ROS-induced R loops trigger a transcription-coupled but BRCA1/2-independent homologous recombination pathway through CSB

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Actively transcribed regions of the genome are protected by transcription-coupled DNA repair mechanisms, including transcription-coupled homologous recombination (TC-HR). Here we used reactive oxygen species (ROS) to induce and characterize TC-HR at a transcribed locus in human cells. As canonical HR, TC-HR requires RAD51. However, the localization of RAD51 to damage sites during TC-HR does not require BRCA1 and BRCA2, but relies on RAD52 and Cockayne Syndrome Protein B (CSB). During TC-HR, RAD52 is recruited by CSB through an acidic domain. CSB in turn is recruited by R loops, which are strongly induced by ROS in transcribed regions. Notably, CSB displays a strong affinity for DNA:RNA hybrids in vitro, suggesting that it is a sensor of ROS-induced R loops. Thus, TC-HR is triggered by R loops, initiated by CSB, and carried out by the CSB-RAD52-RAD51 axis, establishing a BRCA1/2-independent alternative HR pathway protecting the transcribed genome.
Recent advances in our understanding of DNA repair mechanisms have revealed the complexity of how cells respond to DNA damage, particularly in the context of transcription. In transcribed regions, DNA damage can lead to transcriptional stalling and potential repair at non-homologous end-joining (NHEJ), homologous recombination (HR), or a combination of both pathways. The precise choice of repair pathway can depend on various factors, including the type and location of damage, as well as the state of transcription.

The repair of DNA lesions in transcribed regions poses a significant challenge, as transcriptional states and DNA repair mechanisms are tightly integrated. In this context, the role of transcription-coupled repair (TC-R) and transcription-coupled homologous recombination (TC-HR) is crucial. TC-R is initiated by transcription-coupled nucleases, which recognize transcriptional stress and recruit repair factors, while TC-HR is activated by the recruitment of RAD51 to DSBs in BRCA1/2-depleted cells. The presence or absence of APH (Supplementary Fig. 1e, lower panels) indicates that repair had occurred through one or more APH-refractory polymerases. Notably, RAD51 was required for the induction of DSBs by ROS and the activation of TC-HR. To test whether DNA replication affects the function of TC-HR, we followed DNA repair at the TA-KR site in the absence or presence of APH. Although APH increased yH2AX throughout the genome (Supplementary Fig. 1e, upper panel), it did not affect the cleavage of yH2AX at the TA-KR site 24–36 h after damage induction (Supplementary Fig. 1e, lower panels), indicating that repair had occurred through one or more APH-refractory polymerases. Notably, RAD51 was required for the cleavage of yH2AX at the TA-KR locus regardless of the presence or absence of APH (Supplementary Fig. 1e, lower panels). Together, these results suggest that the function of RAD51 in TC-HR is independent of DNA replication.

In the canonical HR pathway, BRCA1 and BRCA2 promote the localization of RAD51 to DSBs. In a previous study, we showed that BRCA1 and BRCA2 were also recruited to sites of ROS-induced DNA damage. Surprisingly, knockdown of BRCA1 and BRCA2 did not affect the recruitment of RAD51 to the locus marked by TA-KR (Fig. 1c and Supplementary Fig. 2a). We then turned our attention to RAD52, which promotes the localization of RAD51 to DSBs in BRCA1/2-deficient cells. We created RAD52 knockout (KO) U2OS TRE cells (Supplementary Fig. 2b). In RAD52 KO and Knockdown cells, RAD51 foci were diminished at the TA-KR marked locus (Fig. 1d and Supplementary Fig. 2c). The defect of RAD51 localization in RAD52 KO cells was suppressed by exogenous RAD52 (Fig. 1e).

Results

RAD52 but not BRCA1/BRCA2 recruits RAD51 in TC-HR. To understand how cells protect the actively transcribed genome against ROS-induced DNA damage, we used KillerRed (KR), a light-excitable and ROS-releasing chromophore, to conditionally activate the transcription activator VP16 and KR (tetR-KR) binds to the TRE array but does not activate transcription. Upon light activation, both TA-KR and tetR-KR release ROS locally, inducing equivalent amounts of DNA damage marked by yH2AX at the locus in the presence and absence of transcription, respectively (Supplementary Fig. 1a). Following damage induction, Ku70 and Ku80 are immediately recruited to KR sites, showing the efficient induction of DSBs by ROS (Supplementary Fig. 1b).

