesion also induced by formaldehyde, and not with DPC repair.

In conclusion, in eukaryotes canonical DNA repair pathways such as NER and HR remove DPCs via the activity of nucleases that cleave DNA near to where a DPC is formed. While the role of NER in the repair of small DPCs is beneficial to cells,
role of HR in removing bulky DPCs requires DSB formation. Thus, activating HR-pathway for DPC repair may be deleterious for cells due the potentially cytotoxic consequences of DSBs. However, it is quite clear that the Fanconi Anemia pathway is not involved in DPC repair pathway.

**Proteolysis Orchestrated DPC repair**

Due to diverse array of DPCs (various proteins attached with different chemistry to DNA) contrasted with the known specificity of the canonical DNA repair pathways postulated to be involved in DPC removal (see above). Thus, it was speculated that cells must contain a specialised DNA repair pathway that is based on direct removal (proteolysis) of DPCs. NER can only excise small DPCs (max size of 8-16 kDa) suggesting that bulky (bigger than 16 kDa) DPCs need to be processed into smaller peptides before the action of NER. Similarly, removal of enzymatic DPCs, TOP1-ccs and TOP2-ccs by Tyrosyl-DNA phosphodiesterase 1 (TDP1) and 2 (TDP2), respectively, requires upstream proteolysis of TOP1 and 2 into smaller peptides [45] as TDP1 and 2 can efficiently process peptides of ~150 amino acid long [46-48] (Box3). Altogether, these data suggest that it must exists a protease that proteolysis large DPCs to small peptide remnants attached to the DNA backbone. These peptide remnants are further processed by NER, TDP1 and TDP2 or bypassed by translesion DNA synthesis (TLS) during DNA synthesis. Without such a protease, bulky DPCs would block the progression of the DNA replication fork and lead to DSBs in proliferative cells. Thus, proteolysis-coupled DPC repair was proposed [49].

**Proteasome in DPC repair**

The proteasome, being the main protease involved in protein degradation, was
considered to be involved in proteolytic DPC repair. However, the role of the proteasome system in DPC processing remains unclear due to contradictory literature reports. In bacteria, inhibition of ATP-dependent proteases, which function like a proteasome, did not affect cell survival after exposure to DPC inducing agents formaldehyde and azacytidine [29]. In human cells, proteasome inhibition prevented the removal of histone DPCs, TOP1ccs, and TOP2ccs [50-52] and sensitised human cells to low doses of formaldehyde [53]. By contrast, in *Xenopus* egg extracts, DPC proteolysis and bypass during DNA replication was inhibited upon depletion of the free pool of ubiquitin, but not by proteasome inhibition [44]. These discrepancies could be explained by the fact that proteasome inhibitors MG132, lactacystin, and bortezemib used in the aforementioned studies can also deplete the nuclear ubiquitin pool in human cells, thus making it hard to conclude whether proteasome is directly involved in DPC repair [54]. Altogether, the data suggest that DPC proteolysis is an ubiquitin-dependent process but a direct role of proteasome in DPC repair is still not clear. A precise understanding of the role of proteasome in DPC repair requires additional studies and different approaches including usage of specific proteasome inhibitors and *in vitro* studies.

**DNA-dependent Proteases in DPC repair**

The proteolysis-dependent model of DPC repair [49] was further supported by several studies, which all demonstrated replication-coupled proteolysis of a specific DPC *in vitro* [44, 55, 56]. However, the protease in question remained unknown. This model was further supported by the discovery of Wss1, a protease in yeast found to cleave TOP1, histone H1 and HMG proteins *in vitro* and contribute to the cellular resistance to formaldehyde and camptothecin [5]. Four recent studies identified a new
mammalian protease, SPARTAN (SPRTN), also known as DNA damage valosin containing factor 1 (DVC1), as a crucial component of DPC repair in human cells [4, 6-8]. Interestingly, even though bacterial species possess SPRT-like proteins (Figure 3A), proteolysis-dependent DPC repair in bacteria has not yet been reported. Like SPRTN [4], yeast protease Wss1 [57] binds zinc and possesses an HEXXH active site, thus making both of these enzymes members of the zinzin family of metallopeptidases [4]. SPRTN is a pleiotropic DNA-dependent protease that cleaves several chromatin-associated substrates, including core histones, H2A, H2B, H3, H4, linker histone H1, HMG1, HLTF, Fan1, TOP1 and TOP2 [4, 6-8]. Both proteases, Wss1 and SPRTN, are linked to DNA replication and share some common characteristics [4, 8, 58-60]. Wss1 and SPRTN need DNA to activate their proteolytic activity, cleave DNA binding proteins, physically interact with AAA ATPase p97/VCP, the central component of the ubiquitin-proteasome system, and inactivation of Wss1 and SPRTN hypersensitises yeast and human cells to formaldehyde, respectively.

