XRCC1 protects transcription from toxic PARP1 activity during DNA base excision repair

Marek Adamowicz, Richard Hailstone, Annie A. Demin, Emilia Komulainen, Hana Hanzlíková, Jan Brazina, Amit Gautam, Sophie E. Wells and Keith W. Caldecott

Genetic defects in the repair of DNA single-strand breaks (SSBs) can result in neurological disease triggered by toxic activity of the single-strand-break sensor protein PARP1. However, the mechanism(s) by which this toxic PARP1 activity triggers cellular dysfunction are unclear. Here we show that human cells lacking XRCC1 fail to rapidly recover transcription following DNA base damage, a phenotype also observed in patient-derived fibroblasts with XRCC1 mutations and Xrcc1<sup>−/−</sup> mouse neurons. This defect is caused by excessive/aberrant PARP1 activity during DNA base excision repair, resulting from the loss of PARP1 regulation by XRCC1. We show that aberrant PARP1 activity suppresses transcriptional recovery during base excision repair by promoting excessive recruitment and activity of the ubiquitin protease USP3, which as a result reduces the level of monoubiquitinated histones important for normal transcriptional regulation. Importantly, inhibition and/or deletion of PARP1 or USP3 restores transcriptional recovery in XRCC1<sup>−/−</sup> cells, highlighting PARP1 and USP3 as possible therapeutic targets in neurological disease.

**Results**

XRCC1 promotes the recovery of transcription following oxidative damage. Given that SSBs can slow or block the physical progression of RNA polymerases<sup>20–22</sup>, we wondered whether SSB repair-defective cells exhibit defects in transcription following DNA damage. We therefore examined the impact of XRCC1 deletion on global transcription in human RPE-1 cells following treatment with H<sub>2</sub>O<sub>2</sub> (a physiologically relevant source of oxidative SSBs) by pulse labelling with 5-ethynyluridine (EU). We observed a robust inhibition of global transcription immediately after treatment with H<sub>2</sub>O<sub>2</sub> in both wild-type (WT) and XRCC1<sup>−/−</sup> RPE-1 cells (Fig. 1a,b). More importantly, whereas the levels of transcription recovered significantly within 2h following H<sub>2</sub>O<sub>2</sub> treatment in the WT RPE-1 cells, they declined further in the XRCC1<sup>−/−</sup> cells (Fig. 1a,b). The lack of transcription recovery in XRCC1<sup>−/−</sup> cells was accompanied by a progressive loss of RNA polymerase I (RNAPI) (RPAl) treated with H<sub>2</sub>O<sub>2</sub>. RNAPI I (RPAl) is a nucleolar enzyme that is involved in the transcription of ribosomal RNA. Its loss indicates that XRCC1<sup>−/−</sup> cells are unable to recover transcription, as evidenced by the lack of RNAPI I (RPAl) recovery (Fig. 1a,c). Given that we did not detect a reduction in the global levels of RNA polymerase II (RNAPII) in the XRCC1<sup>−/−</sup> cells, this most probably reflected the dissociation of RNAPI from ribosomal RNA transcription rather than degradation of the RNAPI (Extended Data Fig. 1a). Although most of the EU signal detected in our pulse-labelling experiments was nucleolar, the recovery of EU incorporation in the XRCC1<sup>−/−</sup> RPE-1 cells was also reduced in the nucleoplasm (Fig. 1a), suggesting that RNA polymerase II (RNAPII)-dependent transcription was impaired in XRCC1<sup>−/−</sup> cells.

**Discussion**

XRCC1 is a scaffold protein that is recruited to SSBs by PARP1 and/or PARP2 activity through direct interaction between poly(ADP-ribose) and the central BRCT domain of XRCC1 (refs. 4,5). XRCC1 in turn interacts with and recruits the enzymes required for SSB repair, including DNA polymerase β<sup>4,13</sup>, DNA ligase III<sup>13</sup>, poly-nucleotide kinase/phosphatase<sup>14,15</sup> and aprataxin<sup>16,17</sup>. Hereditary mutations in XRCC1 and some of these protein partners result in neurodevelopmental disorders and/or progressive neurodegeneration<sup>24–25</sup>. Intriguingly, the deletion and/or inhibition of PARP1 greatly reduces or prevents these pathologies in an Xrccl-defective mouse model of SSB-associated neurological disease, highlighting excessive/aberrant PARP1 activity as a source of SSB-induced neuropathology<sup>16,19</sup>. However, the molecular mechanism by which excessive/aberrant PARP1 activity triggers cellular dysfunction remains unclear.

Here we show that prolonged PARP1 activity at un repaired DNA base damage suppresses the recovery of transcription by promoting excessive recruitment and activity of the ubiquitin-specific protease USP3. Remarkably, excessive USP3 activity reduces global levels of monoubiquitinated histones important for transcriptional regulation, such as histone H2A and H2B monoubiquitination at K119 and K120 (H2AmUb and H2BmUb, respectively). Importantly, either PARP1 or USP3 inhibition/deletion rescue normal levels of histone monoubiquitination and transcription recovery following DNA base damage, highlighting these enzymes as possible therapeutic targets in the treatment of base excision repair (BER)-defective neurological disease.

**Conclusion**

XRCC1 promotes the recovery of transcription following oxidative damage. The molecular mechanism by which this takes place is likely to involve the regulation of PARP1 and USP3 activity during DNA base excision repair. Future studies should focus on understanding the specific role of XRCC1 in transcriptional recovery and identifying the mechanisms by which PARP1 activity is regulated in this process.

**Conflict of interest**

The authors declare no conflict of interest.

**Acknowledgments**

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transcription was also affected. To confirm this, we compared the levels of hyperphosphorylated RNAPII, which is an established measure of the RNAPII transcriptional activity, in WT and XRCC1−/− RPE-1 cells before and after H2O2 treatment. The level of RNAPII hyperphosphorylation was markedly reduced in the XRCC1−/− RPE-1 cells following H2O2 treatment, an observation we verified with antibodies specific to two phosphorylated sites (S2 and S5) within the carboxy (C)-terminal domain of RNAPII that are associated with transcriptional activity (Fig. 1d)25. The loss of hyperphosphorylated RNAPII seemed to involve, at least partly, proteolysis of RNAPII, given that incubation with proteasome inhibitors restored normal levels of the hyperphosphorylated RNAPI (Extended Data Fig. 1b). However, proteasome inhibitors did not restore normal levels of transcription recovery, indicating that degradation of hyperphosphorylated RNAPII was a result of failed transcription recovery rather than a cause (Extended Data Fig. 1b–e). Importantly, the defect in transcription recovery was neither limited to RPE-1 cells nor an off-target effect of gene editing because we observed similar results in XRCCI−/− U2OS cells and the defect in transcriptional recovery in these cells was corrected by stable expression of recombinant human XRCC1 (Extended Data Fig. 2a–d). Collectively, these data indicate that XRCC1 is required for the rapid recovery of transcription by both RNAPI and RNAPII following oxidative stress.

