The cooperative action of CSB, CSA, and UVSSA target TFIIH to DNA damage-stalled RNA polymerase II

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The response to DNA damage-stalled RNA polymerase II (RNAPII) involves the assembly of the transcription-coupled repair (TCR) complex on actively transcribed strands. The function of the TCR proteins CSB, CSA and UVSSA and the manner in which the core DNA repair complex, including transcription factor IIH (TFIIH), is recruited are largely unknown. Here, we define the assembly mechanism of the TCR complex in human isogenic knockout cells. We show that TCR is initiated by RNAPII-bound CSB, which recruits CSA through a newly identified CSA-interaction motif (CIM). Once recruited, CSA facilitates the association of UVSSA with stalled RNAPII. Importantly, we find that UVSSA is the key factor that recruits the TFIIH complex in a manner that is stimulated by CSB and CSA. Together these findings identify a sequential and highly cooperative assembly mechanism of TCR proteins and reveal the mechanism for TFIIH recruitment to DNA damage-stalled RNAPII to initiate repair.
Nucleotide excision repair (NER) is a versatile DNA repair pathway that removes a wide range of helix-distorting DNA lesions from the genome, including ultra-violet (UV) light-induced photolesions. Transcription-coupled repair (TCR) is a specialized NER sub-pathway that specifically removes DNA lesions from actively transcribed DNA strands. It is believed that the TCR pathway is initiated by the stalling of elongating RNA polymerase II (RNAPII) at DNA lesions, which triggers the recruitment of the core NER machinery to repair these lesions. After lesion recognition, the transcription factor TFIH (TFIIF) complex is recruited to unwind the DNA, followed by dual incision, and the release of a 22–30 nucleotide-long DNA strand containing the lesion. The generated single-stranded DNA gap is filled by repair synthesis and the nick is sealed. However, the mechanism through which TCR recognizes transcription-blocking lesions and recruits the repair machinery is unknown.

Inherited defects that selectively impair TCR give rise to Cockayne Syndrome (CS) and UV-sensitive syndrome (UVS). Although cells from both CS and UVS patients show a defect in TCR, the phenotypes associated with these disorders are very different. CS is characterized by severe and progressive neurodegeneration, while UVS is characterized by mild UV sensitivity. The majority of CS patients carry mutations in the CSB or CSA genes, while UVS patients carry mutations in the UVSSA gene.

The CSB protein contains a central SWI2/SNF2-like DNA-dependent ATPase domain, and resides in a complex with RNAPII. Live-cell imaging suggests that CSB monitors the progression of transcription elongation by continuously probing RNAPII complexes. It has been suggested that CSB is involved in the repositioning of RNAPII to make the DNA less accessible for repair proteins. Although the association of CSB with RNAPII is sufficient to recruit TFIH in vitro, it is unknown whether additional factors are required to trigger the recruitment of the repair machinery in vivo.

Like CSB, the CSA and UVSSA proteins also associate with DNA damage-stalled RNAPII. The CSA protein contains seven WD40 repeats that form a seven-bladed β-propeller. Earlier work has shown that CSA is incorporated into a DDB1-CUL4-based E3 ubiquitin ligase complex that becomes transiently activated in response to UV irradiation and targets CSB for proteasomal degradation. Furthermore, the CSA complex also targets the UV-induced transcription repressor ATF3 as a means to regulate transcription restart after UV irradiation. Current models suggest that CSA is dispensable for the recruitment of the excision repair machinery to stalled RNAPII, and that CSA is unlikely to recruit UVSSA to sites of UV-induced DNA damage. Thus, the precise recruitment mechanism and the role of CSA in TCR is currently not clear.

The UVSSA protein contains an N-terminal VHS domain and a C-terminal DUF2043 domain of unknown function. Several studies reported that UVSSA, likely through its binding partner USP7, protects CSB from UV-induced degradation. However, ectopic expression of CSB in UVSSA-deficient cells did not rescue TCR, suggesting that UVSSA has additional functions in this repair mechanism. Moreover, UVSSA was found to associate with RNAPII, but whether UVSSA is constitutively bound to RNAPII, or associates with DNA damage-stalled RNAPII through either CSA or CSB is still a topic of debate.

The TFIH complex consists of seven core subunits, including the XPB and XPD helicases, and three Cdk kinase subunits. While the Cdk complex is crucial during transcription initiation, it inhibits the XPB helicase activity required for repair. The release of the Cdk complex from core TFIH is triggered by the association of repair factors XPA and XPG, which switches TFIH from a transcription factor into a repair factor.

Despite the knowledge that CSB, CSA, and UVSSA are required for TCR, we still know very little about how the interplay between these proteins targets the core repair machinery, including TFIH, to DNA damage-stalled RNAPII. In this study, we demonstrate a sequential and highly cooperative assembly of TCR proteins and unveil the mechanism for TFIH recruitment to DNA damage-stalled RNAPII.

**Results**

**Isolation of active TCR complexes under native conditions.** Our current understanding of the assembly and functioning of multi-protein complexes that mediate transcription-coupled DNA repair (TCR) is fairly limited. This is largely due to a lack of sensitive methods to isolate active TCR complexes and analyze their composition. To overcome this limitation, we set out to establish a new immunoprecipitation-based method to isolate the elongating form of RNAPII (RNAPIIo) and associated proteins from the chromatin fraction of UV-irradiated cells under native conditions (Fig. 1a). To this end, we employed extensive benzyl-xylene treatment to solubilize the chromatin fraction after centrifugation, followed by immunoprecipitation using antibodies that recognize the Ser2-phosphorylated form of RNAPII. This RNAPII modification is absent from transcription start sites (TSS), but increases across gene bodies and is associated with transcription elongation. Immunoprecipitation of RNAPIIo revealed a UV-specific association with the CSB and CSA proteins, as well as with CUL4, DDB1 and RBX1, which are subunits of a DDB1-CUL4 (CRL4) E3 ubiquitin ligase complex that associates with CSA. In addition, several subunits of the TFIH complex (XPD/p80, XBP/p89, GTF2H1/p62, and CDK7) also associated with RNAPIIo after UV irradiation (Fig. 1b).