In TA-KR-expressing cells, the key HR protein RAD51 was readily detected at the locus marked by TA-KR upon light activation (Fig. 1a). In contrast, RAD51 was not efficiently detected at the tetR-KR locus, suggesting that RAD51 is preferentially recruited to the damage site in a transcription-dependent manner. To test whether RAD51 is functionally important for repairing ROS-induced DNA damage, we monitored the clearance of yH2AX over time (Fig. 1b and Supplementary Fig. 1c). At 1 h after light activation, TA-KR induced similar levels of yH2AX in RAD51 knockout cells and control cells. However, yH2AX levels were significantly higher in RAD51 knockout cells and control cells. Furthermore, yH2AX levels were significantly higher in RAD51 knockout cells than in control cells after 36 h, suggesting that DSBs were not efficiently repaired in the absence of RAD51. These results demonstrate that ROS-induced DNA damage at a transcribed locus triggers RAD51-dependent TC-HR.

Rad51 not only functions in HR but also protects stalled replication forks. Blocking DNA replication by Aphidicolin (APH) before damage induction did not affect the formation of yH2AX and RAD51 foci at the TA-KR locus (Supplementary Fig. 1d), showing that DNA replication is not required for the induction of DSBs by ROS and the activation of TC-HR. To test whether DNA replication affects the function of TC-HR, we followed DNA repair at the TA-KR site in the absence or presence of APH. Although APH increased yH2AX throughout the genome (Supplementary Fig. 1e, upper panel), it did not affect the cleavage of yH2AX at the TA-KR site 24–36 h after damage induction (Supplementary Fig. 1e, lower panels), indicating that repair had occurred through one or more APH-refractory polymerases. Notably, RAD51 was required for the cleavage of yH2AX at the TA-KR locus regardless of the presence or absence of APH (Supplementary Fig. 1e, lower panels). Together, these results suggest that the function of RAD51 in TC-HR is independent of DNA replication.

The CSB-RAD52-RAD51 axis promotes TC-HR. To understand how ROS affect DSB repair in transcribed regions.
ROS-induced TC-HR is regulated. CSB is an important regulator of transcription-coupled nucleotide excision repair (TC-NER). We recently showed that CSB was recruited by ROS-induced DNA damage at a transcribed locus. To understand the function of CSB in TC-HR, we created CSB KO U2OS TRE cells. Similarly, in RAD52-depleted cells, the localization of RAD51 to the TA-KR-marked locus was diminished in CSB KO and knockdown cells.

Fig. 1 ROS trigger BRCA2/1-independent, RAD52-, and RAD51-dependent TC-HR. a Schematic diagram of the RAD51 damage response to KillerRed (KR)-mediated ROS-induced damage at transcription on (TA-KR) or off (tetR-KR) genomic loci in U2OS TRE cells (scale bar: 2 μm). b γH2AX foci frequency at TA-KR at early (1 h) or late (36 h) time points after light-induced KillerRed activation in siCtrl and siRAD51-treated cells (scale bar: 2 μm). c RAD51 foci frequency at TA-KR in cells treated with control, BRCA1, or BRCA2 siRNAs (scale bar: 2 μm). d RAD51 foci frequency at TA-KR in WT and RAD52 KO cells (scale bar: 2 μm). e RAD51 foci frequency at TA-KR in RAD52 KO cells and RAD52 overexpressed KO cells. f γH2AX foci frequency at TA-KR at early (1 h) and late (36 h) time points after damage in RAD52 KO cells treated with or without siRAD51. g Model of RAD51 regulation under ROS or nuclease-induced damage at transcribed regions. For a–f, n = 3, 50 cells per replicate. Unpaired t-test, error bars represent SEM. *P < 0.05, **P < 0.01, ***P < 0.001.
Supplementary Fig. 3a-c), suggesting that CSB functions upstream of RAD51. As CSB is also required for the localization of RAD51 to the TA-KR-marked locus⁹, CSB likely promotes RAD51 recruitment through RAD52. The defects of RAD52 and RAD51 localization in CSB KO cells were suppressed by exogenous full-length CSB (CSB-FL) (Fig. 2b-c). Notably, CSB-FL was unable to rescue RAD51 localization in the CSB KO cells treated with RAD52 small interfering RNA (siRNA) (Fig. 2c), confirming that CSB promotes RAD51 localization through RAD52. Knockdown of RAD52, RAD51, and CSB either individually or in combinations resulted in similar defects in knockdown of CSB (Fig. 2d and Supplementary Fig. 3d), suggesting that these proteins function in the same pathway. Knockdown of BRCA1/2 did not affect the localization of CSB and RAD52 to the TA-KR-marked locus (Supplementary Fig. 3e), supporting the notion that the CSB-RAD52-RAD51 axis functions independently of BRCA1/2.

