

## **ScienceDirect**



# Serine ADP-ribosylation in DNA-damage response regulation

Luca Palazzo<sup>1</sup>, Marcin J Suskiewicz<sup>2</sup> and Ivan Ahel<sup>2</sup>



PARP1 and PARP2 govern the DNA-damage response by catalysing the reversible post-translational modification ADPribosylation. During the repair of DNA lesions, PARP1 and PARP2 combine with an accessory factor HPF1, which is required for the modification of target proteins on serine residues. Although the physiological role of individual ADPribosylation sites is still unclear, serine ADP-ribosylation at damage sites leads to the recruitment of chromatin remodellers and repair factors to ensure efficient DNA repair. ADPribosylation signalling is tightly controlled by the coordinated activities of (ADP-ribosyl)glycohydrolases PARG and ARH3 that, by reversing the modification, guarantee proper kinetics of DNA repair and cell cycle re-entry. The recent advances in the structural and mechanistic understanding of ADP-ribosylation provide new insights into human physiopathology and cancer therapy.

#### Addresses

<sup>1</sup> Institute for the Experimental Endocrinology and Oncology, National Research Council of Italy, Via Tommaso de Amicis 95, 80145 Naples, Italy

<sup>2</sup> Sir William Dunn School of Pathology, University of Oxford, South Parks Road, Oxford OX1 3RE, UK

Corresponding author: Ahel, Ivan (ivan.ahel@path.ox.ac.uk)

#### Current Opinion in Genetics & Development 2021, 71:106-113

This review comes from a themed issue on **Mechanisms of Homo- logous Recombination** 

Edited by Eric C Greene and Rodney Rothstein

For a complete overview see the Issue and the Editorial

Available online 31st July 2021

https://doi.org/10.1016/j.gde.2021.07.005

0959-437X/© 2021 The Author(s). Published by Elsevier Ltd. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).

#### Introduction

Reversible post-translational modifications (PTMs) control the spatio-temporal organisation of DNA-damage response (DDR). PTMs trigger and regulate many aspects of DDR, including recognition of DNA damage sites, chromatin remodelling, recruitment of DDR factors, initiation and execution of DNA repair, as well as cell cycle arrest [1]. A variety of PTMs are involved in DDR, including selective protein phosphorylation and dephosphorylation, as well as conjugation of proteinaceous modifiers such as ubiquitin, SUMO, and Nedd8 [2]. Importantly, the different types of modification involved appear to be highly integrated and

even dependent on each other. A critical example is the phosphorylation of the histone variant H2AX (termed  $\gamma$ H2AX) by phosphatidylinositol-3-kinase-related kinases ATM, ATR, and DNA-PK that is required for DDR-linked ubiquitylation [3]. Failure to organise a proper DNA repair response strongly affects maintenance of genome integrity, thus predisposing to various human disorders, such as cancer, neurodegeneration, and immunodeficiency. Therefore, the cellular response to DNA damage as well as the PTMs involved within it have received a great deal of attention in transitional medicine [4].

Prominent among PTMs triggered by DNA damage is ADP-ribosylation, a reversible PTM of proteins that is involved in many cellular processes, such as transcription, cell division, and survival [5]. ADP-ribosylation results in the covalent attachment of a single ADP-ribose unit [namely, mono(ADP-ribosyl)ation (MARylation)] or polymers of ADP-ribose units [namely, poly(ADP-ribosyl)ation (PARylation)] to a variety of amino-acid residues on target proteins [6]. The attached ADP-ribose moieties originate from NAD+, which is cleaved during the reaction, releasing nicotinamide. Interestingly, ADP-ribosylation can also occur on terminal DNA or RNA phosphates, thus representing a novel type of nucleic acids modification during DNA repair [7\*,8–10]. Notably, pharmacological modulation of ADP-ribosylation reactions represents one of the most successful examples of anticancer interventions (i.e. PARP inhibitors; PARPi) [4].

Proteomic studies revealed that one third of the human nuclear proteome in the presence of DNA damage is subject to ADP-ribosylation, mainly on serine residues [11°,12°,13°]. The two types of ADP-ribosylation — MARylation and PARylation — can be recognised by different reader proteins [5], which are also capable, in some cases, of distinguishing between short and long PAR chains [14]. This realisation, together with the indications of the functional relevance of PAR chain branching [15], suggests the existence of a complex 'PAR code' [16] akin to that described for ubiquitin [17] and hints at multiple regulatory roles of ADP-ribosylation in DDR. The investigation of ADP-ribosylation functions in DDR has been facilitated by novel antibodies and antibody-like reagents, which are able to distinguish between different lengths and, in some cases, sites of the modification [18,19°].

The best-studied ADP-ribosylation 'writer' is poly(ADP-ribose) polymerase 1 (PARP1), the founding member of the PARP family of ADP-ribosyltransferases (ARTs)

which in humans consists of 17 core members (PARP1 to PARP16) [20,21].

PARP1, which is thought to account for the greatest part of detectable ADP-ribosylation in human cells under DNA damage, is one of the most abundant nuclear proteins and, in addition to DNA repair, acts as a key player in chromatin remodelling, apoptosis, transcription, telomere maintenance and DNA replication [5.6.21-23,24°°]. Several PARP1 functions in DNA repair are complementary to those of PARP2 [25]. Genetic disruption of PARP1 and PARP2 simultaneously, but not individually, results in embryonic lethality in mice, demonstrating their functional redundancy [26].