**CSA is recruited to DNA damage-stalled RNAPII by CSB.** To acquire more insights into the initial assembly of TCR factors, we generated CSB, CSA, and UVSSA knockout (KO) cells using CRISPR-Cas9-mediated genome editing in U2OS cells equipped with the Flp-In/T-REx system. The knockout of CSB, CSA, and UVSSA was confirmed by western blot analysis and/or DNA sequencing (Fig. 1d). Clonogenic survival assays revealed that all TCR-KO cells were highly sensitive to transcription-blocking DNA damage induced by Illudin S (Fig. 1e), which is a natural compound from mushroom O. illudens causing DNA lesions that are exclusively repaired by TCR.
Fig. 1 CSA is recruited to DNA damage-stalled RNAPIIo by CSB. a Outline of a new IP method to isolate RNAPIIo and associated proteins from mock-treated or UV-irradiated (20 J/m²) U2OS (FRT) cells. b Endogenous RNAPII Co-IPs on WT cells stained for the indicated TCR proteins. Note that it is not possible to stain for all these proteins on one membrane. This panel is a composite of several representative Co-IPs. See Supplementary Fig. 1a for each individual Co-IP. c Endogenous RNAPII Co-IP followed by slot blot analysis of CPDs. d Western blot analysis of CSA, CSB, and UVSSA knockout cells complemented with inducible GFP-tagged versions of these proteins (n = 2). See Supplementary Fig. 2a, b for validation of knockouts by sequencing. e Clonogenic Illudin S survival of WT, CSA, CSB, and UVSSA knockout and rescue cell lines. Each symbol represents the mean of an independent experiment (n = 2 for all except for WT in UVSSA-KO figure which is n = 3) each experiment contains two or three technical replicates. Endogenous RNAPII Co-IP on f WT, CSA, CSB, and UVSSA knockout cells, g CSB-KO stably expressing GFP-CSB, and h WT, XPC, CSA, CSB, and UVSSA knockout cells. The asterisk in e indicates the heavy chain of the RNAPII antibody. At least two independent replicates of each IP experiment were performed obtaining similar results. Source data are provided as a Source Data file.
Importantly, complementation of these TCR-KO cells with inducible GFP-tagged versions of CSB, CSA, and UVSSA fully restored their resistance to Illudin S (Fig. 1d, e). We next applied our immunoprecipitation-based method in the different TCR-KO cells to establish how CSB and CSA recruitment to DNA damage-stalled RNAPIIo is regulated. CSB associated with RNAPIIo in wild-type (WT), CSA-KO, and UVSSA-KO cells specifically after UV irradiation, suggesting that CSB is the first of these proteins to associate with DNA damage-stalled RNAPIIo (Fig. 1f). The association of CSA with stalled RNAPIIo was abolished in CSB-KO cells, but was not affected in cells lacking UVSSA (Fig. 1f). Importantly, re-expressing GFP-tagged CSB in the CSB-KO cells restored the association between RNAPIIo and CSA (Fig. 1f, g). Next, we asked whether CSA mediates the recruitment of the CRL4 complex, including its E3 ubiquitin ligase partner DDB1 to DNA damage-stalled RNAPIIo. As an additional control we also included XPC-KO cells, which are deficient in global genome repair (GGR; Supplementary Fig. 2c). Immunoprecipitation of RNAPIIo revealed a UV-specific interaction with DDB1 in WT, XPC-KO, and UVSSA-KO cells (Fig. 1h). However, this interaction was completely abolished in CSA-KO and CSB-KO cells, showing that CSA indeed mediates the recruitment of the CRL4 complex to lesion-stalled RNAPIIo (Fig. 1h).

Mapping the CSA-interaction motif (CIM) in CSB. In order to gain a better understanding of the CSA recruitment mechanism by CSB, we aimed to identify the region in CSB that is required for the interaction with CSA. To this end, we employed a chromatin-tethering approach making use of the U2OS 2-6-3 cell line harboring an integrated LacO array in the genome39. This cell line enables the analysis of protein–protein interactions by tethering proteins of interest fused to the bacterial LacR and fluorescent protein mCherry to a defined chromosomal region40,41 (Fig. 2a). Expression of mCherry-LacR fused to full-length CSB (Fig. 2b) resulted in clear localization of the fusion protein to the LacO array and triggered the robust recruitment of CSA-GFP (Fig. 2c). In contrast, expression of LacR alone failed to recruit CSA-GFP to the LacO array (Fig. 2c).

To identify the CSA-interaction domain in CSB, we fused various truncated fragments of CSB to mCherry-LacR and examined their ability to recruit CSA-GFP to the LacO array (Fig. 2b, Supplementary Figs. 3, 4). Fragments of CSB spanning the N-terminus or the central region containing the conserved ATPase/helicase domain (N, M, and AC) were unable to recruit CSA-GFP. Conversely, tethering of a LacR-tagged CSB region spanning the C-terminus (C and AN) triggered robust recruitment of CSA-GFP (Fig. 2b–d, Supplementary Fig. 3). These results suggest that the C-terminus of CSB is essential for the interaction with CSA. The C-terminus of CSB contains a ubiquitin-binding domain (UBD; 1400–1428) and a recently identified winged-helix domain (WHD; 1417–1493) that interacts with RIF143. Interestingly, we found that the most N-terminal region (1221–1305) of the CSB C-terminus alone, or fragments containing solely the UBD (1400–1493) or WHD (1417–1493) domains do not support CSA recruitment. However, a region just upstream of the UBD (1306–1399) is sufficient to mediate CSA recruitment to the LacO array (Fig. 2b–d, Supplementary Fig. 3). Importantly, we found that tethering full-length CSB lacking this minimal interaction region (Δ1306–1399) indeed failed to support CSA recruitment (Fig. 2b–d). Further deletion analysis showed that CSB lacking the region just upstream of the UBD (1353–1399) failed to recruit CSA-GFP, whereas CSB lacking the UBD (1400–1428) or amino acids 1306–1352 were fully proficient in interacting with CSA-GFP (Supplementary Fig. 4). Moreover, while CSBΔ1353–1368 and CSBΔ1369–1384 were fully proficient in recruiting CSA-GFP to the LacO array, deleting amino acids 1385–1399 abolished the ability of CSB to interact with CSA-GFP (Fig. 2b–d, Supplementary Fig. 4). These findings identify an evolutionarily conserved CSA-interaction motif (CIM) in CSB that is located between amino acids 1385–1399 (Fig. 2e; Supplementary Fig. 5).