Although Wss1 and SPRTN share similar proteolytic activity in vitro, in vivo they show considerable differences in DPC removal and sensitivity to DPC-inducing agents. SPRTN alone prevents accumulation of endogenous DPCs, as well as formaldehyde-induced DPCs [4, 6]. Concordantly, SPRTN protects cells from formaldehyde-induced DPC toxicity [4, 6, 7]. On the contrary, Wss1 is not involved in the removal of DPCs following formaldehyde treatment, but does partially protect cells from formaldehyde-induced DPC toxicity [5]. Another difference between Wss1 and SPRTN is observed in the repair of TOP1-ccs. SPRTN deficiency results in Top1-cc accumulation and severe sensitivity to CPT [4], while Wss1 depletion does not cause any adverse effects in untreated yeast cells [5]. Only upon co-depletion of
Wss1 with Tdp1 is cell survival affected, while in mammals, TDP1 and SPRTN co-depletion has not an additive effect in comparison to SPRTN depletion alone [4]. The in vivo differences between the two proteases are further demonstrated by the inability of ectopic SPRTN over-expression to rescue the phenotypes of Wss1-deficient yeast cells [7]. However, despite numerous functional differences, both proteases are essential for replication fork progression [4, 8, 58, 61], indicating that Wss1, like SPRTN, removes DPC blocks during replication. Unlike SPRTN in mammals [62-64], Wss1 is not an essential gene in yeast [65], indicating that the function of Wss1 can be compensated. Differences in the phenotypes observed in Wss1- and SPRTN-deficient cells could be due to the existence of other proteases that process DPCs in yeast. Considering the high toxicity of DPCs and the high growth rate of yeast cells, it would not be surprising if yeasts possess other DNA dependent-proteases. Accordingly, in Schizosaccharomyces pombe a Wss2 protease was found to protect cells from acetaldehyde [66].

SPRTN and Wss1 are Two Distinct Proteases

Published literature indicates that SPRTN and WSS1 diverged through evolution from a common ancestor [67]. However, the functional and cellular differences between Wss1 and SPRTN prompted us to phylogenetically analyse the zinzin metallopeptidase superfamily. Phylogenetic analysis suggests that Wss1 and SPRTN are members of two separate families (Figure 3A). The SPRT family, where SPRTN belongs, consists of five subgroups: bacterial, archaea, cyanobacterial, plant and animal. The prokaryote and animal SPRT families map to the same branch of the phylogenetic tree suggesting they share a common ancestor. WLM family, where Wss1 belongs, is equally distant from SPRT family as it is from another gluzincin
family of alanyl aminopeptidases (Figure 3A). This analysis indicates that WLM and SPRT families do not share a common ancestor as was previously suggested [67]. The differences in our methodology compared to previous phylogenetic studies comparing SPRT and WLM families are: (i) an extended number of species were included in the analysis, most importantly prokaryotes (bacteria, archaea, cyanobacteria) and plants which increases the accuracy of tree topologies; and (ii) our analysis included another gluzincin family which enables comparative perspective to the relationship between SPRT and WLM families (Figures S1, S2 and Table S1). Moreover, the SPRT family is present in bacteria, archaea, cyanobacteria, funghi, plants and animals, but is absent from yeast (Figure 3A, Table S1). Wss1 is part of the WLM protein family which, like the SPRT family, consists of zinzin metalloproteases [68]. WLM proteins are present in yeast, funghi and plants, but are absent from animals and bacteria (Fig 3A) [12, 68]. A conserved feature of all zinzin metalloproteases is a short consensus HEXXH motif in their active centres, which includes two zinc-binding histidines and a glutamic acid. SPRT and WLM domains (Figure 3B) are also very different in terms of amino acid sequence identity (5 % identical, 14% similar) and can only be aligned over a short region around the HEXXH motif (Figure 2S) [4]. In addition, SPRT domains have many highly conserved regions, which are not present in WLM domains (Figure S2). To further strengthen our finding that SPRTN and Wss1 are two independently-evolved enzymes, we modeled the structure of the SPRT domain of SPRTN using the recently solved crystal structure of Wss1b in fission yeast (Schizosaccharomyces pombe) (PDB 5JIG) [6] as a template (homology modelling, SWISS-MODEL) [69]. The only part of SPRTN that modelled to Wss1b with high confidence (Supplementary Methods) was a protease core consisting of two α-helices containing the HEXXH motif and the third zinc binding histidine residue (Figure 3C,
The described protein core of SPRTN is shared with other gluzincin metallopeptidases, including Wss1b among others. This is confirmed by a homology model of SPRTN domain using another zinzin protease, abylisin (PDB 4JIU) (Figure 3C, left). Indeed, the region over which the SPRT domain could be modelled with high confidence was longer when aligned to abylisin than to Wss1b. The additional part of SPRTN domain modeled with high confidence includes three β-sheets upstream of the HEXXH active centre.