PARP1 is a multi-domain protein, composed of four DNAbinding domains, namely three Zn-fingers, one tryptophanglycine-arginine (WGR) domain, an auto-modification domain consisting of a BRCT motif, and a catalytic domain [21]. The combination of different DNA-binding domains allows recognition of various DNA breaks including singlestrand and double-strand ones [27°,28°]. Interestingly, PARP1 appears to be released from a DNA end by binding to another, which could allow it to move between DNA ends in a manner that was termed the 'monkey bar' mechanism [29]. While this mechanism does not result in net dissociation of PARP1 from DNA ends, this can be achieved through PARP1 auto-modification, which is thought to counter DNA binding through electrostatic repulsion between negatively charged PAR chains and DNA backbone [30,31]. In cells, PAR chains can additionally contribute to releasing PARP1 from chromatin by recruiting factors, such as the chromatin remodeller ALC1/CHDL1 or the scaffold XRCC1, that facilitate this process [32–34]. It is largely by interfering with PARP1 auto-modification that PARP1 inhibitors induce prolonged residence of PARP1 on chromatin, a phenomenon that is known as 'trapping' and thought to be the key to cytotoxic effects of PARP1 inhibition in cancer [35].

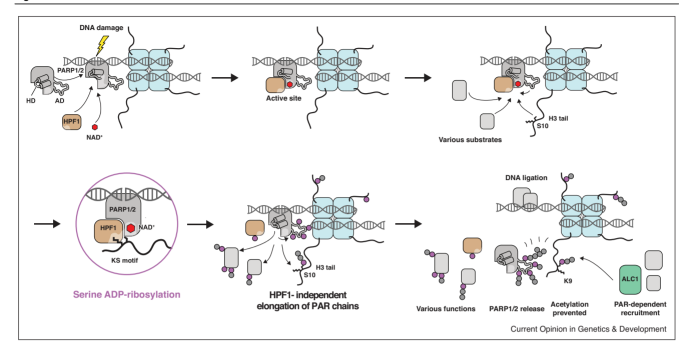
Compared with PARP1, PARP2 has a shorter N-terminal DNA-binding domain, which comprises an unstructured region and a WGR domain, while the catalytic domain is highly homologous to that of PARP1, which results in the majority of PARPi anticancer drugs targeting both PARP1 and PARP2 enzymes [27\*\*]. While the primary role of PARP1 and PARP2 is linked to their ADP-ribosylation activity, the two proteins might also serve other, noncatalytic roles. In particular, recent PARP2 structures suggest that double-strand DNA break recognition not only activates the catalytic domain, but also aligns the two DNA ends such that the DNA appears continuous across the break, seemingly poised for ligation [36,37,38,39].

Here, we discuss the mechanisms by which PARP1 and PARP2-dependent serine ADP-ribosylation controls DDR.

## Histone PARylation factor 1 (HPF1) drives DNA damage repair response by enabling PARP1 and PARP2 to catalyse serine ADPribosylation

PARP1 and PARP2 act as 'first responders' at DNA lesions. Both enzymes are recruited within seconds of the occurrence of various types of DNA damage and rapidly catalyse protein modification [40,41]. PARP1 and PARP2 are allosterically regulated by binding to DNA breaks, which leads to remodelling of an autoinhibitory fragment of the catalytic domain, called the helical domain (HD), which inhibits NAD<sup>+</sup> binding in the resting state [37\*\*,42\*\*]. Biochemical studies accompanied by new mass spectrometry tools have established that the main targets of ADP-ribosylation under both basal and DNA-damage conditions are serine residues in substrate proteins [11°,12°,13°,43°,44°,45]. Targeted sites are enriched for lysine-serine (KS) and, to a lesser extent, arginine-serine (RS) motifs [11°,12°,44°]. Although serine ADP-ribosylation has been detected on a large fraction of human proteins, the bulk of the modification corresponds to a limited number of key sites within PARP1 and PARP2 themselves and histone tails [12°,43°]. Intriguingly, the proximity of ADP-ribosylable serine sites (e.g. S10 of histone 3; H3S10) to lysine residues, which are known targets of acetylation (e.g. H3K9) as well as the overlap with serine phosphorylation sites (e.g. H3S10), suggests an important contribution of histone ADP-ribosylation to a multi-layered cross-talk DDR-relevant **PTMs** with additional histone [44°,46°,47°]. Recent studies revealed that, surprisingly. PARP1 and PARP2 are by themselves insufficient for ADP-ribosylating serine residues. Histone PARylation factor 1 (HPF1), which binds the catalytic domain of PARP1 or PARP2, is an essential participant in this process [12°,37°,43°,48°,49°]. The interaction between HPF1 and PARP1/2 — like NAD+ binding — is negatively regulated by the HD fragment of these PARPs and is therefore enhanced upon their binding to damaged chromatin [37°°,38,49°°] (Figure 1). Strikingly, HPF1 together with PARP1/2 jointly forms the specific peptide-binding site, but also contributes a key residue (E284), termed the 'glutamate finger', to the catalytic core of PARP1 or PARP2 [49\*\*]. Consistent with its catalytic role, E284 is required for in vitro modification of serine-containing substrates and for DNA damageinduced ADP-ribose signalling in human cells despite being dispensable for the HPF1-PARP interaction and HPF1 stability [49°,50,51°]. E284 seemingly acts as a general base that deprotonates the serine hydroxyl group to facilitate the subsequent nucleophilic attack on the 1" carbon of NAD<sup>+</sup>. The importance of acceptor deprotonation for the ADP-ribosylation reaction has long been postulated [52,53] and is underscored by the fact that, in vitro and in the absence of HPF1, PARP1 and PARP2 preferentially modify glutamate and aspartate residues, which are constitutively deprotonated at a neutral pH

Figure 1



Model of HPF1-dependent PARP1 and PARP2 activation and function at the DNA damage foci. Upon DNA damage, an inactive conformation of PARP1 or PARP2 is recruited to DNA. PARP1 and PARP2 undergo a conformational change in the regulatory domain (HD) allowing NAD+ binding. HPF1 binds to the PARP catalytic domain allowing the initiation of ADP-ribosylation of serine residues in target macromolecules, such as histone H3 tails, various DNA repair factors, and PARP1 and PARP2 themselves. Once the initial ADPribose is attached, PARP1/2 can catalyse the PAR chain elongation reaction without HPF1, which may dissociate from the complex. ADPribosylation allows recruitment of PAR-readers (i.e. ALC1) to DNA damage sites, which, in turn, trigger DNA repair and ligation. Furthermore, ADPribosylation at DNA damage sites contributes to modulate proximal chromatin epigenetics, regulates the activity and function of various substrates, and controls PARP1/2 release from DNA.