The C-terminal CIM in CSB recruits CSA to DNA damage-stalled RNAPIIo. We next set out to address the importance of this new CSB motif under more physiological conditions. To this end, we stably expressed GFP-tagged CSBWT or CSBΔCIM in CSB-KO cells (Fig. 3a, b). Pull-down of GFP-tagged CSBWT showed a strong UV-induced interaction with CSA, which was virtually absent after pull-down of CSBΔCIM even though equal amounts of CSB were immunoprecipitated (Fig. 3c). These findings were confirmed by quantitative mass spectrometry (MS) after pull-down of GFP-tagged versions of either CSBWT or CSBΔCIM (Supplementary Fig. 6a–c, see Supplementary Table 7 for link to interactive volcano plots). Immunoprecipitation of endogenous RNAPIIo in these cell lines showed that both CSBWT and CSBΔCIM associated equally with RNAPIIo after UV irradiation. However, CSBΔCIM failed to recruit CSA to DNA damage-stalled RNAPIIo, while a strong association of CSA was observed in cells expressing CSBWT (Fig. 3d). Importantly, the stable expression of GFP-CSBΔCIM in CSB-KO cells failed to restore sensitivity to Illudin S, while expression of GFP-CSBWT almost fully rescued this phenotype (Fig. 3e). To determine whether the CIM can mediate a functional interaction between CSB and CSA, we mixed recombinant Xenopus laevis CSBWT or CSBΔCIM with ubiquitin, E1, E2, and the E3 ubiquitin ligase CRL4CSA consisting of Xenopus laevis CSA, DDB1, CUL4A, and RBX1 (Supplementary Fig. 6d). While xICRL4CSA promoted the efficient ubiquitylation of xCSBWT, it did not ubiquitylate xCSBΔCIM (Fig. 3f). These data suggest that xCSB uses its CIM to interact directly with xCSA. Consistent with this interpretation, immobilized xCSBWT but not xCSBΔCIM interacted with endogenous xCSA from Xenopus egg extract (Fig. 3g). Similar results were observed when xCSB was substituted with hsCSB (Fig. 3f, g). Collectively, these data demonstrate that CSA is recruited to DNA damage-stalled RNAPIIo by CSB through direct interactions with the newly identified C-terminal CIM in CSB.

UVSSA is recruited to DNA damage-stalled RNAPIIo by CSA. Previous studies have demonstrated that UVSSA associates with RNAPIIo, but due to conflicting results, it remains unclear if UVSSA recruitment to RNAPIIo is enhanced by UV irradiation and dependent on the CS proteins17,25,31. Therefore, we monitored GFP-UVSSA recruitment to RNAPIIo in UVSSA-KO cells complemented with GFP-UVSSA (WT) in which we additionally knocked out either CSB or CSA. The knockout of CSB and CSA was verified by western blot analysis, DNA sequencing (Fig. 4a; Supplementary Fig. 2), and Illudin S clonogenic survival assays (Fig. 4b). Immunoprecipitation of endogenous RNAPIIo in these cell lines showed that GFP-UVSSA became readily detectable after UV irradiation in WT cells, whereas this interaction was virtually absent in CSA-KO and CSB-KO cells (Fig. 4c). Thus, GFP-UVSSA is targeted to DNA damage-stalled RNAPIIo in a manner that is dependent on the CS proteins17. Moreover, pull-down of GFP-UVSSA confirmed a robust UV-induced association with RNAPIIo, CSB, and CSA. However, these UV-specific interactions were abolished in CSB-KO and CSA-KO cells. Interestingly, we detected a weak UV-independent interaction between GFP-UVSSA and CSA, which was enhanced after UV irradiation in a manner that required CSB (Fig. 4d). These
findings suggest that the cooperative assembly of the TCR complex is important to mediate efficient targeting of UVSSA to lesion-stalled RNAPIIo.

CSB and CSA are required for the recruitment of the TFIIH complex. CSB, CSA, and UVSSA can each associate with TFIIH23,26,44, but which of these proteins is responsible for the recruitment of TFIIH to DNA damage-stalled RNAPIIo is currently unknown. To directly assess if CSB and CSA are required for the recruitment of TFIIH, we monitored TFIIH (p62 and p89) recruitment in UVSSA-KO complemented with GFP-UVSSA (WT) in which we additionally knocked out either CSB or CSA. Immunoprecipitation of endogenous RNAPIIo revealed a UV-specific interaction with TFIIH in WT cells, while these interactions were severely reduced in the CSB-KO and CSA-KO cells (Fig. 5a). Interestingly, TFIIH also failed to associate with RNAPIIo in CSB-KO cells complemented with GFP-CSBΔCIM (Supplementary Fig. 6e), consistent with our findings that this mutant is not capable of recruiting CSA (Fig. 3c, g). These initial results suggest that the TFIIH complex is recruited in a manner that requires both CS proteins.

UVSSA targets the TFIIH complex to DNA damage-stalled RNAPIIo. It has been reported that UVSSA can interact with TFIIH16,32,44, but whether this reflects a constitutive interaction or a UV-induced association is unclear. To gain more insight into the nature of this interaction, we immunoprecipitated GFP-UVSSA from the solubilized chromatin fraction of mock-treated and UV-irradiated cells followed by mass spectrometry (MS). Following UV irradiation, our MS analysis identified 28 UV-specific UVSSA interactors, including CSB, the CSA-interacting protein DDB1, and RNAPII subunits.
Additionally, among the most prominent UV-specific interactions were the TFIIH subunits XPB/p89 and XPD/p80 (Fig. 5b; see Supplementary Table 7 for link to interactive volcano plots). These findings demonstrate that UVSSA interacts in a UV-specific manner with TFIIH.

Immunoprecipitation of GFP-UVSSA indeed confirmed a UV-specific interaction with TFIIH subunits by western blot analysis (Fig. 5c). Strikingly, these interactions were severely reduced in the CSB-KO and CSA-KO cells, suggesting a cooperative interaction mechanism in which CSB is required to stabilize the interaction.
interaction between CSA and UVSSA, while CSA is required to stabilize the interaction between UVSSA and TFIIH.

We subsequently asked if UVSSA is also required for TFIIH recruitment. To this end, we employed our immunoprecipitation-based method in CSB-KO, CSA-KO, and UVSSA-KO cells to monitor TFIIH recruitment. In addition, we included XPA-KO cells (Supplementary Fig. 2c) as a positive control since XPA recruitment, at least during GGR, occurs downstream of TFIIH45.

Immunoprecipitation of endogenous RNAPIIα in these cell lines revealed a UV-specific interaction with TFIIH in WT and XPA-KO cells (Fig. 5d). These findings suggest that XPA recruitment does not only occur downstream of TFIIH in GGR but also in TCR. Interestingly, similar to CSB-KO and CSA-KO cells, we found that the UV-induced interaction between RNAPIIα and TFIIH was severely reduced in UVSSA-KO cells (Fig. 5d).

Furthermore, complementation of these TCR-KO cells with inducible GFP-tagged versions of CSB, CSA, and UVSSA fully restored the UV-induced association of TFIIH to RNAPIIα (Fig. 5e). These findings demonstrate that CSB, CSA, and UVSSA are equally important for the recruitment of the TFIIH complex to DNA damage-stalled RNAPIIα.