Apart from the differences among the SPRT and WLM domains in terms of amino acid sequence and structure, both proteins differ distinctly in their C-terminal regions. SPRTN has a long C-terminal arm (276 amino acid long), while Wss1 has a comparatively short C-terminus (48 amino acid long) (Figure 3B). Other than both having a p97/Cdc48-binding motifs, the C-terminal regions of both proteins share no similarities: SPRTN contains a PCNA binding motif (PIP) and a ubiquitin-binding motif whereas Wss1 contains SUMO binding motifs (Figure 3B). The C-terminal part of SPRTN is involved in TLS, where it serves as a platform to recruit p97 at sites of stalled DNA replication fork. For detailed role of SPRTN and p97 in TLS please read the following literature [59, 70-75]. Therefore, we suggest that extrapolating similarities between SPRTN and Wss1 with respect to substrate specificity, affinity, mode of substrate binding and recruitment to chromatin should be done with caution. Most importantly, structural extrapolations should not be made before the crystal structure of SPRTN is solved. We conclude that the WLM and SPRT families are two separate families within the gluzincin subgroup of the zinzin superfamily. Like other gluzincins, they share similar properties such as an HEXXH proteolytic active site. Moreover, SPRTN is evolutionary closer to
bacterial SPRT proteases than to Wss1. Thus, we would like to clear up the confusion in the published literature, which occasionally states that yeast Wss1 is an ortholog of SPRTN. Any similar functional properties shared by these two proteases is a result of a convergent rather than divergent evolution [4].

**DNA Replication-coupled DPC Proteolysis Repair**

DPCs constitute strong physical blocks for the progression of DNA replication, causing DNA replication fork stalling and, consequently, fork collapse [4, 8]. Using *in vitro* approach in *Xenopus* egg extract, it was recently demonstrated that in order for replication to progress in the presence of DPCs, DPCs have to be cleaved into smaller peptides on both the leading and lagging DNA strand [44]. However, the protease involved in the processing of DPCs remained unknown until SPRTN was identified as the S-phase specific protease responsible for DPC repair [4] (Figure 2B). Recent findings showed that the majority of DPCs are indeed removed specifically during S-phase [4]. As a part of the replisome, SPRTN prevents fork stalling and DSB formation caused by DPC accumulation and protects replicative cells from DPC-induced toxicity [4, 8]. However, the precise orchestration of replication-coupled DPC-PR is still unknown, specifically: (i) which factors act downstream of SPRTN to remove peptide remnants after DPC proteolysis and whether this occurs only in S-phase; (ii) how SPRTN protease is regulated, and (iii) which other factors are involved in SPRTN-dependent DPC-PR. NER, TLS, HR are all possible candidates for action during S-phase, while NER and MRE11 nuclease activity remain active in non-cycling cells too. Furthermore, it is still an open question as to which factors act upstream of TDP1 in non-cycling cells considering that it is known that TDP1 removes TOP1cc peptide remnants at the sites of transcription stalling in post-mitotic
cells [76, 77]. Overall, we cannot exclude that DPC repair also occurs in non-cycling cells, especially in non-proliferative cells such as neurons where DPC accumulation would pose a threat to transcription. However, work presented here strongly suggests that SPRTN protease, as a component of the DNA replication machinery, forms a unique, proteolysis based DNA repair pathway for DPC repair during DNA synthesis.