Reduced transcriptional recovery in XRCC1-deficient cells and neurons is a result of PARP1. Hereditary mutations in human XRCC1 result in cerebellar ataxia, a phenotype that is also observed in Xrc1lox/− mice in which Xrc1 is conditionally deleted in the brain18,19,26. This phenotype is largely rescued by the deletion of Parp1, the primary sensor of unrepaired SSBs in mammalian cells18,19. We therefore examined the impact of PARP1 deletion on the recovery of transcription in XRCCI−/− cells. Strikingly, PARP1 deletion rescued the transcriptional recovery in XRCCI−/− RPE-1 cells, as measured by EU pulse labelling, RNAPI foci and RNAPII phosphorylation (Fig. 2a–d). In addition, transcription recovery was rescued in both XRCCI−/− RPE-1 and U2OS cells by incubation with PARP inhibitor (PARPI), confirming that the transcription defect was the result of toxic/aberrant ADP-ribosylation (Fig. 2e and Extended Data Fig. 3). Consistent with this idea, the defect observed in the XRCCI−/− cells was recapitulated in WT RPE-1 cells by incubation with PARP inhibitor (PARPI), confirming that the transcription defect was the result of toxic/aberrant ADP-ribosylation (Fig. 2e and Extended Data Fig. 3). Consistent with this idea, the defect observed in the XRCCI−/− cells was recapitulated in WT RPE-1 cells by incubation with PARP inhibitor (PARPI), confirming that the transcription defect was the result of toxic/aberrant ADP-ribosylation (Fig. 2e and Extended Data Fig. 3). Consistent with this idea, the defect observed in the XRCCI−/− cells was recapitulated in WT RPE-1 cells by incubation with PARP inhibitor (PARPI), confirming that the transcription defect was the result of toxic/aberrant ADP-ribosylation (Fig. 2e and Extended Data Fig. 3). Consistent with this idea, the defect observed in the XRCCI−/− cells was recapitulated in WT RPE-1 cells by incubation with PARP inhibitor (PARPI), confirming that the transcription defect was the result of toxic/aberrant ADP-ribosylation (Fig. 2e and Extended Data Fig. 3). Consistent with this idea, the defect observed in the XRCCI−/− cells was recapitulated in WT RPE-1 cells by incubation with PARP inhibitor (PARPI), confirming that the transcription defect was the result of toxic/aberrant ADP-ribosylation (Fig. 2e and Extended Data Fig. 3).
The poly(ADP-ribose)-binding activity of XRCC1 promotes transcriptional recovery by directly regulating PARP1 activity. To understand how XRCC1 prevents toxic PARP1 activity, we identified the protein domain/s important for this process. A truncated Myc-histidine (Myc-His)-tagged fragment of XRCC1 encoding only the central one-third of the protein (Myc-His–XRCC1 161–406) rescued transcription recovery in XRCC1−/− U2OS cells (Fig. 4a,b and Extended Data Fig. 5a–e). This was surprising because this fragment lacks the protein-interaction domains that underpin the role of XRCC1 as a molecular scaffold and so cannot support normal rates of SSB repair (Extended Data Fig. 5f,g). The ability of XRCC1 to bind directly to poly(ADP-ribose) was required to prevent PARP1-mediated transcriptional suppression, as indicated by the failure of a mutant derivative of Myc-His–XRCC1 161–406 harbouning mutations that prevent binding to this polymer (R335A/K369A; denoted Myc-His–XRCC1 161–406 R335A/K369A) to rescue transcriptional recovery, thereby suggesting that XRCC1 might regulate poly(ADP-ribose) synthesis directly (Fig. 4b and Extended Data Fig. 5a–e). The most probable target of this regulation is PARP1 itself, given that this protein was the most abundant, if not the only, ribosylated protein detected in Myc-His–XRCC1 161–406 immunoprecipitates following H2O2 treatment and its recovery was prevented by the mutations that prevent poly(ADP-ribose) binding (Fig. 4c). Consistent with direct regulation of PARP1 activity by the poly(ADP-ribose)-binding domain of XRCC1, both recombinant full-length XRCC1 and XRCC1 161–406 reduced PARP1 auto-ribosylation in vitro, whereas the mutant derivatives of these proteins that cannot bind poly(ADP-ribose) were unable to do so (Fig. 4d).

These data suggest that XRCC1 can suppress aberrant PARP1 activity not only by accelerating SSB repair but also by direct binding to, and suppression of, poly(ADP-ribose) chain extension. To confirm this idea, we examined whether transcriptional recovery could be rescued by the overexpression of APLF, a structurally distinct poly(ADP-ribose)-binding protein that can also negatively regulate poly(ADP-ribose) synthesis. Recombinant APLF suppressed poly(ADP-ribose) chain extension in vitro, albeit less effectively than XRCC1, and overexpression of high levels of APLF restored transcriptional recovery in XRCC1−/− U2OS cells in a manner dependent on the poly(ADP-ribose)-binding zinc finger domain (Fig. 4e,f and Extended Data Fig. 5h–k). These data indicate that the negative regulation of PARP1 activity by direct binding of XRCC1 to poly(ADP-ribose) is both required and sufficient for normal transcription recovery following oxidative stress.

Prolonged transcriptional suppression is triggered by low but persistent PARP1 activity at unrepaired DNA BER intermediates. Collectively, the experiments described so far indicate that although the initial suppression of transcription by H2O2 is independent of PARP1 activity and probably reflects the stalling/inefficient bypass of DNA lesions by RNAPs, the prolonged suppression observed in XRCC1−/− cells is imposed by excessive/aberrant PARP1 activity at persistent SSBs. To address this further we conducted time-course experiments to determine when and for how long transcriptional suppression occurs. Whereas WT RPE-1 cells continued to recover transcription from 2 h following H2O2 treatment, with the transcription levels restored to normal within 24 h, XRCC1−/− cells exhibited little or no transcriptional recovery for at least 9 h following H2O2 treatment (Fig. 5a and Extended Data Fig. 6a,b). Moreover, incubation with PARPi for 1 h rescued transcription even if applied 8 h after the H2O2 treatment, indicating that the failure to recover transcription in the XRCC1−/− cells was a result of prolonged and ongoing ADP-ribosylation (Fig. 5a and Extended Data Fig. 6a,b). Consistent with this idea, although H2O2-induced protein ADP-ribosylation declined below detectable levels in both WT and XRCC1−/− cell extracts 2 h after H2O2 treatment, as measured by western blotting, we detected low but persistent levels of PARP1 auto-ribosylation in the XRCC1−/− cell extracts following enrichment with anti-ADP-ribose antibodies (Extended Data Fig. 6c,d).

We reasoned that the low level of persistent PARP1 activity in the XRCC1−/− cells most probably reflected SSBs arising as intermediates of BER, given that while SSBs induced directly by disintegration of oxidized deoxyribose arise immediately, SSBs arising during BER require the enzymatic excision of oxidized DNA bases and thus arise more slowly. Moreover, we recently showed that PARP1 can become ‘trapped’ on SSB intermediates in XRCC1−/− cells during BER and that low but persistent levels of PARP1 auto-ribosylation is a feature of this phenomenon. Consistent with this idea, short interfering RNA (siRNA)-mediated depletion of NTH1—a DNA glycosylase that excises a broad spectrum of oxidized DNA bases—rescued the recovery of transcription in H2O2-treated XRCC1−/− cells (Fig. 5b,c and Extended Data Fig. 6e). Furthermore, global transcription was substantially suppressed in XRCC1−/− cells following prolonged treatment with methyl methanesulfonate (MMS), an alkylating agent that induces SSBs largely or entirely via BER (Fig. 5d–f). This repression was also accompanied by a loss of RNAPII hyperphosphorylation, which once again was rescued by PARPi (Fig. 5i). Together, these data indicate that transcriptional recovery is suppressed in XRCC1−/− cells by low levels of persistent PARP1 activity at unrepaired BER intermediates.