Genome-wide XR-seq confirms that UVSSA is a core TCR factor. Our findings show that UVSSA, just like CSA and CSB, is required to recruit TFIIH to initiate TCR-mediated repair.
provide further support for a role of UVSSA in TCR, we carried out genome-wide XR-sequencing (XR-seq), which enables the generation of genome-wide repair maps by isolating and sequencing the 30-mers that are generated upon dual incision.46,47 We generated nucleotide-resolution maps of UV-induced CPDs repair in U2OS WT cells (Fig. 5f; Supplementary Fig. 7a; Supplementary Table 8), which revealed that CPD repair under these conditions is enriched on the transcribed strands within gene bodies consistent with TCR-mediated repair.46 Importantly, the bias in CPD repair observed in transcribed strands was completely lost in both CSA-KO (Supplementary Fig. 7a) and UVSSA-KO cells (Fig. 5f). These findings provide...
Fig. 5 CSA, CSB, and UVSSA are equally important for TFIIH recruitment. a Endogenous RNAPII Co-IP in UVSSA-KO, UVSSA/CSA-dKO, and UVSSA/CSB-dKO complemented with GFP-UVSSA. b Volcano plot depicting the statistical differences of the MS analysis on GFP-UVSSA pull-down in mock-treated and UV-irradiated samples. The enrichment (log2) is plotted on the x-axis and the significance (t-test -log p value) is plotted on the y-axis. All significantly UV-induced hits are indicated in green. Several selected hits are shown in red (Link to the interactive volcano plots: GFP-UVSSA vs GFP-UVSSA + UV). c Co-IP of GFP-UVSSA in UVSSA-KO and UVSSA-dKO cells complemented with GFP-UVSSA. d Endogenous RNAPII Co-IP in WT, CSB-KO, CSA-KO, UVSSA-KO, and XPA-KO cells. e Endogenous RNAPII Co-IP in WT and UVSSA-KO cells and CSA-KO, CSB-KO, and UVSSA-KO cells complemented with GFP-tagged versions of these proteins. The asterisk in d and e indicates the heavy chain of the RNAPII antibody. f CPD XR-seq repair signal 3 Kb upstream and 5 Kb downstream of the annotated TSS of 16.088 genes in WT and UVSSA-KO cells. Signal is plotted separately for the transcribed (red) and non-transcribed (black) strands. The data represent the average of two independent experiments with a bin size of 40 nt. See also Supplementary Fig. 7a for additional XR-seq data. At least two independent replicates of each IP experiment were performed obtaining similar results. Source data are provided as a Source Data file.

Discussion

Although it has been recognized for some time that CSA, CSB, and UVSSA are required for TCR, remarkably little is known about how these proteins cooperate to trigger TCR. Our findings suggest a highly cooperative recruitment mechanism that involves the sequential association of CSA, CSB, and UVSSA to target the TFIIH complex to DNA damage-stalled RNAPII to initiate DNA repair (Fig. 7c).

We show that both CSA and CSB associate with RNAPII in a manner that is strongly induced by UV irradiation, and that CSA recruitment is completely dependent on CSB. This is in line with earlier work showing that CSB facilitates the translocation of CSA to the nuclear matrix after UV irradiation48. Moreover, we demonstrate that CSA is required for the association of DDB1 with RNAPII, suggesting that CSA is recruited to DNA damage-stalled RNAPII as part of a CRL4CSA complex24,27. Previous findings suggested that CSB dynamically associates with RNAPII under undamaged conditions and that this interaction is stabilized upon UV irradiation21,49. While our method may not be sensitive enough to capture these transient interactions, our findings do support that the CSB-RNAPII interaction is stabilized after UV irradiation.

Earlier observations suggested that CSB physically interacts with CSA26,28, while other studies failed to detect this association19,20. Our findings fully support a direct UV-induced association between the CS proteins. Importantly, we identified the CIM in the C-terminus of CSB that is essential for targeting CSA to stalled RNAPII. Interestingly, the CIM region in CSB is evolutionary conserved in species that also contain the CSA gene, including mammals, amphibians and fish (Supplementary Fig. 5).
In line with this, we demonstrate that both human and *Xenopus* leavis CSB require its CIM to directly interact with CSA in vitro. However, the CIM is absent in species without CSA, including yeast, nematodes, but also holometabolous insects, which have lost the CSA gene during the course of evolution (Supplementary Fig. 5).

It is striking that even though CSB contains a CIM, the association between these proteins is induced by UV irradiation. Previous studies revealed that the association of CSB with lesion-stalled RNAPII triggers a conformational change that repositions the N-terminus, thereby exposing residues in the C-terminus of CSB\(^\text{49}\). It is conceivable that this conformational

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**Fig. 6** UVSSA is the key protein that recruits TFIIH. **a** A schematic representation of UVSSA WT and deletion mutants. The CSA-interacting region (CIR) and TFIIH-interacting region (TIR) are indicated. **b** Recruitment of CSA-GFP and TFIIH (p89) to the LacO array upon tethering of the indicated mCherry-LacR fusion proteins (scale bar = 5 µm). **c** Quantification of CSA-GFP and endogenous TFIIH (p89) co-localization at the LacO array. Each symbol represents the mean of an independent experiment (\(n = 2\), >50 cells collected per experiment). **d** Western blot analysis of U2OS (FRT) and UVSSA-KO cells complemented with GFP-UVSSA\(^\text{WT}\), GFP-UVSSA\(^\text{ΔCIR}\), and GFP-UVSSA\(^\text{ΔTIR}\) (\(n = 2\)). **e** Co-IP of GFP-UVSSA\(^\text{WT}\), GFP-UVSSA\(^\text{ΔCIR}\), and GFP-UVSSA\(^\text{ΔTIR}\) cell lines. At least two independent replicates of each IP experiment were performed obtaining similar results. See also Supplementary Fig. 7 for additional Co-IP data. Source data are provided as a Source Data file.
change exposes the CIM to facilitate efficient CSA recruitment. Interestingly, while the CIM is located right next to the UBD in CSBΔUBD, we find that CSBΔUBD is fully functional in interacting with CSA. However, it is possible that the CIM and the UBD collaborate, as a tandem protein-interaction module, to enable optimal CSA recruitment. In this scenario, CSA would have protein–protein interactions with the CIM, which would be stabilized by the binding of the UBD to auto-ubiquitylated CSA.