**DPCs and human pathogenesis**

The contribution of DPCs to human pathogenesis was proposed by several studies that associated different DPC-inducing agents with ageing and cancer [78-80]. Mice exposed to formaldehyde accumulate DPCs mostly in their bone marrow and develop squamous cell carcinomas in the nasal passages upon formaldehyde inhalation [81, 82]. DPC accumulation was also correlated with ageing in both mice and humans [83, 84]. However, there was no direct evidence that defective DPC repair could be pathological until the recent characterization of SPARTAN syndrome also known as Ruijs-Aalfs Syndrome (RJALS), which is caused by mutations in the *SPRTN* gene [60, 61]. SPARTAN syndrome is characterised by premature ageing and early-onset hepatocellular carcinoma. At the cellular level, cells from SPARTAN syndrome patients accumulate DPCs and are unable to cope with DPCs during DNA replication, leading to DSB formation during S-phase [4, 7, 61]. Therefore, SPARTAN syndrome constitutes the first direct link between a defective DPC repair and the development of cancer and ageing in humans [4].
Concluding remarks and future perspectives

Repair of DPCs was dogmatically considered to be solely under the jurisdiction of canonical DNA repair pathways like NER and HR. However, recent independent work from several laboratories demonstrates that a specialised DNA repair pathway, which strictly depends on proteolysis, repairs DPCs. We named this novel pathway DNA-protein crosslink proteolysis repair (DPC-PR). The proteases involved in DPC-PR repair are Wss1 in yeast and SPARTAN in metazoans. The discoveries of these two proteases and a human syndrome resulting from defective DPC repair establish this new DNA repair pathway as an essential mechanism for genome maintenance and protection from accelerated ageing and cancer in mammals (see Outstanding questions).

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Glossary

**DNA damage repair (DDR)**: commonly referred to canonical DDR pathways: base excision repair (BER), nucleotide excision repair (NER), and mismatch repair (MMR) which repair single strand DNA damage and homologous recombination (HR) and non-homologous endjoining (NHEJ) which repair DNA double strand breaks.

**DNA damage tolerance (DDT)**: a mechanism of bypassing DNA lesions during DNA replication, thus allowing replication forks to progress, leaving the lesions to be repaired later.

**DNA-protein crosslink (DPC)**: any protein that is irreversibly covalently attached to the DNA.

**Homologous recombination (HR)**: an error-free canonical double strand break repair pathway which acts during S-phase.

**Nucleotide excision repair (NER)**: a canonical DNA damage repair pathway that excises DNA lesions on one DNA strand and is not cell cycle-specific.

**SPARTAN Syndrome also known as Ruijs-Aalfs syndrome (RJALS)**: a monogenic human syndrome with mutations in *SPRTN* gene characterised by premature aging phenotypes and early-onset hepatocellular carcinoma.

**SPRTN/DVC1**: a DNA-dependent metalloprotease that proteolytically digests the proteinaceous part of DPCs during DNA replication fork progression in metazoans.

**Wss1**: a DNA-dependent metalloprotease that proteolytically digests the proteinaceous part of DPCs during S-phase in yeast.

**Translesion DNA synthesis (TLS)**: a DDT pathway involving specialised DNA polymerases that can bypass unrepaired DNA lesions during replication. Some of the TLS polymerases are error-prone and thus TLS can be mutagenic.
Topoisomerase1 and 2 cleavage complexes (TOP1ccs and TOP2ccs): TOP1 or TOP2 which are covalently trapped to the DNA next to a single strand (ss) or a double strand (ds) DNA break, respectively. The strand breaks near TOPccs were created by the enzymes themselves during their normal enzymatic cycles.
Text Boxes