[54]. Indeed, while serine residues are the predominant cellular targets, the alternative activity of PARP1 and/or PARP2 towards glutamate and aspartate residues can be detected in cells, where it becomes relatively more prominent upon HPF1 knockout [43°]. In addition to activating the serine side-chains, HPF1 might use a broader negatively charged region that surrounds E284 to help recognise the lysine or arginine residue that precedes the serine in KS or RS motifs [49\*\*].

The discovery of HPF1 sheds new light on the distinction between MARylation and PARylation. HPF1 binding to PARP1 is sterically incompatible with chain elongation [49°], which results in HPF1 having a negative effect on chain length, especially when it is present at a saturating level in an *in vitro* reaction or overexpressed in cells [48°°]. When present at a high level, HPF1 instead turns PARP1 into an NAD<sup>+</sup> hydrolase by discouraging the attachment of ADP-ribosylation to proteins as soon as available serine sites become singly modified [51°]. However, it is unclear what the relevance of the hydrolase activity is in cells. Paradoxically, HPF1 is at the same time required for normal PARylation by contributing to the attachment of the first ADP-ribosylation unit that is a prerequisite for subsequent chain formation, and appears to be the ratelimiting step of the whole PARylation process [24\*\*]. DNA damage induced PARylation could be seen as consisting of two distinct stages, where the first consists in the attachment of an ADP-ribosylation unit to a protein side-chain or 'chain initiation' (achieved by PARP1 or PARP2 in HPF1-depend manner) and the second in the attachment of consecutive ADP-ribosylation units or 'chain elongation' (catalysed by PARP1 or PARP2 alone and inhibited by HPF1) [24\*\*]. The complex role of HPF1 in this process could explain why the levels of this protein in the cell are kept low [48°] — enough to efficiently initiate chains but not to preclude extension. Interestingly, as explained below, the two stages of serine ADP-ribosylation are also largely distinct when it comes to their regulation by hydrolases, with initiation being counteracted by the serine-specific ARH3 and elongation by PARG [24\*\*]. Dysregulation of ADP-ribosylation initiation by either HPF1 knockout or ARH3 upregulation results in impaired PARP1 auto-modification and increased trapping and inhibitor sensitivity in human cells [35].

## Functional consequences of PARP1/PARP2mediated ADP-ribosylation in response to DNA damage

Functional consequences of ADP-ribosylation reactions are poorly understood and have thus received a great deal of attention in the field in recent years. PARvlation has more profound consequences in the cells compared with MARylation [24\*\*] and can even suppress DNA repair when it is excessive [55]. For these reasons, PARvlation is present only transiently in response to DNA damage. In vitro experiments showed that, within the first ten minutes after DNA damage by H<sub>2</sub>O<sub>2</sub>, PARylation is converted to MARylation by the poly(ADP-ribose)glycohydrolase (PARG) [19°,56,57°°]. Indeed, PARylation, if not reversed in a timely manner, appears to be very toxic to the cells [24°,58,59], for instance by impairing alternative lengthening of telomeres (ALT) and global transcription [24\*\*]. In contrast, MARylation marks are well tolerated, and are hence compatible with recovery after DNA damage and cell division. This observation is unexpected considering that ADP-ribosylation marks were shown to be able to interfere with canonical histone modifications required for cell proliferation (i.e. H3S10 phosphorylation) [24°,43°,60].

As mentioned above, it is only recently that ADP-ribosylation events have been mapped to defined sites in protein substrates and limited information is available on functional consequences of individual modification events. Although it is likely that PARylation affects protein functions and dynamics [61], the only understood consequence of the site specific ADP-ribosylation is the role of PARP1 auto-modification at the 3 serines residues (i.e. 499, 507, and 519). Those serine residues are the predominant in vivo PARP1 auto-modification sites and are crucial for PARP1 release from DNA, therefore their mutation sensitises cells to PARP inhibitors [35]. Nevertheless, the best-understood aspect of proteins' PARylation at DNA foci remains scaffolding. A number of DNA damage response proteins, including specific chromatin modifiers, are rapidly recruited through direct binding of the modification, often mediated by specific protein modules ('readers' domains), such as the macrodomain, PBZ, and WWE domains [5]. Importantly, diverse parts of the modification can be recognised by different proteins readers, thus able to distinguish between MARylation and PARvlation of various lengths [5,14,62]. This observation also suggests a sort of physical and functional compartmentalisation of readers, where some factors may enrich preferentially at long PAR chains, which are mainly observed on auto-modified PARP1 and PARP2. By contrast, other factors may bind with higher affinity short PAR chains or MARylation, mainly present on histone proteins. Biochemical analysis indicates that, upon HPF1-PARP binding to nucleosomes, histone H3 is rapidly modified to saturation at the DNA break, whereas PARP2 continues to accumulate PARylation over time [37°°]. This suggests that, at least for PARP2, the histone modification serves as the initial anchor for recruiting factors to sites of DNA damage. Considering the high overall similarity in the mode of action of the two proteins, PARP1 may act analogously to PARP2; nevertheless, further biochemical and ultimately cellular studies will be needed to investigate this in detail. By contrast to histone ADP-ribosylation, PARP1/PARP2 auto-modification may not present a persistent mark of DNA damage sites, as it ultimately leads to PARP release from DNA, consistent with previous observations [35,63–65]. Of note, PARP2 engages DNA ends in a manner that could prevent access of repair factor access to the break, suggesting its dissociation might be a prerequisite for subsequent repair [37\*\*], consistent with previous observations for PARP1 [66]. In this respect, the ADP-ribosyl modification on histones might serve as a break-proximal recruiting platform and the key determinant of subsequent repair events. However, at this stage it is difficult to pronounce with confidence on the relative functional importance of histone-linked and PARP-linked modification, as both can in theory act as signals that attract DNA repair factors, many of which are recruited at DNA damage foci by ADP-ribosylation, including ALC1, APLF (aprataxin PNK-like factor), and XRCC1 [25,67,68] (Figure 1).