**Aberrant PARP1 activity disrupts global histone monoubiquitination.** Next, we examined the mechanism/s by which aberrant PARP1 activity inhibits transcription recovery during BER. Given that PARP1 can affect transcription by ADP-ribosylating histones, we examined the impact of the deletion of histone parylation factor-1 (HPF1), a protein partner of PARP1 that is required for histone ADP-ribosylation. However, although HPF1 deletion prevented detectable histone ADP-ribosylation in XRCC1−/− U2OS cells as expected, it did not rescue transcription recovery, suggesting...
that histone ribosylation is not the cause of transcriptional suppression (Extended Data Fig. 7).

As PARP1 can also affect transcription indirectly, via the recruitment of chromatin remodelling proteins\(^4\),\(^5\), we examined whether aberrant ADP-ribosylation affected the levels of histone marks known to promote transcriptional activity. Although the global levels of methylation and acetylation on histone H3 (H3K4me3 and H3K9Ac, respectively) were slightly, albeit not statistically
significantly reduced in XRCC1−/− cells compared with WT cells 2 h after H2O2 treatment (Extended Data Fig. 8a,b), the global levels of H2AmUb and H2BmUb were dramatically reduced (Fig. 6a,b, respectively). The reduction in levels of H2BmUb was particularly noteworthy because this modification promotes transcriptional activity by a number of different mechanisms, including the direct...

Fig. 3 | Toxic PARP1 activity suppresses transcriptional recovery in patient-derived fibroblasts with XRCC1 mutations and Xrcc1-deficient mouse cerebellar neurons. a, Representative images of RNAPI foci (RPA194) showing the levels of global transcription (EU pulse labelling) in normal (1BR) and patient-derived fibroblasts with XRCC1 mutations (XD1) following mock treatment or at the indicated times after treatment with 60 μM H2O2 for 5 min. The cells were pulse labelled with EU as in Fig. 1. b, Levels of global transcription (EU immunofluorescence) from the experiment shown in a. Data are the mean ± s.e.m. of three independent experiments. Statistical significance was determined using a two-way ANOVA with Sidak’s multiple comparisons test (significantly different P values are indicated). c, Representative images of the global transcription levels (EU immunofluorescence) in WT and Xrcc1flox-cre mouse cerebellar neurons (NeuN), pretreated with PARPi, following mock treatment or at the indicated times after treatment with 250 μM H2O2 for 5 min. Representative images from one of three independent experiments are shown. Scale bars, 10 μm.
**Fig. 4 | The poly(ADP-ribose)-binding activity of XRCC1 promotes transcriptional recovery following oxidative damage.**

**a.** Cartoon of full-length (FL) and truncated XRCC1 proteins encoded by the Myc- and/or His-tagged expression constructs employed in this work. The interaction partners are shown (top): Polβ, DNA polymerase β; PAR, poly(ADP-ribose); PNKP, polynucleotide kinase phosphatase; LIG3, DNA ligase III. The N-terminal domain (NTD), nuclear localization signal (NLS) and two BRCT domains are also shown; a.a., amino acids.

**b.** Representative images showing the levels of global transcription (EU pulse labelling; bottom) in XRCC1−/− U2OS cells, transiently transfected with expression constructs encoding the indicated XRCC1 proteins, following mock treatment or 2 h after treatment with 1 mM H$_2$O$_2$ for 20 min. The XRCC1 protein levels are also shown (middle). The cells were pulse labelled with EU as in Fig. 1. EV, empty vector.

**c.** Immunoblot of Myc-tagged XRCC1 proteins, PARP1 and ADP-ribose levels in cell extract from the indicated transiently transfected XRCC1−/− U2OS cells before (left, input) and after (right) anti-Myc immunoprecipitation, 2 h after mock treatment or treatment with 1 mM H$_2$O$_2$ for 20 min (right).

**d.** XRCC1 suppresses PARP1 auto-ADP-ribosylation in vitro. Human recombinant PARP1 (100 nM) was incubated in the presence of single-stranded DNA (100 nM), NAD$^+$ (2.5 μM) and with or without 1 μM of full-length XRCC1 or the indicated XRCC1 protein fragment for 30 min. The reaction products were fractionated by SDS–PAGE and detected by western blotting with ADP-ribose detection reagent.

**e.** Representative images showing the levels of global transcription (EU pulse labelling) in XRCC1−/− U2OS cells stably transfected with expression constructs encoding WT APLF (yellow fluorescent protein (YFP)–APLF<sub>WT</sub>) or its ADP-ribose binding mutant (YFP–APLF<sub>ZFD</sub>) following mock treatment or 2 h after treatment with 1 mM H$_2$O$_2$ for 20 min. White and yellow circles indicate examples of cells over-expressing or not YFP–APLF, respectively. For b, e, representative images from one of three independent experiments are shown. Scale bars, 10 μm.

**f.** Representative blots from one of three independent experiments are shown.
relaxation of chromatin structure, facilitation of RNAP elongation and modulation of the activity of different transcriptional regulators. Similar results were observed following MMS treatment, thereby confirming that the loss of histone monoubiquitination in XRCC1−/− cells was a result of DNA base damage (Fig. 6c, d). Moreover, normal levels of histone ubiquitination were restored in
Fig. 6 | Toxic PARP1 activity during BER disrupts global histone monoubiquitination. a, Representative images (left) and levels (right) of H2BmUB in WT and XRCC1−− cells following mock treatment or at the indicated times after treatment with 250 μM H2O2 for 5 min. b, Representative images (left) and levels (right) of H2AmUb in WT and XRCC1−− cells following mock treatment or at the indicated times after treatment with 250 μM H2O2 for 5 min. a, b. The cells were incubated with DMSO vehicle or 10 μM PARPi for 1 h before, during and following H2O2 treatment as indicated. c, Representative images (left) and levels (right) of H2BmUB in WT and XRCC1−− cells following mock treatment or at the indicated times after treatment with 0.1 mg ml−1 MMS (continuous treatment). d, Representative images (left) and levels (right) of H2AmUb in WT and XRCC1−− cells following mock treatment or at the indicated times after treatment with 0.1 mg ml−1 MMS (continuous treatment). c, d. The cells were incubated with DMSO vehicle or 10 μM PARPi during the MMS treatment. a–d. Data are the mean ± s.e.m. of five (a), three (b, d) and four (c) independent experiments. Statistical significance was determined using a two-way ANOVA with Tukey’s multiple comparisons test (significantly different P values are indicated). Scale bars, 10 μm.

XRCC1−− cells by incubation with PARPi, demonstrating that the loss of these important histones modifications was the result of aberrant ADP-ribosylation (Fig. 6a–d). This impact of aberrant PARP activity on histone ubiquitination was particularly pronounced given that we did not detect a reduction in total chromatin ubiquitination, as detected by the anti-ubiquitin antibody FK2 (Extended Data Fig. 8c).