Ionizing radiation (IR) induces ROS²² and generates DSBs and other DNA lesions in both transcribed and untranscribed regions. We observed a mild defect in IR-induced RAD51 focus formation in CSB KO and RAD52 KO cells (Fig. 2e). Furthermore, CSB KO and RAD52 KO cells displayed...
modestly increased IR sensitivity in the clonogenic assay (Fig. 2f), which was also observed in previous studies.\textsuperscript{14,15,23,24} The RNAPII inhibitor 5,6-Dichlorobenzimidazole 1-β-D-ribofuranoside (DRB) sensitized BRCA2 knockdown cells to IR, but did not increase the IR sensitivity of CSB KO and RAD52 KO cells (Fig. 2g and Supplementary Fig. 3f). Therefore, transcription contributes to the repair of IR-induced DNA damage independently of BRCA2 but in the same pathway as CSB/RAD52, which is consistent with the TC-HR pathway (Fig. 2h).

CSB regulates RAD52 through an AD. To understand how CSB regulates RAD52 in TC-HR, we stably expressed a panel
of CSB fragments in CSB KO cells (Supplementary Fig. 4a)\(^25\). Among the CSB fragments, only the fragment containing the CSB-AD (337–509 a.a.) partially rescued the formation of RAD52 foci at the TA-KR-marked locus in CSB KO cells (Fig. 3a). Furthermore, CSB-AD also partially rescued the formation of RAD51 foci in CSB KO cells (Supplementary Fig. 4b). Similar to CSB-FL, CSB-AD rescued RAD51 foci in a RAD52-dependent manner (Fig. 3b). Consistent with the rescue of RAD52 and RAD51 foci, CSB-AD partially suppressed the IR sensitivity of CSB KO cells (Fig. 3c). CSB-AD was able to localize to the TA-KR-marked locus, although less efficiently than CSB-FL (Fig. 3d). Another C terminus fragment of CSB also localized to the TA-KR-marked locus, suggesting that CSB localization is regulated by a bipartite mechanism. Collectively, these results suggest that CSB may function in TC-HR through its AD. As CSB-AD is dispensable for TC-NER upon UV damage\(^28\), the functions of CSB in TC-HR and TC-NER are likely distinct.

An interaction between CSB-AD and RAD52 was observed in cell extracts and in binding assays using purified proteins (Fig. 3e and Supplementary Fig. 4c–d). To understand how CSB-AD interacts with RAD52, we treated purified CSB-AD and RAD52 proteins with disuccinimidyl carbonate (DSS) to crosslink lysine residues in close proximity (Supplementary Fig. 4e). Mass spectrometry (MS) analysis detected multiple crosslinks between RAD52 and the N terminal region of CSB-AD (Fig. 3f and Supplementary Fig. 4f). The N terminal region of CSB-AD is highly enriched for acidic amino acids. To test whether these acidic amino acids are important for RAD52 binding, we mutated 10 or 22 of them into alanines in CSB-AD, resulting in CSB-AD10A and CSB-AD22A mutants (Fig. 3g and Supplementary Fig. 5a). Although CSB-AD22A was able to localize to the TA-KR-marked locus, its binding to RAD52 was significantly reduced compared with CSB-AD and CSB-AD10A (Fig. 3h). Another CSB-AD mutant harboring 12 acidic amino acid mutations outside of the 10A region (CSB-AD12A) still interacted with RAD52 (Supplementary Fig. 5a–b), showing that the 22 acidic amino acids may be redundant for RAD52 interaction. Indeed, CSB-AD, CSB-AD10A, and CSB-AD12A could efficiently rescue RAD52 foci in CSB KO cells, whereas CSB-AD22A could not (Fig. 3i and Supplementary Fig. 5c). Furthermore, CSB-AD and CSB-AD10A, but not CSB-AD22A, rescued RAD51 foci and suppressed the defect of γH2AX clearance in CSB KO cells (Fig. 3j–k). Importantly, the CSB-FL mutant containing the 22A mutations (CSB-22A) also failed to rescue RAD52 foci and γH2AX resolution in CSB KO cells (Supplementary Fig. 5d–e). These results collectively suggest that the interaction of CSB with RAD52 through the acidic amino acids in CSB-AD is critical for TC-HR. Interestingly, CSB homologs in mice and Zebrafish, but not yeast, display an enrichment of acidic amino acids (Supplementary Fig. 5f), suggesting that the regulation of RAD52 has evolved in vertebrates.