Among PAR readers, the helicase ALC1 possesses a macrodomain, which allows PAR binding and recruitment to DNA damage foci, thereby facilitating an open chromatin structure, and, in turn, increasing accessibility of additional repair factors to DNA [68,69]. Depletion of HPF1 abolishes recruitment of DDR factors, as illustrated for recruitment of LIG3 and XRCC1 to Okazaki fragments [70]. Similarly, the PARP1-XRCC1 axis, and possibly HPF1, is involved in short-patch single-strand break (SSB) repair physiologically occurring at enhancers as a response to neuron-specific transcriptional activity [23]. Altogether, these recent observations confirm the importance of ADP-ribosylation in triggering a proper DDR, which may be required in multiple physiological conditions.

#### Reversal of serine ADP-ribosylation

Two enzymes are required for reversal of serine PARvlation, namely PARG and ARH3 [71]. These enzymes function in a cooperative manner; PARG very efficiently cleaves PAR chains [72] but is incapable of removing MARylation [73,74]. De-MARylation is instead catalysed by ARH3, which selectively cleaves the O-glycosidic linkage between the serine of target proteins and ADP-ribose [57\*\*]. PARG is highly active in cells and accounts for the conversion in MARylation of almost all PAR chains within minutes upon DNA damage, therefore in a physiological context MARylation on serine appears to be the persistent form of ADP-ribosylation [19°,24°,57°]. PARG inhibition prevents conversion of PARylation to MARylation [19,75, implying that PARG is required to keep PARylation under control and balance its 'positive' (i.e. recruitment of DNA repair factors) and 'negative' (i.e. toxic effects and pro-apoptosis) roles. ARH3 instead prevents chain formation by removing the priming unit of ADP-ribose covalently linked to modified substrates, without which a PAR chain cannot be extended [24°,57°]. Thus, PARG and ARH3, although primarily active at different stages of de-ADPribosylation reaction, show synergy and their joint inactivation is required to unleash widespread, and ultimately toxic, PARylation [24°].

#### **Conclusions**

The understanding of DDR control by PARP1-dependent or PARP2-dependent ADP-ribosylation at the molecular level has improved significantly in the last decade. With the development of new biochemical tools, proteomic approaches, electron microscopy, and genetic screens, new key players have been identified, providing novel mechanistic insights. Major breakthroughs in the field in the last years were the identification of HPF1 and the dissection of the molecular mechanisms of the HPF1-PARP1/2 complex catalytic activity, the characterisation of serine residues as the main targets of ADP-ribosylation in response to DDR, better elucidation of recruiting and scaffolding roles of this modification, and, not least, the cooperation between PARG and ARH3 in counteracting PARP1 and PARP2 activity.

Considering that the PARP inhibitors, currently in clinical use, were identified before the discovery of such mechanisms, it may be of interest to re-evaluate the potency and in vitro selectivity of existing PARPi in the context of HPF1-PARP1/2 complexes, although one must bear in mind that such complexes might be relatively scarce and short-lived in the cell, PARP1/2 presumably remaining predominantly unbound at any given moment. The emerging picture is that while HPF1 might increase affinity of PARP1 for some existing inhibitors [35,76°], this regulatory factor actually promotes resistance to PARPi in the cell by stimulating PARP1 auto-modification, which in turn counteracts inhibitor-induced trapping [35]. Future studies of existing and design of novel PARPi should take into account recent insights into allosteric communication between DNA-binding and catalytic domains of PARP1 [77\*\*].

From the 'erasers' point of view, arising therapeutic potential of PARG inhibition (PARGi) in cancer therapy has also been investigated. PARGi is indeed very efficient in inducing toxic accumulation of cellular PARylated proteins, thus promising advantages for treatment of PARPi-resistant tumours [78\*\*]. Moreover, our studies suggest the potential for ARH3 as a novel drug target for cancer therapy, for instance in improving PARGi cytotoxicity or in selectively killing PARG-null cancer

cells [24\*\*]. On the latter point, PARG depletion is a major PARPi resistance mechanism of serous ovarian and triple-negative breast cancers [75\*\*] and could be targeted with ARH3 inhibitors. Conversely, downregulation of ARH3 confers resistance to PARPi [24\*\*,35], which phenocopies the effect of *PARG* loss or PARG inhibition [75°,78°]. These major breakthroughs open new and fascinating scenarios regarding development of new therapeutic strategies for PARPi-resistant cancers.

As the understanding of HPF1 and ARH3 molecular functions provided insights into the elucidation of human diseases and their treatment [24°,35,48°], many other DDR factors controlled by ADP-ribosylation signalling may equally impact therapeutics and/or prognosis of human disorders. For instance, the central role of ALC1 in allowing access of DDR factors at damaged foci may explain the crucial impact of its expression levels on sensitivity to PARPi in cancer cells [32–34,79,80]. These studies provide the rationale for designing novel therapeutics focused on inhibiting the recruitment of ADPribosylation readers, such as ALC1, to DNA damage foci to further sensitise cancer cells to PARPi.

Altogether, ADP-ribosylation is of vital importance in DDR and its further investigation promises new insights, which may significantly contribute to the understanding of human pathophysiology and treatment of human diseases.

## **Conflict of interest statement**

Nothing declared.

### **CRediT** authorship contribution statement

Luca Palazzo: Writing - original draft, Writing - review & editing. Marcin J Suskiewicz: Writing - original draft, Writing - review & editing. Ivan Ahel: Supervision, Writing - review & editing.

## **Acknowledgements**

We are grateful to Alessandra Peters, Johannes Rack, and Roberta Visconti for comments and suggestions, and for proof reading the article. The authors apologise to those whose work was omitted due to space restrictions.