The recently identified UVSSA protein can be isolated as part of a chromatin-bound stalled RNAPIIo complex. Our current findings shed light on its recruitment mechanism by demonstrating that the association of UVSSA with RNAPIIo is strongly induced by UV irradiation and fully dependent on both CSA and CSB. Moreover, knockout of UVSSA did not affect CSA or CSB recruitment to DNA damage-stalled RNAPIIo, suggesting that UVSSA is the last of these proteins to be recruited. Consistent with a reported association between CSA and UVSSA, we find that CSA targets UVSSA to DNA damage-stalled RNAPIIo by interacting with a region in the N-terminal VHS domain (CIR; amino acids 100–200) of UVSSA. Intriguingly, the robust UV-induced association between CSA and UVSSA is stabilized by CSB, suggesting a cooperative assembly mechanism of the TCR complex.

In contrast to our observation that the CS proteins are required for the recruitment of UVSSA to DNA damage-stalled RNAPIIo, live-cell imaging experiments showed that UVSSA is recruited to sites of UV-C-induced laser damage independently of the CS proteins. There could be several reasons for these seemingly conflicting results. Firstly, the methodology is very different. We isolate RNAPIIo-associated TCR proteins from the chromatin-bound fraction after UV, while live-cell imaging studies monitor the recruitment of GFP-tagged TCR proteins to local UV-C laser

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**Fig. 7 The CIR and TIR of UVSSA are crucial for TCR.**

a Clonogenic Illudin S survival of WT and UVSSA-KO cell lines and the GFP-tagged UVSSA rescue cell lines. Each symbol represents the mean of an independent experiment (n = 3 for all except for GFP-UVSSAΔCIR which is n = 2), each of which is based on two technical replicates. Note that the same survival data for UVSSA-KO is also shown in Fig. 4b. b CPD XR-seq repair signal 3 Kb upstream and 5 Kb downstream of the annotated TSS of 16,088 genes in GFP-UVSSAWT, GFP-UVSSAΔCIR, and GFP-UVSSAΔTIR. Signal is plotted separately for the transcribed (red) and non-transcribed (black) strands. The data represent the average of two independent experiments for GFP-UVSSAWT and one single experiment for GFP-UVSSAΔCIR and GFP-UVSSAΔTIR with a bin size of 53 nt. See also Supplementary Fig. 9 for additional XR-seq data. c Model of how the assembly of CSB, CSA, and UVSSA targets the TFIIH complex to DNA damage-stalled RNAPIIo. Source data are provided as a Source Data file.
damage. Therefore, it is possible that the observed recruitment of CSB and UVSSA could, in part, be triggered by something other than DNA damage-stalled RNAPII. In line with this hypothesis, GFP-CSA could not be detected at sites of local UV-C laser damage, even though CSA is essential for TCR and showed a robust UV-specific association with RNAPII under our conditions. Secondly, the time frame during which UVSSA association is measured is different. While we isolate RNAPII-associated UVSSA 1 h after UV irradiation, the recruitment studies visualized UVSSA binding in the first 40 s after UV-C laser irradiation. It cannot be excluded that UVSSA transiently associates with UV-damaged chromatin independently of the CS proteins, but that the stable association with stalled RNAPII during productive TCR is fully dependent on CSA and CSB. In line with this, we find that mutants of TCR proteins that display a clear assembly defect under our conditions also show a strong sensitivity to Icladin S reflecting impaired TCR. In conclusion, our findings favor a model in which UVSSA is recruited by CSA and argues for a cooperative assembly mechanism in which CSB stabilizes the association between CSA and UVSSA to ensure efficient targeting to stalled RNAPII.

A major unresolved question is how the core NER machinery, likely starting with the TFIIH complex, is recruited to DNA damage-stalled RNAPII to initiate repair. In vitro experiments have shown that the association of CSB with RNAPII is sufficient to recruit TFIIH. In addition, CSA was shown to associate with the p44 subunit of TFIIH, while UVSSA can interact with the p62 subunit of TFIIH. In agreement, we found that GFP-USVSSA associates with several subunits of the TFIIH complex in a UV-specific manner in vivo. Furthermore, our data reveals that CSB, CSA, and UVSSA are equally important for the recruitment of TFIIH to DNA damage-stalled RNAPII in vivo. Indeed, similar to previous results with CSB-deficient cells, our high-resolution repair maps fully support a crucial role of both CSB and UVSSA in the TCR-mediated clearing of UV-induced lesions on a genome-wide level. Importantly, we found that UVSSA contains a TFIIH-interacting region (TIR; amino acids 400-500), which is crucial for the association of TFIIH with DNA damage-stalled RNAPII. Consistently, it has been shown that the PH domain of p62 (1-108) associates with a small fragment in UVSSA (400-419) in vitro and that mutations within this region causes a defect in recovery of RNA synthesis in vivo. Moreover, we found that the UVSSAACR mutant was not only unable to associate with CSA, but also with the TFIIH complex. Our findings favor a model in which CSA not only recruits UVSSA to stalled RNAPII but also stabilizes the direct interaction between UVSSA and TFIIH, resulting in the recruitment of TFIIH to DNA damage-stalled RNAPII. In this regard, it would be interesting to examine if this interaction between UVSSA and the p62 subunit of TFIIH is the sole mechanism through which TFIIH is recruited to DNA damage-stalled RNAPII in vivo, or whether other subunits and regions also contribute.

Here we show that UVSSA is essential to bridge the TFIIH complex to CSB/CSA-bound RNAPII to initiate TCR. Importantly, these findings also suggest that neurodegeneration seen in CS is not caused by the inability to remove transcription-blocking DNA lesions, since neurodegeneration is not a feature in UVSSA. Previous findings revealed that CS fibroblasts fail to ubiquitylate and subsequently degrade DNA damage-stalled RNAPII, while UVSSA fibroblasts displayed even faster degradation of RNAPII after UV, possibly due to a failure to ubiquitylate RNAPII by the UVSSA binding partner USP7. These findings suggest that features of CS are caused by toxicity associated with prolonged RNAPII stalling at DNA lesions rather than the inability to remove transcription-blocking DNA lesions.

It remains unresolved how RNAPII is repositioned while TCR is in progress to enable access to the DNA lesion. We speculate that the TCR complex that we capture contains an inactive TFIIH complex, which is bound to the CAK complex. Indeed, we detect the association of CAK subunit CKD7 with RNAPII after UV. The association of repair factors XPA and XPG activates TFIIH, which could cause the backtracking, or even removal of RNAPII. This model would explain why we do not detect the accumulation of downstream repair factors with DNA damage-stalled RNAPII.