Box 1. Types of DPCs

DPCs are intricately complex DNA lesions. The complexity is a result of the diversity of crosslink formations in terms of protein size and physicochemical properties (charge, level of disorder), type of crosslink (crosslinked to one or both intact DNA strands) and number of covalent bonds (stability and structure of DPC), as well as temporal distribution of crosslink occurrence (cell cycle dependence). DPCs are commonly divided into two groups: (A) general, non-enzymatic DPCs (Type 1) which include any protein found in proximity to DNA at the moment of exposure to endogenous or exogenous DPC-inducing agents and (B) specific, enzymatic DPCs (Type 2) such as Topoisomerase 1 and 2 cleavage complexes (TOP1-ccs and TOP2-ccs), DNA polymerase (Pol) β and poly(ADP-ribose) polymerase 1 (PARP1) crosslinks. During their normal enzymatic cycles, these enzymes reversibly bind to DNA. However, upon treatment with anti-cancer drugs, camptothecin and/or etoposide, TOP1ccs and TOP2ccs are formed [24, 25], while abortive DNA repair can lead to Pol β and PARP1 crosslink formation [38]. Type 1 DPCs are the most prevalent under physiological conditions and they always include proteins crosslinked to an undisrupted DNA strand(s) [12]. Conversely, Type 2 DPCs include proteins crosslinked to a broken DNA backbone. TOP1 is crosslinked to the 3’ end of a ssDNA break, TOP2 to the two 5’ ends of a dsDNA break [46], while PARP1 is crosslinked to the 3’ end of a ssDNA break resulting from deficient DNA repair [85, 86]. In the case of topoisomerases, the breaks are created during the normal enzymatic decatenation reactions of topoisomerases and persist after crosslinking. The chemistry of both types of DPCs is explained in BOX 2. Altogether, this highlights that genomic DNA is constantly exposed to various DPCs, induced either by endogenous and
exogenous sources. Indeed, DPCs are among the most common DNA lesions. Thus, cells have to have specialised DPC recognition and repair mechanisms to prevent DPC-induced genotoxicity.

Box 2. Chemistry of DPC formation

Aldehydes react with proteins in the vicinity of DNA resulting in the formation of DPCs and base adducts [87-89] and to a lesser extent intra- and inter-strand crosslinks (ICLs), SSBs and DSBs [90]. Firstly, aldehydes react with lysine, cysteine and histidine residues on the protein forming a protein adduct which then reacts with the amino group on DNA bases resulting in crosslinks of variable stability. Target sites for protein crosslinking on DNA molecule include: N7 of guanine, C-5 methyl group of thymine and the exocyclic amino groups of guanine, cytosine and adenine [23], thus further expanding the complexity and diversity of crosslinks. Reactive oxygen and nitrogen species (ROS and RNS) can react with DNA (guanine, cytosine and thymine bases) and/or proteins (lysine and tyrosine side chains) resulting in the formation of free radicals or electrophilic lesions, which in turn react with another protein and DNA molecule thus creating the crosslink. For detail review on the chemistry of DPCs we refer the readers to Tretyakova et al. [23].

TOP1 and 2 relax DNA supercoils during DNA transactions (replication, transcription, recombination and repair) by decatenating DNA through single or double strand incisions, respectively, and re-ligating the nicked ends. The chemistry behind it is as follows: the catalytic tyrosine residue in topoisomerase becomes transiently covalently attached to the DNA phosphate at the 3' (TOP1) or 5' (TOP2) end of the broken DNA as a result of nucleophilic attack by the catalytic tyrosine on
the DNA phosphodiester bond. Normally, this topoisomerase-DNA covalent reaction intermediate would then dissociate and topoisomerase would re-ligate the broken DNA ends. However, camptothecin (TOP1) and etoposide (TOP2) bind to the catalytic site at the enzyme-DNA interface, thus preventing re-ligation and leaving the enzyme trapped to DNA and the strand break/s unsealed [24, 25, 91]. Topoisomerases can also become trapped to the DNA backbone during their normal enzymatic cycles if in proximity to oxidative radicals or upon encountering bulky DNA adducts, ribonucleotides or abasic sites [46]. Given the high demand for DNA relaxation by topoisomerases during all DNA transactions, it is expected that the number of topoisomerases bound to DNA is high. Indeed, it was recently confirmed that the incidence of endogenous TOP2ccs is much higher than previously thought and that they cause severe genomic instability if left unrepaired [39].