This work was supported by the Wellcome Trust [grant numbers 101794 and 210634] to IA; BBSRC, [grant number BB/R007195/1] to IA; Cancer Research UK, [grant number C35050/A22284] to IA; the Italian Foundation for Cancer Research, [grant number 14895] to LP; the POR Campania FESR 2014/2020, [grant IDs SATIN and RECOVER-COVID19] to L.P; the Ovarian Cancer Research Alliance [grant number 813369] to IA and LP.

#### References and recommended reading

Papers of particular interest, published within the period of review, have been highlighted as:

- · of special interest
- of outstanding interest
- Polo SE, Jackson SP: Dynamics of DNA damage response proteins at DNA breaks: a focus on protein modifications. Genes Dev 2011, 25:409-433.

- Dantuma NP, van Attikum H: Spatiotemporal regulation of posttranslational modifications in the DNA damage response. EMBO J 2016, 35:6-23.
- Kolas NK, Chapman JR, Nakada S, Ylanko J, Chahwan R, Sweeney FD, Panier S, Mendez M, Wildenhain J, Thomson TM et al.: Orchestration of the DNA-damage response by the RNF8 ubiquitin ligase. Science 2007, 318:1637-1640.
- Lord CJ, Ashworth A: PARP inhibitors: synthetic lethality in the clinic. Science 2017, 355:1152-1158.
- Gupte R, Liu Z, Kraus WL: PARPs and ADP-ribosylation: recent advances linking molecular functions to biological outcomes. *Genes Dev* 2017, **31**:101-126.
- Suskiewicz MJ, Palazzo L, Hughes R, Ahel I: Progress and outlook in studying the substrate specificities of PARPs and related enzymes. FEBS J 2021, 288:2131-2142.
- 7. Talhaoui I, Lebedeva NA, Zarkovic G, Saint-Pierre C, Kutuzov MM, Sukhanova MV, Matkarimov BT, Gasparutto D, Saparbaev MK, Lavrik OI, Ishchenko AA: Poly(ADP-ribose) polymerases covalently modify strand break termini in DNA fragments in vitro. Nucleic Acids Res 2016, 44:9279-9295

The study showed that PARP1 and PARP2 can modify DNA ends in vitro.

- Munnur D, Ahel I: Reversible mono-ADP-ribosylation of DNA breaks. FEBS J 2017, 284:4002-4016.
- Munnur D, Bartlett E, Mikolčević P, Kirby IT, Rack JGM, Mikoč A, Cohen MS, Ahel I: Reversible ADP-ribosylation of RNA. Nucleic Acids Res 2019, 47:5658-5669.
- 10. Groslambert J, Prokhorova E, Ahel I: ADP-ribosylation of DNA and RNA. DNA Repair 2021, 105 http://dx.doi.org/10.1016/j. dnarep.2021.103144.
- Leidecker O, Bonfiglio JJ, Colby T, Zhang Q, Atanassov I, Zaja R,
   Palazzo L, Stockum A, Ahel I, Matic I: Serine is a new target residue for endogenous ADP-ribosylation on histones. Nat

Chem Biol 2016, 12:998-1000 The paper described for the first time DNA damage-dependent ADPribosylation of histone proteins on serine residues.

- Bonfiglio JJ, Fontana P, Zhang Q, Colby T, Gibbs-Seymour I,
   Atanassov I, Bartlett E, Zaja R, Ahel I, Matic I: Serine ADP-ribosylation depends on HPF1. Mol Cell 2017, 65:932-940 ADP-ribosylation on serine residues involves many other proteins in response to DNA damage beyond histone proteins.
- 13. Hendriks IA, Larsen SC, Nielsen ML: An advanced strategy for comprehensive profiling of ADP-ribosylation sites using mass spectrometry-based proteomics. Mol Cell Proteomics 2019,

The paper showed that one third of the human nuclear proteome in presence of DNA damage is subject to ADP-ribosylation mainly on serine

- 14. Dasovich M, Beckett MQ, Bailey S, Ong SE, Greenberg MM, Leung AKL: Identifying poly(ADP-ribose)-binding proteins with photoaffinity-based proteomics. J Am Chem Soc 2021, **143**:3037-3042.
- 15. Aberle L, Krüger A, Reber JM, Lippmann M, Hufnagel M, Schmalz M, Trussina IREA, Schlesiger S, Zubel T, Schütz K et al.: PARP1 catalytic variants reveal branching and chain lengthspecific functions of poly(ADP-ribose) in cellular physiology and stress response. Nucleic Acids Res 2020, 48:10015-10033
- 16. Krietsch J, Rouleau M, Pic É, Ethier C, Dawson TM, Dawson VL Masson JY, Poirier GG, Gagné JP: Reprogramming cellular events by poly(ADP-ribose)-binding proteins. Mol Aspects Med 2013, 34:1066-1087.
- 17. Komander D, Rape M: The ubiquitin code. Annu Rev Biochem 2012, **81**:203-229.
- 18. Gibson BA, Conrad LB, Huang D, Kraus WL: Generation and characterization of recombinant antibody-like ADP-ribose binding proteins. Biochemistry 2017, 56:6305-6316.
- Bonfiglio JJ, Leidecker O, Dauben H, Longarini EJ, Colby T, San Segundo-Acosta P, Perez KA, Matic I: An HPF1/PARP1-based chemical biology strategy for exploring ADP-ribosylation. Cell 2020, 183:1086-1102.e23

The paper described novel antibody-based tools to investigate protein serine ADP-ribosylation and demonstrated the prevalence of MARylation