ROS induce R loops at a transcribed locus. The transcription dependency of TC-HR prompted us to investigate whether DNA is involved in activating the CSB-RAD52-RAD51 axis. We previously showed that expression of RNase H1, which cleaves the RNA in DNA:RNA hybrids, inhibited TC-HR\(^9,27\). This result raised the possibility that R loops, which contain DNA:RNA hybrids and displaced single-stranded DNA, are induced by ROS\(^28\). Indeed, using a monoclonal antibody (S9.6) that specifically recognizes DNA:RNA hybrids, we detected R loops at the locus marked by TA-KR (Fig. 4a). The R loop signals at the TA-KR-marked locus were reduced by expression of RNase H1, but not the catalytically inactive mutant D210N (Fig. 4b)\(^29\). In contrast to TA-KR, tetR-KR did not induce R loops (Fig. 4a), suggesting that R loop formation is transcription-dependent. A fusion protein that activates transcription but does not induce ROS (TA-mCherry) and another fusion protein that lacks both transcription activation and ROS-releasing activities (tetR-mCherry) did not generate R loops (Fig. 4a), suggesting that both transcription and DNA damage are needed for R loop formation. In addition, inhibition of RNAPII reduced the R loops at the TA-KR-marked locus (Fig. 4c). DNA–RNA immunoprecipitation (DRIP) confirmed that R loops were induced at the TA-KR locus in a damage- and transcription-dependent manner (Fig. 4d and Supplementary Fig. 6a). In addition, super-resolution imaging of the TA-KR-marked locus with stochastic optical reconstruction microscopy (STORM) confirmed that R loops were indeed in close proximity to TA-KR (Fig. 4e). These results demonstrate that R loops are robustly induced by ROS when TC-HR is activated. ROS induce both SSBS and DSBS. In vitro, SSBS in transcribed DNA are a potent inducer of R loops\(^30\). The DSBS generated by an endonuclease only modestly increased DNA:RNA hybrids in transcribed regions\(^31,32\). Together, these findings suggest that ROS may induce R loops by generating SSBS and DSBS in transcribed regions, providing a possible explanation for the unique ability of ROS to induce TC-HR efficiently.

CSB binds R loops at damage sites through its C terminus. We next asked how R loops contribute to TC-HR activation. As expected, both CSB and RAD52 colocalized with R loops at the TA-KR-marked locus (Fig. 5a). Importantly, the localization of CSB and RAD52 at the TA-KR-marked locus was reduced by RNase H1, but not the D210N mutant (Fig. 5b and Supplementary Fig. 6b), showing that CSB and RAD52 are recruited to the damage site in an R loop-dependent manner. The role of R loops in recruiting CSB and RAD52 suggests that R loops may be functionally important for TC-HR. Indeed, removal of R loops by RNase H1 expression increased the γH2AX signal at the locus marked by TA-KR (Fig. 5c), suggesting that TC-HR was compromised. These results suggest that R loops are required for activating the CSB-RAD52-RAD51 axis in TC-HR. Interestingly, a recent paper suggested that the HR function of BRCA1 is impaired by its interaction with the transcription elongation...
machinery in cells with high levels of R loops, raising the possibility that ROS-induced R loops may promote a switch from canonical HR to TC-HR in transcribed regions.