**Box 3. The repair of Topoisomerase 1 and 2 cleavage complexes (TOP1- and 2-ccs)**

The repair of TOP-ccs (enzymatic DPCs) has been extensively studied in recent decades due to their importance for cancer therapy. Many therapeutic approaches rely on the inhibiton of topoisomerases through the action of topoisomerase poisons, camptothecin and etoposide, which induce TOPccs with the aim to stop cancer proliferation and/or induce cancer cell death. Indeed, therapy for more than 30-50% of cancers relies on the use of topoisomerase inhibitors [46]. Until very recent advances, it was thought that TOP1cc and TOP2cc repair relies mainly on the action of phosphodiesterase enzymes, tyrosyl-DNA-phosphodiesterase 1 and 2 (TDP1 and TDP2), respectively. These enzymes act on the DNA backbone by excising small
peptide remnants left from TOPccs, thus leaving ss (TDP1) or ds breaks (TDP2) which are subsequently repaired by canonical DDR pathways. More specifically, TDP2 cleaves DNA by hydrolyzing 5’ tyrosine phosphodiester bonds which converts them into 5’phosphate ends which are subsequently repaired by NHEJ [91]. Although the mechanism of action of TDPs was known, the upstream factors involved in TOPcc repair remained unknown. More specifically, how TOPccs are cleaved into smaller peptides, thus enabling TDPs to act was only resolved recently when the protease SPRTN was found to remove TOPccs in vitro and in vivo [4, 7]. SPRTN proteolytically digests TOP1 and TOP2 thus reducing the size of the DPCs and enabling the action of TDPs. Indeed, it was shown that SPRTN and TDP1 act epistatically in the repair of TOP1ccs [4]. Recently, a separate mechanism for TOP2cc repair was discovered, one which relies on the action of the nuclease Mre11, otherwise known as a part of the MRN complex with a main role in HR [39-41]. Mre11 removes TOP2ccs independently of HR, while NHEJ acts downstream of Mre11 activity to repair ds breaks left after nuclease action [39]. Additionally, Mre11-mediated TOP2cc removal is a dominant and separate pathway to TDP2-mediated TOP2cc repair [39]. In light of these recent discoveries it is possible that SPRTN and TDP1 and TDP2 form a separate pathway for TOPcc repair, distinct from Mre11-mediated TOP2cc removal. However, we cannot exclude the possibility that SPRTN is also needed in Mre11-mediated TOP2cc removal to reduce the size of the proteinaceous part of the DPC. The precise coordination of these two pathways remains to be determined.
Figure legends

**Figure 1. DNA repair pathways.** Schematic model of various DNA lesions caused by different genotoxic agents and respective DNA repair pathways. Specialised DNA repair pathways cope with specific type of DNA lesions. DNA lesions caused by the covalent attachment of bulky protein, DNA-protein crosslink (DPC) are predominantly repaired by DNA-protein crosslink proteolysis repair (DPC-PR). DDR; DNA damage response, DDT; DNA damage tolerance, BER; base excision repair, NER; nucleotide excision repair, HR; homologous recombination, NHEJ; non-homologous end joining, MMR; mismatch repair, FA; Fanconi Anemia, TLS; translesion DNA synthesis, DPC; DNA-protein crosslink.

**Figure 2. (A) Comparison of NER, HR and DPC-PR in respect to repair of DPCs.** Upper panel; Colour code represents activity of three known repair pathways for DPC repair during the cell cycle. Green = NER, Blue = HR and Violet = DPC-PR. Lower panel: DPC repair pathways and their cellular characteristics and consequences. Mutations in these pathways cause several inborn diseases; XP; Xeroderma Pigmentosum, TTD; Trichothiodystrophy, CS; Cockayne Syndrome, ATLD; Ataxia-teleangiecstasia-like disorder, NBS; Nijmegen breakage syndrome, SPARTAN syndrome (also known as Ruijs-Aalfs) **(B) SPRTN is a central player in replication-coupled proteolysis repair pathway of DPCs.** SPRTN proteolytically digests the proteinaceous part of bulky DPCs which block DNA replication fork progression. After digesting the DPC, SPRTN inactivates itself by self-cleavage.