PARP1 suppresses histone monoubiquitination and transcription recovery by triggering aberrant recruitment of the ubiquitin protease USP3. Finally, to identify the mechanism by which aberrant PARP1 activity reduces histone monoubiquitination, we depleted enzymes that are known to deubiquitinate histone H2A and/or H2B46–48. Whereas siRNA directed against either of the deubiquitylases USP22 and USP36 failed to rescue transcription recovery in XRCC1−− cells following H2O2 treatment, siRNA to USP11 resulted in a small but significant rescue of transcription recovery (Extended Data Fig. 9a). More importantly, however, siRNA directed against USP3 resulted in almost complete rescue of transcription recovery in XRCC1−− RPE-1 cells following H2O2 treatment, whether measured by EU pulse labelling (Fig. 7a and Extended Data Fig. 9a) or RNAPII hyperphosphorylation (Fig. 7b). Consistent with these data, USP3 siRNA prevented the PARP1-dependent reduction in histone monoubiquitination following H2O2 treatment (Fig. 7c and Extended Data Fig. 9b). The USP3 siRNA also reduced the loss of histone monoubiquitination in XRCC1−− cells during continuous treatment with MMS (Fig. 7d and Extended Data Fig. 9c) throughout the 3 h time course. We also detected the accumulation of green fluorescent protein (GFP)-tagged USP3 in the chromatin of the XRCC1−− cells during MMS treatment, which was prevented by treatment with PARPi, providing support for PARP1 activity as a source of aberrant USP3 recruitment during BER (Fig. 8a). Interestingly, the amino (N)-terminal zinc finger domain of USP3 was sufficient for PARP-dependent recruitment at BER intermediates. However, we were unable to detect any evidence of direct binding of this domain to poly(ADP-ribose) (Extended Data Fig. 10a–f), which suggests that PARP1 mediates USP3 recruitment indirectly, perhaps via its well-established role in regulating chromatin compaction16,44,55.

Discussion

XRCC1 is a molecular scaffold protein that interacts with many of the enzymatic components of SSB repair47. At SSBs arising during BER, as obligate intermediates of base excision, XRCC1 interacts with and stabilizes DNA polynuclerase β and DNA ligase III46. Another critical role of XRCC1 during SSB repair is to prevent excessive activation of PARP1, which can otherwise lead to neurological dysfunction50,51. However, the mechanisms by which excessive/aberrant PARP1...
activity triggers this pathology have been unclear. One recent clue emerged from the discovery that the assembly of DNA repair protein complexes by XRCC1 is necessary to prevent excessive engagement of PARP1 on BER intermediates and consequently their reduced access and repair by other BER enzymes. The data presented in this study demonstrate that the low but persistent levels of PARP1 activity associated with this ‘trapped’ PARP1 in XRCC1-defective cells, even in the continued presence of unrepaired BER intermediates. The recovery of transcription at BER intermediates can thus be separated mechanistically from the completion of BER. This result may indicate that although SSBs may initially impede RNAP progression, these lesions can eventually be bypassed, as long as PARP1 signalling at the unrepaired SSBs is suppressed. Alternatively, the prolonged suppression of transcription by PARP1 detected in our experiments may reflect an impact on RNAP initiation and/or elongation in trans at sites distal to unrepaired SSBs.

PARP1 is a dynamic regulator of gene activity and is able to either repress or activate gene transcription in response to different physiological stimuli. PARP activity can regulate transcription in a number of ways, such as by relaxing higher-order chromatin compaction and/or regulating RNAP activity. PARP1 can also recruit polycomb and NuRD transcriptional repressor complexes, thereby facilitating the initial suppression of transcription following DNA damage—the initial reduction in EU pulse labelling observed following H2O2 treatment was not prevented by PARP1 deletion or inhibition. Thus, in our experiments the initial suppression of transcription probably reflects a PARP-independent mechanism, such as the pausing of RNAPII by BER intermediates.
A striking feature of our work is the discovery that excessive/aberrant PARP1 activity prolongs transcriptional suppression via recruitment of the ubiquitin protease USP3. We do not yet know how poly(ADP-ribose) synthesis recruits USP3 into chromatin at unrepaired BER intermediates. Although the N-terminal zinc finger of USP3 is sufficient for this recruitment, we have so far not detected binding of this domain to poly(ADP-ribose). PARP1 might therefore mediate USP3 recruitment indirectly, such as via its ability to regulate chromatin compaction. For example, the ability of poly(ADP-ribose) to relax chromatin structure directly or indirectly via the recruitment of one or more chromatin remodelling enzymes may allow the USP3 zinc finger to detect its ubiquitylated substrate. USP3 deubiquitinates multiple substrates, including the histones H2A, γH2AX and H2B, and is implicated in transcriptional suppression. Note that PARP1 inhibition prevents USP3 recruitment and rescues global transcription recovery, most probably reflecting the recruitment, extensive/excessive protein deubiquitination (including histones) at damaged and proximal/nearby undamaged sites, and prolonged complexes and transcription resumption. In XRCC1-defective cells, persistent PARP1 activity at unrepaired BER intermediates leads to aberrant USP3 activity.

These modifications are not known to regulate transcriptional activity.

USP3 also deubiquitinates H2A and H2B monoubiquitination at K119 and K120, respectively, both of which are critical for normal transcriptional control and are regulated in response to DNA damage. Of these two modifications, it is the loss of H2BmUb that is a more likely explanation for the prolonged transcriptional suppression observed in our experiments. This is because this modification promotes transcriptional activity, whereas loss of H2AmUb might dominate the transcriptional response in XRCC1-defective cells.

In our experiments but it may relate to the different mechanisms and/or kinetics by which H2AmUb and H2BmUb regulate transcription. For example, loss of H2BmUb can reduce RNAP progression directly, and thus rapidly, by affecting nucleosome unwinding mediated by the histone chaperone FACT ahead of RNAP. In contrast, loss of H2AmUb may increase gene transcription more slowly via the progressive loss of polycomb-repressive complexes from chromatin. Consistent with this idea, we did not detect changes in the levels of H3K4me3, which is a downstream target of H2AmUb, in our experiments (Extended Data Fig. 8).

H2BmUb is tightly coupled with RNAP elongation and promotes transcription in several ways, including facilitating histone
H3K4 and H3K79 methylation and cooperating with FACT. H2BmUb is also regulated in response to DNA damage. For example, H2BmUb is introduced at DNA double-strand breaks as a result of ATM-mediated phosphorylation and recruitment of the RNF20–RNF40 heterodimer, the ubiquitin E3 ligase that promotes this modification. More intriguingly, with respect to the current work, H2BmUb is deubiquitinated at sites of DNA damage that have stalled RNAP2. In yeast, H2BmUb is deubiquitinated by Ubp8 and Ubp10, the former of which is the ubiquitin protease most closely related to USP3. It is thus tempting to speculate that a similar process occurs at sites of stalled RNAP in human cells and that the prolonged transcriptional suppression by USP3 in BER-defective cells is a pathological extension of this process (Fig. 8b).

Finally, hereditary XRCC1 mutations result in cerebellar ataxia and deletion of XRCC1 in the mouse brain results in cerebellar ataxia, seizures and juvenile mortality. The impact of excessive/aberrant PARP1 activity on these phenotypes is illustrated by PARP1 inhibition and/or deletion, which alleviate the neuropathology in Xrcc1-defective neurons and mice. Xrcc1-defective neurons also exhibit a defect in pre-synaptic Ca2+ signalling, which is rescued by PARP inhibition/deletion, which in light of our recent data might reflect an impact of aberrant PARP1 activity on the expression of genes critical for normal neuronal activity.