To address how R loops promote the recruitment of CSB and RAD52, we performed electrophoresis mobility shift assays using synthetic DNA:RNA hybrids and purified CSB-FL and RAD52 proteins (Supplementary Fig. 4c, 6c). Consistent with previous studies, RAD52 has an affinity for DNA:RNA hybrids (Fig. 5d). Strikingly, CSB-FL displayed a much higher affinity for DNA:RNA hybrids than RAD52 (Fig. 5d). These results suggest that both CSB and RAD52 may directly recognize the R loops at sites of DNA damage, and that CSB may enhance the recruitment of RAD52 to R loops by binding to both R loops and RAD52. The strong affinity of CSB to DNA:RNA hybrids in vitro, as well as its dependency on DNA:RNA hybrids to localize to sites of DNA damage in cells, suggest that CSB is likely a sensor of ROS-induced R loops that initiates the CSB-RAD52-RAD51 functional cascade.

To further understand how CSB senses R loops, we tested the binding of CSB fragments to DNA:RNA hybrids in cell extracts. Consistent with the bipartite mechanism of CSB localization to the TA-KR site, CSB-AD and the CSB C-terminal fragments 961–1493 and 1200–1493 were captured by DNA:RNA hybrids (Supplementary Fig. 6d). Similar to CSB-AD, CSB 961–1493 and 1200–1493 were able to localize to the TA-KR site (Supplementary Fig. 6e). The CSB C-terminal fragments bound to DNA:RNA more efficiently than CSB-AD, suggesting that they contain the major DNA:RNA-binding domain of CSB. Importantly, using purified protein, we found that CSB 1200–1493 bound to DNA:RNA hybrids directly (Fig. 5e and Supplementary Fig. 6f), thereby revealing the DNA:RNA-binding domain of CSB. Unlike CSB 1200–1493, purified CSB-AD did not bind to DNA:RNA hybrids directly in vitro (Supplementary Fig. 6g), indicating that its localization to R loops might be mediated by other R loop-binding proteins.

**Discussion**

Although the role of transcription in TC-NER has been long appreciated, its role in TC-HR has just begun to unfold. In this study, we used a unique experimental system to characterize ROS-induced TC-HR at a transcribed locus. We discovered that ROS induce R loops robustly in transcribed regions, recruiting CSB to sites of DNA damage through its C-terminal domain (CTD). Once recruited by R loops, CSB uses its AD to interact with RAD52 and promote its localization to DNA damage sites.
The function of CSB in TC-HR is distinct from its function in TC-NER, revealing a new functional mode of CSB in transcription-coupled DNA repair. Why do ROS preferentially activate TC-HR? Although high levels of R loops promote the recruitment of CSB and RAD52, they may inhibit the functions of certain canonical HR proteins. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair. For example, in Ewing sarcoma, EWS-FLI1 induces R loops, which increases the interaction between RNAPII and BRCA1, and decreases the interaction between RNAPII and BRCA2. Together, a functional cascade of CSB, RAD52, and RAD51 is triggered by ROS-induced R loops, protecting the transcribed genome from transcription-coupled DNA repair.
ROS and nuclease damage induction. U2OS TRE cells were cultured in 35 mm glass-bottom dishes (MatTek, P35GC-1.5-14-C) at 60% confluence 24–36 h before the transfection. For ROS-induced damage, cells were transfected with plasmids containing KR (TA-KR/tetR-KR). Light-induced KR activation was done by exposing cells to a 15W Sylvania cool white fluorescent bulb for 25 min. Cells were recovered at around 1 h before live-cell observation or fixation. For γH2AX staining, cells were recovered for 36 h before fixation. For nuclease-induced damage, cells were co-transfected with pCMV-I-SceI plasmid and tetR-Cherry, and incubated for 36 h before harvest.