**Figure 3. (A) Evolutionary analysis of SPRT and WLM families.** Alanyl aminopeptidase family, another member of the gluzincins subgroup of zinzin
metalloproteases, was used to compare the evolutionary relationship between the
SPRT and WLM families. SPRT is equally distant to the WLM group as it is to
Alanyl aminopeptidase group, thus there is no indication of common ancestry
between the SPRT and WLM families. The SPRT family is present in bacteria,
arachnea, cyanobacteria, plants and animals, while the WLM family emerged in yeast,
fungi and plants. Accession numbers of sequences used in the analysis, methodology
for phylogenetic analysis as well as an expanded phylogenetic tree including branch
support values are available in the supplementary material. (B) Domain organization
of SPRTN and Wss1. SPRTN and Wss1 share a short consensus HEXXH motifs in
the active centre of the protease, a common property of all zinzin metalloproteases,
and a p97/Cdc48 binding motif (SHP and VIM, in blue). C-terminal regions
(downstream of SPRT and/or WLM domain) differ substantially between the proteins,
with the long C-terminal arm of SPRTN (276 amino acids) bearing a PCNA binding
(PIP, in light blue) and ubiquitin binding motifs (UBZ, in orange) and the short C-
terminal arm of Wss1 (48 amino acids) bearing sumo interaction motifs (SIM, in red).
(C) Homology model of human SPRT domain. The human SPRT domain was
modelled according to yeast Wss1b (left panel) (5JIG) and another gluzincin member,
abylysin (right panel) (4JIU) using the SWISS-MODEL workspace (see
supplementary methods). Both models show a conserved protease core of SPRTN
with two α-helices (in green) that contain the catalytic active centre, including three
zinc binding histidines (in red) and a glutamic acid (in blue). Abylysin-based model
gave broader coverage of SPRTN domain (66. – 142. amino acids of full length
SPRTN) and higher model confidence (see supplementary methods) compared to the
Wss1b-based model coverage (103.-144. amino acids), thus confirming that SPRTN
is equally structurally similar to Wss1 as to any other gluzincin protein. Three β-
sheets (orange) upstream of the catalytic core were modeled with high confidence according to the abylysin structure (right panel), while two β-sheets between the two α-helices in the Wss1b-based model are modeled with too low confidence in order to be considered reliable (see supplementary material).
**Outstanding questions**

Which factors act downstream of SPRTN proteolysis? How is the whole DPC repair pathway orchestrated and the choice between different downstream pathways made, e.g. NER, HR, Mre11, TDPs and TLS?

How are DPCs removed in non-cycling, lowly proliferative cells, where they pose a threat to transcription progression?

How are TOP1ccs and TOP2ccs predominantly repaired, via SPRTN cleavage followed by TDP-mediated peptide excision or via Mre11 through the excision of ssDNA overhang bearing the TOP2cc? Can Mre11 also act in the removal of other DPCs in vivo?

How is SPRTN protease regulated? What is the signal to trigger SPRTN-mediated proteolysis and how is its pleiotropic protease activity kept in check to prevent unspecific substrate cleavage?

What is the exact mechanism behind the link between DPC accumulation and the carcinogenesis and premature aging observed in SPARTAN syndrome patients?
DNA repair pathways

Damage Source
- X-rays
- Oxygen radicals
- Alkylating agents
- Anti-tumor agents (cis-DDP, MMC)
- Replication fork collapse
- UV light
- Polycyclic aromatic hydrocarbons
- Formaldehyde
- Anti-tumor agents (cis-DDP, MMC)
- DNA replication of damaged template base

DNA lesions
- Uracil
- Abasic site
- 8-Oxo guanine
- SSB
- DSB
- (6-4)PP
- Bulky adduct
- CPD
- A-G Mismatch
- T-C Mismatch
- Insertion
- Deletion
- DNA Interstrand crosslinks (ICs)
- DNA-Protein crosslinks (DPCs)
- UV-induced lesions
- Abasic site
- Base lesion

Pathway
- BER
- HR
- NHEJ
- NER
- MMR
- FA
- DPC-PR
- HR
- NER
- TLS
- Template switch

Figure 1
(A) Repair of DPCs throughout the cell cycle

(B) DPC-PR pathway

**Repair Pathway**
- Proteolytic dependent
- Size limitation
- DSB formation
- Replication dependent
- Recombinogenic

**Impaired repair**
- DPCs accumulation
- DSB formation
- Cancer and ageing

**Inborn disease**
- XP
- TTD
- CS
- ATLD
- NBS
- SPARTAN syndrome (RJALS)

**Outcome**
- DPCs removal
- Normal replication
- No DSB
- Genomic stability

Figure 2
Figure 3

(A) Evolutionary tree

(B) Domain organization

(C) Homology models of SPRT domain

Wss1b template
Abylsin template
Inventory of Supplementary Information

1. Supplementary methods include methodology used to construct phylogenetic tree in Figure 3A and to perform homology modelling of the SPRT domain of human SPRTN based on the newly published structure of yeast Wss1b and zinzip metallopeptidase abylysin shown in Figure 3C.