- Vyas S, Matic I, Uchima L, Rood J, Zaja R, Hay RT, Ahel I, Chang P: Family-wide analysis of poly(ADP-ribose) polymerase activity. Nat Commun 2014, 5:4426.
- 21. Langelier MF, Eisemann T, Riccio AA, Pascal JM: PARP family enzymes: regulation and catalysis of the poly(ADP-ribose) posttranslational modification. Curr Opin Struct Biol 2018, **53**:187-198.
- 22. Hoang SM, Kaminski N, Bhargava R, Barroso-González J, Lynskey ML, García-Expósito L, Roncaioli JL, Wondisford AR, Wallace CT, Watkins SC et al.: Regulation of ALT-associated homology-directed repair by polyADP-ribosylation. Nat Struct Mol Biol 2020, 27:1152-1164.
- 23. Wu W, Hill SE, Nathan WJ, Paiano J, Callen E, Wang D, Shinoda K, van Wietmarschen N, Colón-Mercado JM, Zong D et al.: Neuronal enhancers are hotspots for DNA single-strand break repair. Nature 2021, 593:440-444.
- Prokhorova E, Agnew T, Wondisford AR, Tellier M, Kaminski N, Beijer D, Holder J, Groslambert J, Suskiewicz MJ, Zhu K et al.: Unrestrained poly-ADP-ribosylation provides insights into chromatin regulation and human disease. Mol Cell 2021,

81:2640-2655.e8

Synergism between PARG and ARH3 to unleash widespread, and ultimately toxic. PARvlation in response to DNA damage.

- Hanzlikova H, Gittens W, Krejcikova K, Zeng Z, Caldecott KW: Overlapping roles for PARP1 and PARP2 in the recruitment of endogenous XRCC1 and PNKP into oxidized chromatin. Nucleic Acids Res 2017, 45:2546-2557.
- Ménissier de Murcia J, Ricoul M, Tartier L, Niedergang C, Huber A, Dantzer F, Schreiber V, Amé JC, Dierich A, LeMeur M et al.: Functional interaction between PARP-1 and PARP-2 in chromosome stability and embryonic development in mouse. EMBO J 2003. 22:2255-2263.
- 27. Langelier MF, Planck JL, Roy S, Pascal JM: Structural basis for
   DNA damage-dependent poly(ADP-ribosyl)ation by human PARP-1. Science 2012, 336:728-732

This seminal paper described the crystal structure of a large portion of PARP1 in complex with DNA.

- 28. Eustermann S, Wu WF, Langelier MF, Yang JC, Easton LE, Riccio AA, Pascal JM, Neuhaus D: Structural basis of detection and signaling of DNA single-strand breaks by human PARP-1.
- Mol Cell 2015, 60:742-754 An NMR structural study showing the involvement of PARP1 zinc fingers in recognising a single-strand break.
- Rudolph J, Mahadevan J, Dyer P, Luger K: Poly(ADP-ribose) polymerase 1 searches DNA via a 'monkey bar' mechanism. el ife 2018. 7:e37818.
- 30. Zahradka P, Ebisuzaki K: A shuttle mechanism for DNA-protein interactions. The regulation of poly(ADP-ribose) polymerase. Eur J Biochem 1982, 127:579-585.
- 31. Krüger A, Bürkle A, Hauser K, Mangerich A: Real-time monitoring of PARP1-dependent PARylation by ATR-FTIR spectroscopy. Nat Commun 2020, 11:2174.
- 32. Juhász S, Smith R, Schauer T, Spekhardt D, Mamar H, Zentout S, Chapuis C, Huet S, Timinszky G: The chromatin remodeler ALC1 underlies resistance to PARP inhibitor treatment. Sci Adv 2020, 6:eabb8626.
- 33. Hewitt G, Borel V, Segura-Bayona S, Takaki T, Ruis P, Bellelli R, Lehmann LC, Sommerova L, Vancevska A, Tomas-Loba A et al.: Defective ALC1 nucleosome remodeling confers PARPi sensitization and synthetic lethality with HRD. Mol Cell 2021, 81:767-783.e11.
- 34. Demin AA, Hirota K, Tsuda M, Adamowicz M, Hailstone R, Brazina J, Gittens W, Kalasova I, Shao Z, Zha S *et al.*: XRCC1 prevents toxic PARP1 trapping during DNA base excision repair. Mol Cell 2021, 81:3018-3030.
- Prokhorova E, Zobel F, Smith R, Zentout S, Gibbs-Seymour I, Schutzenhofer K, Peters A, Groslambert J, Zorzini V, Ágnew T

et al.: Serine-linked PARP1 auto-modification controls PARP inhibitor response. Nat Commun 2021, 12:4055.

 Obaji E, Haikarainen T, Lehtiö L: Structural basis for DNA break
 recognition by ARTD2/PARP2. Nucleic Acids Res 2018, 46:12154-12165

A crystal structure demonstrating for the first time the potential of the PARP2 WGR domain to hold together two DNA ends.

 Bilokapic S, Suskiewicz MJ, Ahel I, Halic M: Bridging of DNA
 breaks activates PARP2-HPF1 to modify chromatin. Nature 2020, 585:609-613

Cryo-EM structure showing the activated HPF1-PARP2 complex bridging a DNA break in the context of nucleosomes.

- Gaullier G, Roberts G, Muthurajan UM, Bowerman S, Rudolph J, Mahadevan J, Jha A, Rae PS, Luger K: Bridging of nucleosomeproximal DNA double-strand breaks by PARP2 enhances its interaction with HPF1. PLoS One 2020, 15:e0240932.
- Obaji E, Maksimainen MM, Galera-Prat A, Lehtiö L: Activation of PARP2/ARTD2 by DNA damage induces conformational changes relieving enzyme autoinhibition. Nat Commun 2021, 12:3479.
- 40. Haince JF, McDonald D, Rodrigue A, Déry U, Masson JY, Hendzel MJ, Poirier GG: PARP1-dependent kinetics of recruitment of MRE11 and NBS1 proteins to multiple DNA damage sites. *J Biol Chem* 2008, 283:1197-1208.
- Chen Q, Kassab MA, Dantzer F, Yu X: PARP2 mediates branched poly ADP-ribosylation in response to DNA damage. Nat Commun 2018, 9:3233.
- 42. Dawicki-McKenna JM, Langelier MF, DeNizio JE, Riccio AA,
   Cao CD, Karch KR, McCauley M, Steffen JD, Black BE, Pascal JM: PARP-1 activation requires local unfolding of an autoinhibitory domain. Mol Cell 2015, 60:755-768

The paper described that the remodelling of an autoinhibitory fragment of the PARP1 catalytic domain, called the helical domain (HD), inhibits NAD+binding.