Consistent with this idea, we recently demonstrated that neuronal enhancers are hotspots of SSB repair, most probably as a result of cytosine demethylation during the epigenetic (re)programming of neuronal gene expression; a process in which XRCC1-mediated BER is strongly implicated. It is also possible that PARP1 activity might impact on transcription in other pathophysiological conditions that induce DNA base damage, such as chronic inflammation.

Online content

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Methods

Cell lines, treatments and culture conditions. All cell lines were cultured in a low-oxygen (3%) incubator (37 °C and 5% CO₂). RPE-1 cells (American Type Culture Collection) were cultured in DMEM-F12 GlutaMAX medium containing 10% FBS supplemented with penicillin–streptomycin and l-glutamine. Primary fibroblasts from patients with XRCC1 mutations (XD1) as well as control counterparts 1BR.3 (denoted as 1BR) were cultured in MEM medium containing 15% FBS supplemented with penicillin–streptomycin and l-glutamine. ShRNA-transfected cells were selected with puromycin (2 μg/mL) and single colonies were isolated, amplified and verified by western blotting after one week. For the transient transfection experiments, U2OS cells were incubated with poly(ethyleneimine) solution (Sigma) for 6 h in serum-free medium and the cells were fixed and analysed after 24 h. HPFI1 and XRCC1 were transfected by transfection of WT U2OS or XRCC1−/− U2OS cells with a Cas9–GFP plasmid (Addgene, 48138) and plasmids USP36 or XRCC1161–406RK from pTWO-E-His–XRCC1161–406 and pTWO-E-His–XRCC1161−/−. APLF and APLF ZFD (harbouring the N-terminal His-tag) were expressed from pET16b-USP3 in E. coli strain BL21 (DE3). The cells were harvested by trypsinization and resuspended in sample buffer and heated for 5 min at 90 °C and subjected to SDS–PAGE. The proteins were transferred to nitrocellulose and immunoblotted with ADP-ribose-binding reagent (Millipore; Supplementary Table 1).

In vitro analysis of PARP1 auto-ribosylation. Recombinant high-specific-activity PARP1 (100nM; Trevigen) was either mock ribosylated in the absence of NAD+ or auto-ribosylated in the presence of 1 mM NAD+. PARP1 auto-ribosylation was performed using the indicated primers (Supplementary Table 1) and cloned in-frame with the use of EGFP. The recombinant proteins were expressed and purified using affinity chromatography and gel filtration before the buffer was exchanged to 150 mM NaCl, 50 mM Tris–HCl pH 7.5, 10% glycerol and 1 mM DTT. Aliquots were then snap-frozen in liquid nitrogen and stored at −80 °C.

Protein purification. N-terminal His-tagged APLF and APLF(1-110) (harbouring the mutations C379A and C385A, and C421A and C427A in the PAR-binding zinc fingers ZF1 and ZF2, respectively) were expressed from the pET16b-APLF and pET16b-APLF(1-110) plasmids in BL21 (DE3) E. coli (NEB) by overnight induction with 0.5 mM isopropyl β-D-1-thiogalactoside (IPTG) at 20 °C. Full-length C-terminal His-tagged XRCC1 and XRCC1Δ1–35 were expressed from pET16b-XH (ref. 1) and pET16b-XH(Δ1; ref. 1) and N-terminal His-tagged XRCC1(1–406) and XRCC1(1–406K) from pTWO-E-His–XRCC1(1–406) and pTWO-E-His–XRCC1(1–406K) (ref. 1) by induction with 1 mM IPTG in Luria–Bertani medium at 30 °C for 3 h. N-terminal His-tagged human USP3 was expressed from pET16b-USP3 in BL21 (DE3) E. coli (NEB) by overnight induction with 0.5 mM IPTG at 20 °C. The proteins were purified by metal-chelate affinity chromatography and gel filtration before the buffer was exchanged to 150 mM NaCl, 50 mM Tris–HCl pH 7.5, 10% glycerol and 1 mM DTT. Aliquots were then snap-frozen in liquid nitrogen and stored at −80 °C.

Immunofluorescence. For immunofluorescence, cells were fixed after transfection and seeded into 96-wells plates for the amplification of single-cell clones, which were subsequently validated by immunoblotting. Gene editing was confirmed by Sanger sequencing. Primary cerebellar neurons were established from seven-day-old (P7) male and female C57BL/6J and C3H/HeJ (ref. 1) mouse pups using the Papain Dissociation System (Worthington). The mice from which these neurons were obtained were maintained in accordance with the UK Animal (Scientific Procedures) Act 1986 and satisfied local institutional regulations at the University of Sussex and under the auspices of UK Home Office licence number P3CDBCBA8. The cells were seeded on poly-l-lysine-coated coverslips at 6 × 10^4 cells per well and maintained in Neurobasal A medium (Sigma) supplemented with 2% B27 (Invitrogen), 5.25 mM KCl, 0.5 mM glutamine and penicillin–streptomycin in a humidified incubator (3% O₂ and 5% CO₂ at 37 °C). The cultures were employed for transcription experiments between 4 and 11 days in vitro.

Western blotting. RPE-1 cells were harvested and lysed in TEB150 lysis buffer (50 mM HEPEs pH 7.4, 150 mM NaCl, 2 mM MgCl₂, 5 mM EGTA pH 8, 1 mM dithiothreitol (DTT), 0.5% Triton X-100 and 10% glycerol) containing protease inhibitor cocktail (Roche), phosphatase inhibitor cocktail (PhosSTOP; Sigma) and 1:1,000 benzazene (Sigma) for 45 min on ice. Lysis was stopped by the addition of SDS–PAGE sample buffer. The other samples were harvested directly in sample buffer and boiled as described for the RPE-1 cells. The protein extracts were fractionated by SDS–PAGE, transferred to nitrocellulose, blocked in 0.1% TBS–TWEEN containing 5% non-fat dried milk and incubated with the indicated appropriate antibodies (Supplementary Table 1).

ADP-ribose immunoprecipitation. Cells were harvested by trypanosinization and lysed in TEB250 lysis buffer (50 mM HEPEs pH 7.4, 250 mM NaCl, 2 mM MgCl₂, 5 mM EGTA pH 8, 1 mM DTT, 0.5% Triton X-100, 10% glycerol, protease inhibitor cocktail (Roche), phosphatase inhibitors (PhosSTOP; Sigma), 30 μM PARPi). The cells were washed once with PBS before incubation in CSK buffer (10 mM PIPES, pH 7.0, 100 mM NaCl, 300 mM sucrose, 3 mM MgCl₂, and 0.7% Triton X-100) for 3 min at room temperature. The cells were subsequently washed in PBS and fixed in 4% PFA for 10 min. Next, the cells were counterstained with DAPI, mounted in Moviol and subjected to immunofluorescence. Between 50 and 100 H2B–mCherry-expressing cells were counted on a confocal microscope.

In vitro ADP-ribose binding assay. First, 96-well plates were washed with H₂O and then incubated overnight with 0.1 μg/mL of calf thymus histones (Sigma) at

SDS–PAGE. The proteins were transferred to nitrocellulose and immunoblotted with ADP-ribose-binding reagent (Millipore; Supplementary Table 1).