Immunofluorescence staining and microscopy. Cells in a 35 mm dish were rinsed with phosphate-buffered saline (PBS) and fixed in 4% paraformaldehyde (PFA; Affymetrix, 19943-1 LT) for 15 min at room temperature. They were washed three times by PBS, permeabilized by 0.2% Triton X-100 in PBS for 8 min, then washed three times by PBS. Then they were blocked by 5% bovine serum albumin (BSA) (SIGMA, A-7030) in 0.1% PBS-Tween (PBST) for 30 min at room temperature. Primary antibodies were diluted in blocking buffer and incubated with cells overnight at 4 °C. Then the cells were washed three times with 0.05% PBST and incubated with secondary antibodies for 1 h at room temperature, including Alexa Fluor 405/488/594 goat anti-mouse/rabbit IgG conjugate (1:10,000). Finally, they were washed three times by 0.05% PBST and optionally stained with DAPI (4′,6-diamidino-2-phenylindole; 1:1000 in PBS) for 5 min at room temperature. The primary antibodies for immunostaining are RAD51 (ab63801, Abcam, 1:100), γH2AX ser139 (JBW301, 05-636, EMD Millipore, 1:400), Myc-tag (ab9106 and ab32, Abcam, 1:200), HA (12CA5, 11666606001, Roche, 1:300), and S9.6 (ENH001, Kerafast, 1:200). APH was purchased from Abcam (ab142400).

For S9.6 staining, the cells were fixed and permeabilized in a 35 mm glass-bottom dish using a standard protocol, then incubated in buffer (10 mM Tris-HCl, 2 mM EDTA, pH = 9) and steamed on a 95 °C heating block for 20 min to expose the antigen. Then the dish was cooled, washed three times with PBS, and treated with RNase A (100 μg/mL) in buffer (5 mM EDTA, 300 mM NaCl, 10 mM Tris-HCl, pH 7.5) for 15 min at room temperature. Then the cells were washed and blocked using 5% BSA in 0.1% PBST for 0.5 h at room temperature. The first and secondary antibodies were diluted in the same buffer (5% BSA in 0.1% PBST) and the standard immuno-fluorescence protocol was followed. This protocol was

![Fig. 6 Model of CSB-, RAD52-, and RAD51-mediated TC-HR at ROS-induced R loops in transcribed regions. The current model is composed of the following sequential steps: ROS at transcribed regions trigger R loop formation; CSB senses R loops via its CTD; AD of CSB binds RAD52 and facilitates RAD52 recruitment at R loops; CSB-RAD52 promotes RAD51 loading at R loops; repair completion and transcription restart.](image-url)
modified from the classical heat-induced antigen retrieval method for PFA-fixed tissues using Tris-EDTA buffer. For the RNAPII inhibitor, DRB (D1916; Sigma) or α-amanitin (A2263; Sigma) was added with a final concentration of 20 μM or 100 μg/mL for 24 h or 2 h before light irradiation, respectively.

The images were acquired using the Olympus FX100 confocal microscopy system (Cat. F10PROIR-V1-R, Olympus) and Fiji (an open source software). For quantification of the percentage of foci positive cells at sites of KR, 50 cells were counted for each sample experiment and representative data are shown. For quantification of relative foci intensity, the intensity of foci and background was acquired by ImageJ 1.50i.*

Co-immunoprecipitation and western blots. FIp in 293 cells were co-transfected with expression vectors 36–48 h after transfection; cell lysates were collected in 1 mL of lysis buffer (10 mM Hepes, pH 7.6; 50 mM Tris-HCl, pH 7.4; 150 mM NaCl; 1% NP-40, 3 mM EDTA; protease inhibitor from Roche). For anti-fluorescent protein (GFP) immunoprecipitation, 2 μg anti-GFP monoclonal antibody (1181446001, Roche), and 25 μL of G-Sepharose protein beads (GE Healthcare Bio-Sciences) were added to each lysate. Mixtures were incubated at 4 °C overnight with rotation; the supernatant was removed and protein beads were washed four times with 4 mL of lysis buffer. For western blotting analysis, samples were boiled at 95 °C for 5–8 min in SDS loading buffer. Then they were subjected to electrophoresis in 10–12% SDS-polyacrylamide gels and transferred to the polyvinylidene difluoride membrane. The membranes were blocked with 5% non-fat milk in PBS for 1 h before being incubated with primary antibody at 4 °C overnight. The primary antibodies for western blotting used in this study are GFP (1181446001, Roche, 1:2000), β-Actin (60008-1-Ig, Cell Signaling Technology, 1:20,000), and β-Gal (512124, Promega, 1:10,000). The secondary antibody was donkey anti-rabbit IgG-HRP (A10354, Thermo Fisher, 1:10,000) for 1 h at room temperature. The membranes were washed in 0.1% PBST for four times before exposure. Chemiluminescent HRP substrate was purchased from Millipore (Catalog#: WBKLS0500). Images were acquired in a BIO-RAD Universal Hood II machine with ImageLab software. The uncropped scans of the important blots are displayed in Supplementary Fig. 7.