43. Palazzo L, Leidecker O, Prokhorova E, Dauben H, Matic I, Ahel I:
 Serine is the major residue for ADP-ribosylation upon DNA damage. eLife 2018, 7:e34334

The paper showed that ADP-ribosylation in response to DNA damage is mainly serine-linked and depends on HPF1 and ARH3 in cells.

44. Larsen SC, Hendriks IA, Lyon D, Jensen LJ, Nielsen ML: Systemswide analysis of serine ADP-ribosylation reveals widespread occurrence and site-specific overlap with phosphorylation. Cell Rep 2018, 24:2493-2505.e4

A mass spectrometric analysis demonstrating the prevalence of serine ADP-ribosylation and its potential cross-talk with phosphorylation.

- Buch-Larsen SC, Hendriks IA, Lodge JM, Rykær M, Furtwängler B, Shishkova E, Westphall MS, Coon JJ, Nielsen ML: Mapping physiological ADP-ribosylation using activated ion electron transfer dissociation. Cell Rep 2020, 32:108176.
- 46. Bartlett E, Bonfiglio JJ, Prokhorova E, Colby T, Zobel F, Ahel I,
  Matic I: Interplay of histone marks with serine ADP-ribosylation. Cell Rep 2018, 24:3488-3502.e5

The study demonstrates that serine ADP-ribosylation interferes with some canonical histone markers and *vice versa*.

 47. Liszczak G, Diehl KL, Dann GP, Muir TW: Acetylation blocks DNA
 damage-induced chromatin ADP-ribosylation. Nat Chem Biol 2018, 14:837-840

First demonstration of a cross-talk between histone H3 serine ADP-ribosylation and acetylation.

- 48. Gibbs-Seymour I, Fontana P, Rack JGM, Ahel I: HPF1/C4orf27 is
- a PARP-1-interacting protein that regulates PARP-1 ADPribosylation activity. Mol Cell 2016, 62:432-442

Discovery of HPF1 as a PARP1-interacting protein involved in DNA damage response.

49. Suskiewicz MJ, Zobel F, Ogden TEH, Fontana P, Ariza A, Yang JC,
 Thu K, Bracken L, Hawthorne WJ, Ahel D et al.: HPF1 completes the PARP active site for DNA damage-induced ADP-

the PARP active site for DNA damage-induced ADPribosylation. Nature 2020, 579:598-602
HPF1 completes the PARP2 catalytic domain. This paper describes the molecular details of serine ADP-ribosylation.

- Sun FH, Zhao P, Zhang N, Kong LL, Wong CCL, Yun CH: HPF1 remodels the active site of PARP1 to enable the serine ADPribosylation of histones. Nat Commun 2021, 12:1028.
- 51. Rudolph J, Roberts G, Muthurajan UM, Luger K: HPF1 and
   nucleosomes mediate a dramatic switch in activity of PARP1 from polymerase to hydrolase. eLife 2021, 10:e65773

This study highlights the importance of the HPF1 residue E284 for serine ADP-ribosylation catalysis and demonstrates the puzzling HPF1-dependent switch of PARP1 into an NAD<sup>+</sup> hydrolase.

- 52. Marsischky GT, Wilson BA, Collier RJ: Role of glutamic acid 988 of human poly-ADP-ribose polymerase in polymer formation. Evidence for active site similarities to the ADP-ribosylating toxins. *J Biol Chem* 1995, 270:3247-3254.
- Kleine H, Poreba E, Lesniewicz K, Hassa PO, Hottiger MO, Litchfield DW, Shilton BH, Lüscher B: Substrate-assisted catalysis by PARP10 limits its activity to mono-ADPribosylation. Mol Cell 2008, 32:57-69.
- Chapman JD, Gagné JP, Poirier GG, Goodlett DR: Mapping PARP-1 auto-ADP-ribosylation sites by liquid chromatography-tandem mass spectrometry. J Proteome Res 2013, 12:1868-1880.
- Chen SH, Yu X: Targeting dePARylation selectively suppresses DNA repair-defective and PARP inhibitor-resistant malignancies. Sci Adv 2019, 5:eaav4340.
- Brochu G, Duchaine C, Thibeault L, Lagueux J, Shah GM, Poirier GG: Mode of action of poly(ADP-ribose) glycohydrolase. Biochim Biophys Acta 1994, 1219:342-350.
- 57. Fontana P, Bonfiglio JJ, Palazzo L, Bartlett E, Matic I, Ahel I: Serine
  ADP-ribosylation reversal by the hydrolase ARH3. *eLife* 2017, 6:e28533

The paper discovered ARH3 as the main enzyme responsible for terminal removal of ADP-ribosylation on serine residues.