DNA strand-break measurements. To measure DNA strand breaks under the conditions at which transcription is inhibited in XRCC1−/− cells, the cells were treated with 100 μM H₂O₂ for 20 min (the H₂O₂ was refreshed after 10 min) in serum-free medium at 37 °C and then in drug-free medium for 2 h (Extended Data Fig. 4a). To measure the kinetics of SSBR repair, the cells were treated with 50 μM H₂O₂ for 10 min on ice and then in drug-free medium at 37 °C for the indicated times (Extended Data Fig. 4b). The DNA strand breaks were measured using alkali comet assays, as described previously, using the Comet Assay IV software (Perceptive Instruments).
4 °C. Next, the plates were blocked with 5% BSA in PBST (PBS containing 0.1% Tween-20) for 30 min. The absorbed histones were mock ADP-ribosylated in the absence of NAD+ or ADP-ribosylated in the presence of 50 μM NAD+ (Sigma) in PARP1 reaction buffer (50 mM Tris–HCl pH 8, 0.8 mM MgCl2, 1% glycerol and 1.5 mM DTT) containing 40 nM single-stranded oligodeoxyribonucleotide (5′-CATATGCGGGGATCCGCTCC-3′) and 10 nM high-specific-activity PARP1 (Trevigen) for 2 h. After subsequent washes with PBST, 500 nM of XRCC1-His, XRCC1-HisRK, His–XRCC1 161–406, His–APLF, His–APLFZFD and His–USP3 were added and incubated for 30 min on ice in Dilution buffer (20 mM Tris–HCl pH 7.5 and 130 mM NaCl). The plates were washed with PBST and incubated for 30 min with Dilution buffer containing mouse anti-polyhistidine antibody (Sigma) followed by three washes with PBS and subsequent incubation with horseradish peroxidase-conjugated secondary antibody (DAKO) for 30 min. After washing out the secondary antibody, 3,3′,5,5′-tetramethylbenzidine liquid substrate (slow kinetic form; Sigma) was added to the wells for 5 min. The reactions were stopped by the addition of 0.2 M HCl. Absorbance was read at 450 nm using a CLARIOstar microplate reader (BMG Labtech).

Statistics and reproducibility. No statistical method was used to predetermine the sample size, but we routinely employed at least three biological repeats for each experiment, in each case scoring as many technical replicates as possible (typically several hundred/thousand cells) using the applicable automated microscope software (scanR or Comet Assay IV). All replicates were successful and included in the data. No data were excluded from the analyses. The samples/ experiments were not randomized because all samples were specific genetic cell lines or samples derived from such and so randomization was not appropriate. The investigators were not blinded to allocation during the experiments and outcome assessment because all numerical data were software automated and independent of investigator subjectivity. All data presented in the manuscript are represented as the mean ± s.e.m., unless stated otherwise, and were analysed using GraphPad Prism (version 8/9). The statistical test employed for each dataset has been specified in the figure legends.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
Because of their large number, all microscope quantification datasets from this work, such as Excel and scanR/comet fluorescence quantification tables, are available from the corresponding author on reasonable request, citing the experiments of interest. Source data are provided with this paper.

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Author contributions
K.W.C. and M.A. conceived the study and designed the experiments. A.A.D. conducted the PARP1 biochemical experiments. E.K. established the primary neurons. J.B. generated the HPF1−/− gene-edited U2OS cell lines. S.E.W. performed the laser tracing experiments. A.G. generated the USP3−/− and XRCC1−/− USP3−/− RPE-1 cell lines. H.H. generated the XRCC1−/−, XRCC1−/− PARP1−/− and PARP1−/− RPE-1 cell lines published previously8,19. M.A. conducted all of the other experiments. K.W.C. and M.A. wrote the manuscript. K.W.C. coordinated the project. All authors analysed the data.

Competing interests
The authors declare no competing interests.

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Correspondence and requests for materials should be addressed to Keith W. Caldecott.
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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | XRCC1 promotes the recovery of transcription following oxidative damage. a, Immunoblot of RPA194 in WT and XRCC1−/− RPE-1 cells following mock treatment or 2 h after treatment with 250 μM H2O2 for 5 min. A representative image from one of three independent experiments is shown. b, Immunoblot of RNAPII hyperphosphorylation in WT and XRCC1−/− RPE-1 cells following mock treatment or 2 h after treatment with 250 μM H2O2 for 5 min. Where indicated, cells were pretreated with either 10 μM MG132 or 1 μM Bortezomib for 1 h before H2O2 treatment. A representative blot from one of three independent experiments is shown. c, RNAPI foci (RPA194) and levels of global transcription (eU immunofluorescence) in WT and XRCC1−/− RPE-1 cells pretreated for 1 h with 10 μM MG132 or 1 μM Bortezomib following mock treatment or at 2 h after treatment with 250 μM H2O2 for 5 min. d and e, Quantification of the levels of global transcription (EU immunofluorescence) and RNAPI foci (RPA194) shown in (c). Data are means (± s.e.m.) of three independent experiments.
Extended Data Fig. 2 | XRCC1 promotes the recovery of transcription following oxidative damage. a, Representative images of the RNAPI foci (RPA194) and levels of global transcription (EU immunofluorescence) in WT and XRCC1−/− U2OS cells stably transfected with either empty vector (EV) or expression construct encoding full length C-terminal histidine-tagged XRCC1 (XRCC1WT), following mock treatment or at the indicated times after treatment with 1 mM H2O2 for 20 min. Scale bars, 10 μm. b and c, Quantification of the RNAPI foci (RPA194) and levels of global transcription (EU immunofluorescence) shown in (a). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey’s multiple comparisons test (p values are indicated). d, Immunoblot of RNAPII hyperphosphorylation in WT U2OS cells, XRCC1−/− U2OS cells, and XRCC1−/− U2OS cells stably transfected with either empty vector (EV) or expression construct encoding full-length histidine-tagged XRCC1 (XRCC1WT), following mock treatment or 2 h after treatment with 1 mM H2O2 for 20 min. A representative blot from one of three independent experiments is shown.
Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | Reduced transcriptional recovery in XRCC1-deficient cells is a result of toxic PARP1 activity. 

**a,** Representative images of the RNAPI foci (RPA194 immunofluorescence) and levels of global transcription (EU immunofluorescence) in WT and XRCC1−/− RPE-1 cells pretreated with PARPi inhibitor (PARPi) following mock treatment or at the indicated times after treatment with 250 μM H2O2 for 5 min. Scale bars, 10 μm. 

b and c, Quantification of the RNAPI foci (RPA194) and levels of global transcription (EU immunofluorescence) shown in (a). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey’s multiple comparisons test (p values are indicated).

d, Representative images of the RNAPI foci (RPA194) and levels of global transcription (EU immunofluorescence) in WT and XRCC1−/− U2OS cells pretreated with PARPi inhibitor (PARPi) following mock treatment or at the indicated times after treatment with 1 mM H2O2 for 20 min. Scale bars, 10 μm.

e and f, Quantification of the RNAPI foci (RPA194) and levels of global transcription (EU immunofluorescence) shown in (d). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey’s multiple comparisons test (p values are indicated).
Extended Data Fig. 4 | Toxic PARP1 activity suppresses transcriptional recovery in XRCC1 patient fibroblasts and Xrcc1-deficient mouse cerebellar neurons. 