DNA-RNA immunoprecipitation. The DRIP assay was performed according to the literature with minor modifications.55. U2OS TRE cells in a 10 cm dish were transfected with TA-KR or TA-Cherry, exposed to light for 20 min in PBS, and recovered for 1 h. The cells were washed and pelleted, resuspended in 1.6 mL of TE, and 41.5 μL of 20% SDS and 0.5 μL of Proteinase K (Roche Life Sciences) were added, and then the samples were incubated overnight at 55 °C. After 24 h of incubation, 800 μL of insect cells were lysed in Buffer A (25 mM Tris-Cl pH 7.5, 10% Glycerol, 0.5 mM EDTA, 300 mM KCl, 0.1% IGEPAL, 1 mM dithiothreitol (DTT), 1 mM phenyl-methylsulfonyl fluoride (PMSF), and protease inhibitors (5 μg/mL each of leupeptin, chymotrypsinogen, aprotinin, and peptatin)) and sonicated for 30 s on a sonicator. The lysate was subjected to ultracentrifugation for 60 min at 100,000 × g. The supernatant was added to 3 mL of anti-flag M2 Affinity Agarose Gel (Sigma) pre-washed in Buffer B and rotated for 2 h. Beads were then washed four times, each with with 20 mL Buffer B (25 mM Tris-Cl pH 7.5, 10% Glycerol, 0.5 mM EDTA, 300 mM KCl, 0.1% IGEPAL, 1 mM DTT), followed by elution in 15 mL Buffer B with 200 μg/mL Flag peptide. The eluate was added to 3 mL of Nickel-NTA beads (Qiagen), pre-washed in Buffer B with 20 mM imidazole, and rotated for 2 h. Nickel-NTA beads were washed four times each with 10 mL Buffer B and 20 mM imidazole, followed by elution in 10 mL Buffer B with 200 mM imidazole. The eluate was subjected to size exclusion chromatography on a Superdex column in Buffer B (20 mM Tris-Cl pH 7.5, 10% Glycerol, 0.5 mM EDTA, 300 mM KCl, 0.1% IGEPAL, 1 mM DTT), followed by elution in 15 mL Buffer B with 200 μg/mL Flag peptide. The eluate was pooled, concentrated using 100 kDa cutoff concentrator, and stored in small aliquots at –80 °C.

Rosetta cells harboring pET28b-RAD52 that encode hRAD52 with a C-terminal 6–4× histag were grown to an optical density of 0.6 and induced by adding 0.3 mM IPTG for 4 h at 37 °C. Ten grams of overexpressed cell mass were lysed in buffer ly$15$ (25 mM Tris-HCl (pH 7.5), 500 mM KCl, 1 mM EDTA, 10% glycerol, 1 mM DTT, 0.01% Igepal, 1 mM PMSF, and a mixture of protease inhibitors) and sonicated. The lysed sample was centrifuged at 1 h at 16,000 g (11,000 g, 10 min). Cleared supernatant was diluted five fold and applied on Affihive Nickel-NTA agarose beads. The beads were washed three times with Buffer B containing 100 mM KCl, 1% CHAPS (11814460001, Roche, 1:2000), CSB (H-300, sc-25370, Santa Cruz Biotechnology, 1:1000), β-Actin (60008-1-Ig, Cell Signaling Technology, 1:20,000), and β-Gal (512124, Promega, 1:10,000). The eluate was concentrated and stored in small aliquots at –80 °C.