We propose a model in which CSB is the first protein to be recruited to DNA damage-stalled RNAPII (Fig. 7c). This binding of CSB could bring about a conformational change, thereby exposing the newly identified CIM to facilitate efficient CSA recruitment through direct protein–protein contacts. Once bound, CSA targets UVSSA to DNA damage-stalled RNAPII, and this interaction is stabilized by CSB. UVSSA, in turn, mediates the recruitment of the TFIIH complex in a cooperative manner that is stabilized by both CSB and CSA. Although both CSA and UVSSA could interact with TFIIH, it is likely that only CSA contributes directly to this stabilization, while CSB contributes indirectly through ensuring the association of CSA itself and stabilizing the interaction between CSA and UVSSA. At the stage when TFRH is bound, it is likely that RNAPII and CSB/CSA/UVSSA are displaced and that the TCR-specific pre-initiation complex is assembled starting with XPA. It is interesting to note that the yeast orthologue of CSB, RAD26, is bound to the DNA upstream of RNAPII, while human TFIIH in the transcription pre-initiation complex is bound downstream of RNAPII. If TFIIH is recruited to the same side of RNAPII during TCR, it suggests that CSB/CSA/UVSSA extend from the upstream to the downstream DNA around RNAPII to position TFIIH. It will be very interesting to gain structural insights into these molecular events. In conclusion, our findings reveal the recruitment mechanism of the TFIIH complex to DNA damage-stalled RNAPII, which involves the sequential and cooperative assembly of the CSB, CSA, and UVSSA proteins.

**Methods**

**Cell lines.** Cell lines (listed in Supplementary Table 1) were cultured at 37 °C in an atmosphere of 5% CO₂ in DMEM (Thermo Fisher Scientific) supplemented with penicillin/streptomycin (Sigma) and 10% Fetal bovine serum (FBS; Bodinco BV). Sp cells were cultured in ESF 921 insect cell culture medium (Fisher Scientific). U2OS 2-6-3 cells containing 200 copies of a LacO-containing cassette (~4 Mbp) were a gift from Susan Janicki. UVSSA-deficient KPS3-hTERT cells and their UVSSA-rescued counterparts were a gift from Tomoo Ogi. U2OS Flp-In/T-Rex cells, which were generated using the Flp-In/T-Rex system (Thermo Fisher Scientific), were a gift from Daniel Durocher.

**Methods**

**Generation of knockout cell lines.** To generate stable knockouts, U2OS Flp-In/T-Rex cells were co-transfected with pLV-U6-VPFP encoding a guide RNA from the LUMC/Sigma-Aldrich sgRNA library (see Supplementary Table 2 for plasmids, Supplementary Table 3 for sgRNA sequences) together with an expression vector encoding Cas9-2A-GFP (pX458; Addgene #48138) using lipofectamine 2000 (Invitrogen). Transfected cells were selected on puromycin (1 μg/mL) for 3 days, plated at low density after which individual clones were isolated. To generate double knockouts, single knockout clones were transfected with pLV-U6-VPFP encoding a sgRNA together with pX458 encoding Cas9, cells were FACs sorted on GFP (GFP), plated at low density after which individual clones were isolated. Isolated knockout clones were verified by western blot analysis and/or sanger sequencing. The absence of Cas9 integration/stable expression was confirmed by western blot analysis.

**PCR analysis of knockout clones.** Genomic DNA was isolated by suspending cell pellets in WCE buffer (50 mM KCl, 10 mM Tris pH 8.0, 25 mM MgCl₂, 0.1 mg/mL gelatin, 0.45% Tween-20, 0.45% NP-40) containing 0.1 mg/mL Protease K (E00491; Thermo Fisher Scientific) and incubating for 1 h at 56 °C followed by a 10 mm heat inactivation of Protease K by 96 °C. Fragments of ~1 kb, containing the sgRNA sequence, were amplified by PCR (sequencing primers are listed in Supplementary Table 4) followed by Sanger sequencing using either the forward or the reverse primer.
Generation of stable cell lines. Selected knockout clones of CBX, CSA, and UVSSA (Supplementary Table 1) were passaged to stably express GFP-CSPβVT, GFP-CSβVT, CSAWT, GFP-UVSSAΔC, GFP-UVSSAΔΔC, and GFP-UVSSAΔΔD by co-transfection of pcDNAs/FRT/TO-Puro plasmid encoding these CBX, CSA, and UVSSA variants (2 μg), together with pOG44 plasmid encoding the Flp recombinase (0.5 μg). After selection on 1 μg/mL pur- emycin and 4 μg/mL blasticidin, single clones were isolated and expanded. Clones were confirmed by Sanger sequencing and were then compared with parental U2OS Flp-In/T-Rex cells. Expression of these GFP-tagged TCR proteins was induced by the addition of 2 μg/mL doxycycline for 24 h.

Plasmid constructs. The Neomycin resistance gene in pcDNAs/FRT/TO-Neo (Addgene #41000) was replaced with a Puromycin resistance gene. Fragments spanning GFP-N1 (onlatch) and GFP-C1 (onlatch) including the multiple cloning sites were inserted into pcDNAs/FRT/TO-Puro. CSPβVT, CSAWT, and UVSSAΔ were amplified by PCR (see Supplementary Table 5 for primers) and inserted into pcDNAs/FRT/TO-Puro-GFP-N1 or pcDNAs/FRT/TO-Puro-GFP- C1 and in m-Cherry-LacR-NSL-C1/G3. Deletion constructs of CBX and UVSSA were generated by site-directed mutagenesis PCR. All sequences were verified by sequencing.

Ilidin S survival assay. Knockout and rescue cell lines were trypsinized, seeded at low density, and mock-treated or exposed to a dilution series of Illidin S (Santa cruz; sc-391575) for 72 h (30, 60, 100 pg/mL or 50, 100, and 200 pg/mL). On day 10, the cells were washed with 0.9% NaCl and stained with methylene blue. Colonies of more than 20 cells were scored.

Immunoprecipitation for Co-IP. Cells were irradiated with UV-C light (20 μm2) or mock-treated and harvested 1 h after UV. Chromatin-enriched fractions were prepared by incubating the cells for 20 min on ice in IP buffer (IP-130 for endo- genous RNAPII IP and IP-150 for GFP-IP), followed by centrifugation, and removal of the supernatant. For Co-IP of endogenous RNAPII, the chromatin- enriched cell pellets were lysed in IP-130 buffer (30 mM Tris pH 7.5, 130 mM NaCl, 2 mM MgCl2, 0.5% Triton X-100, protease inhibitor cocktail (Roche), 250 U/mL Benzonase Nuclease (Novagen)), and 2 μg RNAPII-S2 (ab9595, Abcam) for 2–3 h at 4 °C. For GFP IPs, the chromatin-enriched cell pellets were lysed in IP-150 buffer (50 mM Tris pH 7.5, 150 mM NaCl, 0.5% NP-40, 2 mM MgCl2, protease inhibitor cocktail (Roche), and 500 U/mL Benzonase Nuclease (Novagen)), for 1 h at 4 °C. Protein complexes were pulled down with 1.5 h incubation with Protein A agarose beads (Millipore) or GFP-Trap A beads (Chromotek). For subsequent analysis by western blotting, the beads were washed six times with IP-130 buffer for endogenous RNAPII IP and EBC-2 buffer (50 mM Tris pH 7.5, 150 mM NaCl, 1 mM EDTA) for GFP-IPs. The samples were prepared by boiling in Laemmli-SDS sample buffer. Unless indicated otherwise, all IP experiments were performed on the chromatin fraction. At least two independent replicates of each IP experiment were performed.