**a**, Quantification of the RNAPI foci (RPA194) shown in (Fig. 3a). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated).

**b**, Immunoblot of RNAPII hyperphosphorylation in normal fibroblasts (1BR) and XRCC1 patient fibroblasts (XD1) following mock treatment or 2 h after treatment with 60 μM H2O2 for 5 min. Cells were incubated with DMSO vehicle or with 10 μM PARP inhibitor (PARPi) for 1 h prior to, during, and following H2O2 treatment as indicated. A representative image from one of three independent experiments is shown.

**c**, XRCC1 and NeuN immunofluorescence in WT and Xrcc1nes-cre mouse cerebellar neurons. Scale bars, 10 μm.

**d**, Quantification of the levels of global transcription (EU immunofluorescence) shown in (Fig. 3c). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey’s multiple comparisons test.
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | The poly(ADP-ribose) binding activity of XRCC1 promotes transcriptional recovery following oxidative damage. a, Quantification of the levels of global transcription (EU immunofluorescence) shown in Fig. 4b. Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated). b, Representative images of the RNAPI (RPA194) and XRCC1 immunofluorescence in WT U2OS cells, XRCC1−/− U2OS cells, and XRCC1−/− U2OS cells transiently transfected with empty vector (EV) or expression construct encoding the indicated XRCC1 proteins, following mock-treatment or 2 h after treatment with 1 mM H2O2 for 20 min. Scale bars, 10 μm. c, Quantification of the RNAPI foci (RPA194) shown in (b). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated). d, Immunoblot of XRCC1 in XRCC1−/− U2OS cells stably expressing the indicated XRCC1 proteins. A representative blot from three independent experiments is shown. e, Immunoblot of RNAPII hyperphosphorylation in XRCC1−/− U2OS cells stably expressing the indicated XRCC1 proteins, following mock-treatment or 2 h after treatment with 1 mM H2O2 for 20 min. A representative image from one of three independent experiments is shown. f and g, Quantification of DNA strand breaks measured by alkaline comet assays in XRCC1−/− U2OS cells stably expressing the indicated XRCC1 proteins, following mock treatment (not treated; ‘NT’) or at the indicated times after treatment with either 50 μM H2O2 for 10 min on ice (conditions to measure SSB repair kinetic; panel f) or with 100 μM H2O2, for 20 min (refreshed every 10 min) in serum-free medium at 37 °C followed by recovery for 2 h (conditions, at which transcription recovery is perturbed in XRCC1−/− cells; panel g). Data show individual tail moments (an arbitrary measure of DNA breaks) from 50 cells per sample for three independent experiments. For each sample, the fifty tail moments are stacked vertically and the three experiments are positioned side-by-side (exp1, exp2, exp3) with the tick mark centred on exp2. Statistical significance was determined by one way ANOVA with Sidak’s two tailed Student’s t-test (p values are indicated). h, Quantification of the levels of global transcription (EU immunofluorescence) shown in Fig. 4d. Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey’s multiple comparisons test (p values are indicated). i, Immunoblot of APLF in the XRCC1−/− U2OS cells stably expressing the indicated APLF proteins. Single validation of SDS-PAGE mobility shift. j, Immunoblot of PARP1 auto-ADP-ribosylation in an in vitro ADP-ribosylation assay. Human recombinant PARP1 (100 nM) was incubated for 30 min in the presence of ssDNA (100 nM), NAD+ (2.5 μM) and with or without 1.5 μM of wild type or Zn-finger double mutant APLF (APLFWT, APLFZFD respectively). A representative blot from one of three independent experiments is shown. k, Binding of full length XRCC1 and APLF, as well as their PAR binding mutants denoted as XRCC1RK and APLFZFD respectively, to the indicated mock-ribosylated (-NAD+) or ribosylated (50 μM NAD+) histones. Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey’s multiple comparisons test (p values are indicated).
Extended Data Fig. 6 | Persistent PARP1 activation on the BER intermediates is a source of toxic PARP1 activity. a, Representative images of the RNAPI foci (RPA194) and levels of global transcription (EU Immunofluorescence) in WT and XRCC1−/− RPE-1 cells following mock treatment or at the indicated times after treatment with 250 μM H2O2 for 5 min. b, Quantification of the RNAPI foci (RPA194) shown in (a). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated). c and d, Immunoblot of PARP1 and/or ADP-ribose levels in total cell extract (panel c) and following ADP-ribose immunoprecipitation (panel d) prepared from WT and XRCC1−/− RPE-1 cells following mock treatment or at 2 h after treatment with 250 μM H2O2 for 5 min. Representative blots from one of three independent experiments are shown. e, Immunoblot of NTH1 in WT and XRCC1−/− RPE-1 cells treated for 72 h with 10 nM siRNA against NTH1. A representative blot from one of two independent experiments is shown.
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | Deletion of the histone ADP-ribosylation factor HPF1 does not rescue transcription recovery in XRCC1−/− RPE-1 cells. a, Model of the impact of HPF1 on PARP1-mediated histone ADP-ribosylation as well as PARP1 auto-ADP-ribosylation. b, Immunoblot of HPF1 and XRCC1 in WT, HPF1−/−, XRCC1−/−, and HPF1−/−/XRCC1−/− U2OS cells. A representative blot from one of three independent experiments is shown. c, Immunoblot of ADP-ribosylation upon H2O2 treatment (1 mM, 5 min) in WT, HPF1−/−, XRCC1−/−, and HPF1−/−/XRCC1−/− U2OS cells. A representative blot from one of three independent experiments is shown. d, Global transcription (EU immunofluorescence) in WT, HPF1−/−, XRCC1−/−, HPF1−/−/XRCC1−/− U2OS cells following mock treatment or at the indicated times after treatment with 1 mM H2O2 for 20 min. Scale bars, 10 μm. e, Quantification of the EU signal shown in (d). Data are means (±s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Tukey's multiple comparisons test (p values are indicated).
Extended Data Fig. 8 | Levels of histone acetylation, methylation, protein ubiquitylation in XRCC1+/− RPE-1 cells. 

**a,** Levels of H3K9Ac in RPE-1 cells, following treatment with 250 μM H₂O₂ for 5 min. Cells were incubated with 10 μM PARP inhibitor (PARPi) for 1 h prior to, during, and following H₂O₂ treatment. Quantification of H3K9Ac immunofluorescence shown on the right. Data are means (±s.e.m.) of four independent experiments. Scale bars, 10 μm.

**b,** Levels of H3K4me3 in RPE-1 cells, following treatment with 250 μM H₂O₂ for 5 min. Cells were incubated with 10 μM PARP inhibitor (PARPi) for 1 h prior to, during, and following H₂O₂ treatment. Quantification of H3K4me3 immunofluorescence shown on the right. Data are means (±s.e.m.) of three independent experiments. Scale bars, 10 μm.