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Hybrid substrate preparation. The 5′-end of the oligo 1 (Supplementary Table 3) was labeled using a 5′-oligonucleotide end labeling kit (Vectorlabs) and a maleimide-IR800 probe (LI-COR Biosciences). RNA–DNA substrate was prepared by annealing oligo 1 and 2 (Supplementary Table 3) and confirmed. 22 Briefly, 5′-end labeled oligo 1 was mixed with oligo 2 in buffer H [90 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, and 50 mM NaCl], heat denatured, and annealed by slow cooling. Annealed substrates were separated by 10% native polyacrylamide gel electrophoresis (PAGE)-Tris-accetate-EDTA. The corresponding gel bands were excised and eluted. RNA–DNA hybrid substrate was confirmed by mobility in native PAGE, heat denaturation, and RNAseH treatment. 22

Electrophoretic mobility shift assay. 5′-End maleimide-IR800-labeled hybrid substrate was incubated with RAD52 or CSB in Buffer B [25 mM Tris-HCl (pH 7.5), 1 mM MgCl₂, 1 mM DTT, 50 µg/mL BSA] with 50 mM NaCl for 15 min at 37°C. Reaction mixtures were loaded on 6% PAGE-TBE gel and resolved at 4°C. Gels were imaged using an Odyssey scanner (LI-COR Biosciences) and quantitated.

Biotin-labeled hybrid pulldown assay. 3′-Biotin-labeled single ssRNA (oligo 3) and complementary ssDNA (oligo 4) were synthesized from IDT and annealed (Supplementary Table 3). The 293 Flp-in cells overexpressed with GFP-CSB 337–509, 510–960, 961–1493, 1206–1493, and 1400–1493 were lysed in Pierce IP lysis buffer (25 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA; 1% NP-40; 500 µM NaCl; 1% Tween-20) and eluted using SDS loading buffer. The binding supernatants and elution fractions were analyzed by western blotting.

Chemical crosslink and MS analysis. To crosslink the protein complex, the recombinant His-RAD52 and GST-CSB-AD (residues 337–509) were incubated with 1 mM DMS in an amine-free buffer (100 mM Hepes, pH 7.5, 50 mM KCl). The cross-linking reactions were performed at 25°C for 1 h with constant agitation (1300 r.p.m.) before subsequent quenching with 50 mM ammonium bicarbonate (final concentration) for 10–15 min. The cross-linked samples were then briefly centrifuged to remove any potential insoluble products and the supernatants were collected for SDS-PAGE analysis. After protein reduction and alkylation, the cross-linked samples were separated by a 4–12% SDS-PAGE gel (NuPAGE, Thermo Fisher). The regions (> 250 kDa) corresponding to the cross-linked species were cut and digested with trypsin. 36–38 After proteolysis, the peptide mixtures were desalted and analyzed with a nano-LC 1200 system that is coupled online with a QExactive basic mass spectrometer (Thermo Fisher). The detailed peptides were loaded onto a picochip column (C18, 3 µm particle size, 300 Å pore size, 50 µm × 10.5 cm; New Objective) and eluted using an 80 min LC gradient. The QE instrument was operated in the data-dependent mode. The target resolution was 75,000 for MS and 15,000 for tandem MS (MS/MS) analyses. The quadrupole isolation window was 1.8 Th and the maximum injection time for MS/MS was set at 800 ms. The data were searched by pLink (39) for identification of cross-linked peptides. The mass accuracies were specified as 10 ppm for MS and 50 ppm for MS/MS, respectively. Other search parameters included cysteine carboxymethylation as a fixed modification and methionine oxidation as a variable modification. A maximum of two trypsin missed-cleavage sites was allowed.

STORM microscopy. The samples for STORM imaging were labeled using the same immunofluorescent staining method described above. 39, 67, 83, and RAD52 was labeled by Alexa Fluor 647; TA-KR was labeled by Cy3B. Alexa Fluor 647 conjugated donkey anti-mouse antibody and Cy3B-conjugated donkey anti-rabbit antibody, which were synthesized in our laboratory (1:5000), were used for two-color STORM imaging. The two channels were imaged sequentially at 200 frames. The reconstruction of the super-resolution image was performed using our custom program written in Matlab 2015 (MathWorks) 41. Multi-color fluorescence beads (TetraSpeck microspheres, 0.1 µm diameter, blue/green/orange/dark red fluorescence, Fisher Scientific) were used to correct the chromatic aberration error across different color channels.

Data availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author upon request. The proteomics data of chemical crosslink and mass spectrometric analysis (CX-MS) analysis has been deposited into MIRVE data repository and the accession code is MSV000082876.

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