2. Figure S1 contains phylogenetic tree with branch support values and full protein names which is complementary to phylogenetic tree shown in Figure 3A. Related to Figure 3A.

3. Table S1 contains protein accession codes from NCBI database of sequences used to construct phylogenetic tree in Figure 3A. Related to Figure 3A.

4. Figure S2 contains multiple sequence alignment of SPRT and WLM domains across species. Related to Figure 3C.

5. Supplementary references

1. Supplementary methods

PSI-BLAST (Position-Specific Iterated Blast) was used to identify orthologs of SRTN in bacteria, archaea, cyanobacteria, yeast, fungi, plants and animals by blasting the SPRT domain protein sequence of human SPRTN through the NCBI database (National Center for Biotechnology Information) [1]. The same approach was used to identify WLM domain-containing proteins using the WLM domain protein sequence from S. cerevisiae Wss1. Alanyl aminopeptidases (leukotriene A-4 hydrolase) protein sequences were downloaded from NCBI, and M1-LTA4H domains containing a HEXXH protease core were used for alignment and tree construction. Multiple sequence alignments were done using MAFFT [2]. Quality of alignment was estimated with Guidance software (alignment score was 0.532896) [3]. Phylogenetic tree was constructed using Maximum Likelihood method in PhyML 3.0.1 software (LG model, 10 rate categories, best of NNI and SPR for tree searching operations) [4]. Confidence of nodes was estimated by approximate likelihood ratio test (aLRT) [5]. Homology modelling of human SPRT domain was done using the crystal structure of Wss1b in fission yeast (Schizosaccharomyces pombe) (PDB: 5JIG) as a
template or using zinzin protease, ablyysin (PDB: 4JIU) in the SWISS-MODEL workspace [6, 7]. Confidence of the models was estimated using the QMEAN scoring function [8].
2. **Figure S1.** Phylogenetic tree with branch support values and full protein names which is complementary to phylogenetic tree shown in Figure 3A.
### Table S1. Protein accession codes for sequences used to construct phylogenetic tree in Figure 3A.

| Abbreviation | Accession | Species |
|--------------|-----------|---------|
| NP_000886.1  | HsLTA4H   | Homo sapiens |
| XP_002711258.1 | OcLTA4H | Oryctolagus cuniculus |
| NP_001025202.1 | RnLTA4H | Rattus norvegicus |
| XP_011282463.1 | FcLTA4H | Fels catus |
| NP_000664289.1 | SoLTA4H | Sus scrofa |
| XP_001309392.1 | OaLTA4H | Ornithorhyncus anatinus |
| NP_01006234.1  | GgLTA4H   | Gallus gallus |
| XP_002189060.1 | TgLTA4H | Taeniopygia gutata |
| NP_001006234.1 | HsSPRTN | Homo sapiens |
| NP_001104611.1 | MmSPRTN | Mus musculus |
| XP_003496855 | CgSprt | Cricetulus griseus |
| XP_008122534.1 | AcSPRTN | Anolis carolinensis |
| XP_005173863.1 | DrSprtn | Danio rerio |
| NP_59307 | SpWss1 | Saccharomyces pombe |
| NP_589231 | SpWss2 | Saccharomyces pombe |
| XP_003710670.1 | SySprtn1 | Synechococcus sp. |
| XP_003714265.1 | MoWss1 | Magnaporthe oryzae |
| XP_003709311 | MoWss2 | Magnaporthe oryzae |
| NP_001143131 | ZmWss1 | Zea mays |
| XP_008659182 | ZmWss2 | Zea mays |
4. Figure S2. Multiple sequence alignment of SPRT domain and WLM domain across species

Conserved regions specific for SPRT domains are shown in grey, active site motif HEXXH is shown in a black frame.
5. Supplementary References

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