- Koh DW, Lawler AM, Poitras MF, Sasaki M, Wattler S, Nehls MC, Stöger T, Poirier GG, Dawson VL, Dawson TM: Failure to degrade poly(ADP-ribose) causes increased sensitivity to cytotoxicity and early embryonic lethality. Proc Natl Acad Sci U S A 2004, 101:17699-17704.
- Wang Y, Kim NS, Haince JF, Kang HC, David KK, Andrabi SA, Poirier GG, Dawson VL, Dawson TM: Poly(ADP-ribose) (PAR) binding to apoptosis-inducing factor is critical for PAR polymerase-1-dependent cell death (parthanatos). Sci Signal 2011. 4:ra20.
- Hanzlikova H, Prokhorova E, Krejcikova K, Cihlarova Z, Kalasova I, Kubovciak J, Sachova J, Hailstone R, Brazina J, Ghosh S et al.: Pathogenic ARH3 mutations result in ADP-ribose chromatin scars during DNA strand break repair. Nat Commun 2020, 11:3391.
- Malanga M, Althaus FR: Poly(ADP-ribose) reactivates stalled DNA topoisomerase I and induces DNA strand break resealing. J Biol Chem 2004, 279:5244-5248.
- Fahrer J, Kranaster R, Altmeyer M, Marx A, Bürkle A: Quantitative analysis of the binding affinity of poly(ADP-ribose) to specific binding proteins as a function of chain length. Nucleic Acids Res 2007, 35:e143.
- Langelier MF, Riccio AA, Pascal JM: PARP-2 and PARP-3 are selectively activated by 5' phosphorylated DNA breaks through an allosteric regulatory mechanism shared with PARP-1. Nucleic Acids Res 2014, 42:7762-7775.
- Kim MY, Mauro S, Gévry N, Lis JT, Kraus WL: NAD\*-dependent modulation of chromatin structure and transcription by nucleosome binding properties of PARP-1. Cell 2004, 119:803-814.
- 65. Muthurajan UM, Hepler MR, Hieb AR, Clark NJ, Kramer M, Yao T, Luger K: Automodification switches PARP-1 function from chromatin architectural protein to histone chaperone. *Proc Natl Acad Sci U S A* 2014, **111**:12752-12757.
- Satoh MS, Lindahl T: Role of poly(ADP-ribose) formation in DNA repair. Nature 1992, 356:356-358.

- 67. Ahel I, Ahel D, Matsusaka T, Clark AJ, Pines J, Boulton SJ, West SC: Poly(ADP-ribose)-binding zinc finger motifs in DNA repair/checkpoint proteins. Nature 2008, 451:81-85.
- **68.** Ahel D, Horejsí Z, Wiechens N, Polo SE, Garcia-Wilson E, Ahel I, Flynn H, Skehel M, West SC, Jackson SP *et al.*: **Poly(ADP-ribose)**dependent regulation of DNA repair by the chromatin remodeling enzyme ALC1. Science 2009, 325:1240-1243.
- 69. Sellou H, Lebeaupin T, Chapuis C, Smith R, Hegele A, Singh HR, Kozlowski M, Bultmann S, Ladurner AG, Timinszky G, Huet S: The poly(ADP-ribose)-dependent chromatin remodeler Alc1 induces local chromatin relaxation upon DNA damage. Mol Biol Cell 2016, 27:3791-3799.
- Kumamoto S, Nishiyama A, Chiba Y, Miyashita R, Konishi C, Azuma Y, Nakanishi M: HPF1-dependent PARP activation promotes LIG3-XRCC1-mediated backup pathway of Okazaki fragment ligation. Nucleic Acids Res 2021, 21:5003-5016.
- 71. Rack JGM, Palazzo L, Ahel I: (ADP-ribosyl)hydrolases: structure, function, and biology. Genes Dev 2020, 34:263-284.
- 72. Lin W, Amé JC, Aboul-Ela N, Jacobson EL, Jacobson MK: Isolation and characterization of the cDNA encoding bovine poly(ADP-ribose) glycohydrolase. J Biol Chem 1997, **272**:11895-11901.
- 73. Slade D, Dunstan MS, Barkauskaite E, Weston R, Lafite P, Dixon N, Ahel M, Leys D, Ahel I: The structure and catalytic mechanism of a poly(ADP-ribose) glycohydrolase. Nature 2011, 477:616-
- 74. Barkauskaite E, Brassington A, Tan ES, Warwicker J, Dunstan MS, Banos B, Lafite P, Ahel M, Mitchison TJ, Ahel I, Leys D: Visualization of poly(ADP-ribose) bound to PARG reveals inherent balance between exo- and endo-glycohydrolase activities. Nat Commun 2013, 4:2164.

75. Gogola E, Duarte AA, de Ruiter JR, Wiegant WW, Schmid JA, de Bruijn R, James DI, Guerrero Llobet S, Vis DJ, Annunziato S et al.: Selective loss of PARG restores PARylation and counteracts PARP inhibitor-mediated synthetic lethality. Cancer Cell 2018, 33:1078-1093.e12

Loss of PARG activity is a major resistance mechanism to PARP inhibitors.

76. Rudolph J, Roberts G, Luger K: Histone parylation factor 1 contributes to the inhibition of PARP1 by cancer drugs. Nat Commun 2021, 12:736

A novel. sensitive PARP inhibitor binding assay reveals an influence of HPF1 on the binding of some inhibitors.

Zandarashvili L, Langelier MF, Velagapudi UK, Hancock MA, Steffen JD, Billur R, Hannan ZM, Wicks AJ, Krastev DB, Pettitt SJ et al.: Structural basis for allosteric PARP-1 retention on DNA breaks. Science 2020, 368:eaax6367

Demonstration of the potential for allosteric enhancement of PARP1 retention by inhibitors.

- Pillay N, Tighe A, Nelson L, Littler S, Coulson-Gilmer C, Bah N, Golder A, Bakker B, Spierings DCJ, James DI et al.: DNA
- replication vulnerabilities render ovarian cancer cells sensitive to poly(ADP-Ribose) glycohydrolase inhibitors. Cancer Cell 2019, 35:519-533.e8

The paper reveals the rationale for employing PARG inhibitor as a novel therapeutic strategy for PARPi-resistant ovarian cancer cells.

- Blessing C, Mandemaker IK, Gonzalez-Leal C, Preisser J, Schomburg A, Ladurner AG: The oncogenic helicase ALC1 regulates PARP inhibitor potency by trapping PARP2 at DNA breaks. Mol Cell 2020, 80:862-875.e6.
- 80. Verma P, Zhou Y, Cao Z, Deraska PV, Deb M, Arai E, Li W, Shao Y, Puentes L, Li Y et al.: ALC1 links chromatin accessibility to PARP inhibitor response in homologous recombinationdeficient cells. Nat Cell Biol 2021, 23:160-171.