**c,** Levels of chromatin mono/poly-ubiquitination (FK2) in RPE-1 cells, following treatment with 250 μM H₂O₂ for 5 min. Cells were incubated with 10 μM PARP inhibitor (PARPi) for 1 h prior to, during, and following H₂O₂ treatment. Quantification of chromatin mono/poly-ubiquitination immunofluorescence shown on the right. Data are means (±s.e.m.) of three independent experiments. Scale bars, 10 μm.
Extended Data Fig. 9 | Activated PARP1 recruits USP3 to induce histone deubiquitination and transcription inhibition. a, Global transcription (EU Immunofluorescence) in RPE-1 cells, following treatment with 250 μM H₂O₂ for 5 min. Representative images of cells 2 h after H₂O₂ are shown. Cells were pre-treated for 72 h with control or siRNA targeting USP11, USP22, USP36 and USP3. Quantifications are shown below images. Data are means (±SEM) of three or four independent experiments with statistical significance determined by Two-way ANOVA, Sidak’s multiple comparisons test (p values are indicated).

b, Levels of H2BK120mUb (H2BmUb; left) and H2AK119mUb (H2AmUb; right) in RPE-1 cells, following treatment with 250μM H₂O₂ for 5 min. Cells were pre-treated for 72 h with control or siRNA targeting USP3. Data are means (±SEM) of five independent experiments with statistical significance determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated).

c, Levels of H2BmUb (left) and H2AmUb (right) in RPE-1 cells, following treatment with 0.1 mg/ml MMS for 3 h. Cells were pre-treated for 72 h with control or siRNA targeting USP3. Data are means (±SEM) of four independent experiments with statistical significance determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated).
Extended Data Fig. 10 | USP3 does not interact directly with Poly-ADP-ribose chains. a, Cartoon of GFP-tagged USP3 constructs employed in this work. 

b, Representative ScanR images of chromatin retention of USP3 FL-GFP and USP3 ZnF-GFP in WT and XRCC1−/− U2OS cells, following mock-treatment or for 4 h with 0.1 mg/ml MMS in the presence/absence of 10 μM PARPi as indicated. Cells were transfected with H2B–mCherry and USP3–GFP constructs 24 h before experiment. Cells were fixed with PFA to show total protein levels or were pre-extracted with detergent prior to fixation to show chromatin-bound proteins. c, Quantification of experiments depicted in b. Data are means (± s.e.m.) of three independent experiments, and statistically significant differences were determined by two-way ANOVA with Sidak’s multiple comparisons test (p values are indicated). d, Immunoblot of PARP1 and USP3 ZnF-GFP levels in anti-ADP-ribose immunoprecipitates from WT and XRCC1−/− U2OS cell extracts following mock treatment or 1 h after treatment with 1 mM H2O2 for 20 min. A representative blot from one of two independent experiments is shown. e, Coomassie staining of purified recombinant human USP3 employed in the ADP-ribose binding assay shown in f. f, In vitro PAR binding assay. Data are means (± s.e.m.) of three independent experiments.
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Our web collection on statistics for biologists contains articles on many of the points above.

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Data analysis: scanR software, Comet Assay IV software, Graphpad Prism 9

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Data availability: Source data are provided with this study. Because of their large number, all raw microscope quantification data sets involved in this work such as Excel and scanR/comet fluorescence quantification tables (which generated the provided source data) are available from the corresponding author on reasonable request, citing the experiments of interest.
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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size          | No statistical method was used to predetermine sample size, but we routinely employed at least three biological repeats for each experiment, in each case scoring as many technical replicates as possible (typically several hundred/thousand cells) using the applicable automated microscope software (scanR/comet assay IV). |
|----------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Data exclusions      | no data was excluded                                                                                                                                                                               |
| Replication          | all findings were independently replicated at least 3 times. All replicates were successful and included in the data                                                                                     |
| Randomization        | this is not appropriate because all of our samples/experiments were specific genetically defined cell lines                                                                                           |
| Blinding             | Samples/investigators were not blinded during experiments and outcome assessment because all numerical data was software automated and independent of investigator subjectivity.                             |

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We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | n/a                   |
| ☑   | Antibodies            |
| ☑   | Eukaryotic cell lines |
| ☑   | Palaeontology and archaeology |
| ☑   | Animals and other organisms |
| ☑   | Human research participants |
| ☑   | Clinical data         |
| ☑   | Dual use research of concern |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | n/a                   |
| ☑   | ChIP-seq              |
| ☑   | Flow cytometry        |
| ☑   | MRI-based neuroimaging |

Antibodies

Antibodies used

- see below
- Primary abs:
  - Antibody: Provider: Use: Cat. No.: Species:
  - XRCC1 Novus IF 1:300 WB 1:8000 NBP1-87154 Rabbit
  - TUBULIN Sigma WB 1:20000 T6074 Mouse
  - PARP1 BioRad WB 1:1000 MCA1522G Mouse
  - RNAPII Santa Cruz WB 1:100 sc-9001 Rabbit
  - RNAPII Santa Cruz WB 1:100 sc-55492 Mouse
  - NeuN Millipore IF 1:500 MAB377 Mouse
  - RNAPII pS2 Abcam WB 1:30000 ab5095 Rabbit
  - RNAPII pS5 Abcam WB 1:30000 ab5131 Rabbit
  - ADP-ribose binding reagent Millipore IF 1:1000 WB 1:10000 MABE1016 Rabbit
  - RPA194 Santa Cruz IF 1:200 WB 1:100 sc-48385 Mouse
  - NTH1 Santa Cruz WB 1:200 sc-130644 Mouse
  - HPF1 Novus WB 1:1000 NBP1-93973 Rabbit
  - H2BK120mUb Cell Signaling IF 1:800 5546T Rabbit
  - H2AK119mUb Cell Signaling IF 1:1000 8240T Rabbit
  - H3K9Ac Cell Signaling IF 1:1000 9649T Rabbit
  - H3K4me3 Abcam IF 1:400 Ab8580 Rabbit
  - USP11 Proteintech WB 1:1000 10244-1-AP Rabbit
  - FK2 BioRad IF 1:300 MCA6035 Mouse
KU80 Invitogene WB 1:10000 MAS-12933 Mouse

Secondary abs:
Alexa Fluor 488 Invitogene IF 1:10000 A21206 Donkey, (anti-Rabbit)
Alexa Fluor 488 Invitogene IF 1:1000 A11001 Goat, (anti-Mouse)
Alexa Fluor 647 Invitogene IF 1:1000 A21245 Goat, (anti-Rabbit)
Alexa Fluor 647 Invitogene IF 1:1000 A21235 Goat, (anti-Mouse)
APLF Invitogene WB 1:1000 PAS39226 Rabbit
USP3 Abcam WB 1:500 Ab229348

Validation
antibodies were chosen based on published literature/recommendation and were either not further validated or were internally validated by the use of cell lines lacking the target antigen

Eukaryotic cell lines
Policy information about cell lines
Cell line source(s) RPE1 cells and U2OS cells were both from ATCC. Mouse cerebellar neurons were generated from mice in-house and human fibroblasts from our previous work (Hoch et al Nature 2017)
Authentication these are all authenticated by fingerprinting in our cell culture facility
Mycoplasma contamination all cells were confirmed mycoplasma negative.
Commonly misidentified lines (See ICLAC register) no commonly misidentified cell lines were employed in this study

Animals and other organisms
Policy information about studies involving animals, ARRIVE guidelines recommended for reporting animal research
Laboratory animals no live laboratory animals were used in this study. We used only isolated neurones.
Wild animals no wild animals were employed
Field-collected samples not applicable
Ethics oversight we are governed by the UK Home Office and by the University Ethics committee. Our animal and research licenses are fully approved by such.

Note that full information on the approval of the study protocol must also be provided in the manuscript.