Generation of mass spectrometry samples. For the generation of mass spectrometry samples, the beads were washed four times with EBC-2 buffer without NP-40 (1.5 ml each) and sonicated for 2 h followed by overnight digestion using 2.5 μg trypsin at 37 °C under constant shaking. The bead suspension was loaded onto a 0.45 μm filter column (Millipore) to elute the peptides. The peptides were passed through a C-18 stage tips for desalting. The stage tips were activated by washing with methanol followed by washing with buffer B (80% acetonitrile and 0.1% formic acid) and 1% formic acid. Peptides were acidified with 2% trifluoroacetic acid and loaded on the stage tips. The peptides were eluted twice with 25 μL 60% acetonitrile/0.1% formic acid and lyophilized. At least three biological repeats for each condition were grouped and only protein groups identified by site and reverse peptide sequences from default settings: the maximum number of Cleavages allowed, number of allowed modifications (fixed and variable), dynamic exclusion of proteins, and dynamic exclusion of peptides.

Mass spectrometry. Mass spectrometry analysis was performed on a Q-Exactive Orbitrap mass spectrometer (Thermo Scientific, Germany) coupled to an EASY- nanoLC 1000 system (Proxeon, Odense, Denmark). The mass spectrometer was operated in positive-ion mode at 2.9 kV with the capillary heated to 250 °C in a data-dependent acquisition mode with a top 7 method, Precursor ions with a charge state of one and greater than six were excluded from triggering MS/MS events. For the UV-dependent UVSSA interactors (Fig. 3b), samples were analyzed essentially as previously described25–27. Digested peptides were separated using a 15 cm fused silica capillary (ID: 75 μm, OD: 375 μm, Polymicro Technologies, California, US) in-house packed with 1.9 μm C18-AQ beads (Reprospher-DE, Pur, Dr. Maisch, Ammerbuch-Entringen, Germany). Peptides were separated by liquid chromatography using a gradient from 2% to 95% acetonitrile with 0.1% formic acid in water, through a 0.3 mm × 15 cm analytical column ( Bucksport, ME) and monitored in poly mount (Polysciences; 18606) or subjected to immuno- nfluorescent labeling.

Western blot. Proteins were separated on 4–12% Criterion XT Bis-Tris gels (Bio- Rad, #3450124) in NuPage MOPS running buffer (NP0001-02 Thermo Fisher Scientific) and blotted onto PVDF membranes (Bio-Rad). The membrane was blocked with blocking buffer (Rockland, MB-070-003) for 2 h at RT. The membrane was then probed with antibodies (listed in Supplementary Table 6) and subjected to ECL reagent (Sigma, GERP2232).

Slot blot for CPDs after IP. Cells were irradiated with UV-C light (20 μm2) or mock treated and crosslinked with 0.5 mg/mL dicyclosinimidyl glutarate (DSG; Thermo Fisher) in PBS for 45 min at room temperature. Cells were washed with PBS and crosslinked with 1% PFA for 20 min at room temperature. Fixation was stopped by adding 1.25 M glycine in PBS to a final concentration of 0.1 M for 3 min at room temperature. Cells were washed cold with PBS and lysed in a buffer containing 0.25% Triton X-100, 10 mM EDTA (pH 8.0), 0.5 mM EGTA (pH 8.0), and 20 mM Hepes (pH 7.6). Chromatin was pelleted in 5 min at 400 g and incubated for 1 h at room temperature. The chromatin was re-suspended in a buffer containing 0.5 mM EGTA (pH 8.0), and 50 mM Hepes (pH 7.6) for 10 min at 4 °C. Chromatin was again pelleted for 5 min at 400 g and resuspended in Chip-buffer (0.1% SDS, 1% Triton X-100, 150 mM NaCl, 1 mM EDTA (pH 8.0), 0.5 mM EGTA (pH 8.0), and 20 mM Hepes (pH 7.6)) to a final concentration of 20 × 106 cells/mL. Chromatin was sonicated to approximately one nucleosome length using a Bioruptor water bath sonicator (Diagenode). Chromatin of 20 × 106 cells was incubated with 6 μg RNAPII-S2 (ab9595, Abcam) overnight at 4 °C, followed by a 1.5 h protein- chromatin pull-down with a 1:1 mix of protein A and protein G Dynabeads (Thermo Fisher; 10001D and 10003D). The beads were washed extensively, followed by decrosslinking for 4 h at 65 °C in the presence of proteinase K and RNAse A. The DNA was then purified with concentrated acetone (Sigma) and the concentration was measured using a dsDNA HS Qubit assay (Thermo Fisher Scientific). 15 ng of DNA for each sample was denatured for 15 min at 98 °C and blotted on Hybond N+ (RP20385, GE healthcare). The membrane was dried for at least 1 h at 70 °C and blocked overnight in 5% milk in PBS – 0.1% tween. CPDs were detected using anti-CPD mouse monoclonal antibody followed by incubation with an anti-mouse HRP conjugated antibody (Supplementary Table 6). CPD signals were detected using an ECL reagent (Sigma, GERP2232).

Chromatin tethering. U2OS 2–6-3 cells containing 200 copies of a LacO- containing cassette were co-transfected with lipofectamine 2000 (Invitrogen) and plasmid DNA for 6 h at 37 °C in an atmosphere of 5% CO2, 24 h after transfection the cells were fixed with 4% paraformaldehyde (Sigma) in PBS for 15 min. The cells were rinsed once in PBS for 5 min and mounted in poly mount (Polysciences; 18606) or subjected to immunofluorescent labeling.

Mass spectrometry data analysis. Raw mass spectrometry files were analyzed with MaxQuant software (version 1.5.3.30) as described26, with the following modifications from default settings: the maximum number of missed cleavages was set to 4, Label Free Quantification (LFQ) was enabled thereby disabling the Fast LFQ feature. Match-between-runs feature was enabled with a match time window of 0.7 minutes and an alignment time window of 20 min. We performed the search against an in silico digested UniProt reference proteome for Homo sapiens (14th December 2017). Analysis output from MaxQuant was further processed in the Perseus (version 1.5.3.3) computational platform57. Proteins identified as common contaminants, only identified by site and reverse peptide sequences were filtered out, and then all the LFQ intensities were log2 transformed. Different biological repeats of each condition were grouped and only protein groups identified by site and reverse peptide sequences were considered for comprehensive visualization and analysis of the data. A web app (VolcaNoseR) was made with R/Shiny for generating and sharing interactive volcano plots. Links to the interactive volcano plots are listed in Supplementary Table 7 and in the relevant figure legends.

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Immunofluorescent labeling. Cells were permeabilized with 0.5% triton X-100 (Sigma) in PBS for 10 min, followed by treatment with 100 mM glycine in PBS for 10 min. All polyacrylamide gels were rinsed with PBS and equilibrated in wash buffer (WB: PBS containing 0.5% BSA, and 0.05% Tween-20 (Sigma-Aldrich)) for 10 min. Antibody stops and washes were in WB. The primary antibody rabbit-p89 (1/100; Santa Cruz; SC-293; S19) was incubated for 2 h at RT. Detection was done using goat-rabbit Ig coupled to Alexa 488 (1/1000; Invitrogen). Cells were incubated with 0.1 μg/mL DAPI and mounted in Poly mount (Polysciences; 18606).

Microscopic analysis of fixed cells. Images of fixed samples were acquired on a Zeiss Axioslager M2 or D2 widefield fluorescence microscope equipped with a 63x PLAN APO (1.4 NA) oil-immersion objectives (Zeiss) and an HXP 120 metal-halide lamp used for excitation. Fluorescent probes were detected using the following filters: DAPI (excitation filter: 350–380 nm, dichroic mirror: 400 nm, emission filter: 460–490 nm); GFP/Alexa 488 (excitation filter: 470/40 nm, dichroic mirror: 495 nm, emission filter: 525/50 nm), mCherry (excitation filter: 560/40 nm, dichroic mirror: 585 nm, emission filter: 630/70 nm). Images were recorded using ZEN 2012 software (blue edition, version 1.1.0.0).

Genome-wide XR-seq alignment. Cell lysis was performed using ZRN 2012 software (blue edition, version 1.1.0.0). Briefly, briefly, cells were harvested 3 hr after treatment with 20 J/m² UVC. Primary excision products were pulled down by TTHH immunoprecipitation (Santa Cruz Biotechnology sc25329 and sc271500), and ligated to both 2′ and 3′ adaptors. Ligation products containing CPD were purified by immunoprecipitation with the anti-CPD antibody (Cosmo Bio NM-DND-001) and repaired in vitro by Drosophila melanogaster GFP photolyase. Repaired DNA was PCR-amplified with Index primers and purified on Sephadex G-50. The polyacrylamide gels were poured and sequenced either on a HiSeq 2500 or on a NextSeq 500 to produce at least 4 million reads per sample. Quality score for each nucleotide was analyzed using the Fastx-toolkit to ensure only high-quality reads are processed. Adapter sequence was trimmed from each read using Trimmomatic (version 0.36)6. Reads were aligned to the genome using Bowtie (version 1.1.2)6. Following alignment reads that were mapped to chromosome Y or mitochondrial chromosome were filtered (U2OS cell line is derived from female bone tissue) and PCR duplicates were removed using Picard/CommandLineMarkDuplicates (version 2.8.1) (http://broadinstitute.github.io/picard/). There were high levels of PCR duplicates due to low efficiency of excision. The PCR duplicates were removed using the PRIdE partner repository6.2 (https://www.ebi.ac.uk/pride/) with the dataset identifiers PXD013572 and PXD017329. Processed mass spectrometry proteomics data used for interactive Volcano plots is deposited on Zenodo (https://doi.org/10.5281/zenodo.3625858). XR-seq data are deposited in main Figs. 5b, 7b, and Supplementary Figs. 5a, 5c. XR-seq data are deposited in the Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/) under GSE132840. Additional data and custom code will be made available upon reasonable request. The source data underlying Figs. 1b, d–h, 2b, d–h, 3b–d, 4a–d, 5a, 5c, 6b–f, and 7a and Supplementary Figs. 1a–c, 2c, 3c, 4d, 6b–e, and 7b–c are provided as a Source Data file.

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Protein expression and purification. Coding sequences of Xenopus laevis CSB and CSA-DDB1-CUL4A/RBX1 (CRL4CSA)64, as well as human CSB were amplified from cdNA clones or ordered as codon-optimized gene blocks from Integrated DNA Technologies. All open reading frames were cloned into pAcCBac vectors for insect expression, plasmids were transformed into chemically competent DH10Bac cells (Invitrogen) and PCR duplicates were removed using the PRIdE partner repository6.2 (https://www.ebi.ac.uk/pride/) with the dataset identifiers PXD013572 and PXD017329. Processed mass spectrometry proteomics data used for interactive Volcano plots is deposited on Zenodo (https://doi.org/10.5281/zenodo.3625858). XR-seq data are deposited in main Figs. 5b, 7b, and Supplementary Figs. 5a, 5c. XR-seq data are deposited in the Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/) under GSE132840. Additional data and custom code will be made available upon reasonable request. The source data underlying Figs. 1b, d–h, 2b, d–h, 3b–d, 4a–d, 5a, 5c, 6b–f, and 7a and Supplementary Figs. 1a–c, 2c, 3c, 4d, 6b–e, and 7b–c are provided as a Source Data file.

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Author contributions
Y.v.d.W. generated knockout cells, constructs and stable cell lines, performed LacR-based tethering assays, clonogenic survivals, PCR and western blot analysis to validate knockouts, Co-IP experiments for western blot analysis, Co-IP experiments for mass spectrometry, slot blot assays, and wrote the paper. K.A. generated stable cell lines, western blot analysis to validate knockouts, and Co-IP experiments. R.G.-P. and A.C.O.V. analyzed the mass spectrometry samples. H.G.-B. performed XR-seq. H.G.-B., E.E.H., and S.A. analyzed the XR-seq samples. J.G. developed VolcaNoseR. T.E.T.M. generated recombinant CSB proteins and xLCRL4CSA, and performed pull-down and in vitro ubiquitylation assays. D.v.d.H. generated knockout cells, constructs, and performed western blot analysis to validate knockouts, and Co-IP experiments. J.C.W. supervised T.E.T.M. M.S.L. supervised the project and wrote the paper.

Competing interests
The authors declare no competing interests.

Additional information

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