The PSO4 Protein Complex Associates with Replication Protein A (RPA) and Modulates the Activation of Ataxia Telangiectasia-mutated and Rad3-related (ATR)*

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Background: The function of the PSO4 complex in response to DNA damage remains unclear.

Results: The PSO4 complex is required for efficient accumulation of ATRIP at DNA damage sites and the subsequent CHK1 activation and RPA2 phosphorylation.

Conclusion: The PSO4 complex modulates ATR activation through interaction with RPA.

Significance: We reveal a role for RNA processing factor PSO4 in ATR activation.

The PSO4 protein complex is composed of PSO4/PRP19/SNEV, CDC5L, PLRG1, and BCAS2/SPF27. Besides its well defined functions in pre-mRNA splicing, the PSO4 protein complex has been shown recently to participate in the DNA damage response. However, the specific role for the PSO4 complex in the DNA damage response pathways is still not clear. Here we show that both the BCAS2 and PSO4 subunits of the PSO4 complex directly interact and colocalize with replication protein A (RPA). Depletion of BCAS2 or PSO4 impairs the recruitment of ATR-interacting protein (ATRIP) to DNA damage sites and compromises CHK1 activation and RPA2 phosphorylation. Moreover, we demonstrate that both the RPA1-binding ability of BCAS2 and the E3 ligase activity of PSO4 are required for efficient accumulation of ATRIP at DNA damage sites and the subsequent CHK1 activation and RPA2 phosphorylation. Our results suggest that the PSO4 complex functionally interacts with RPA and plays an important role in the DNA damage response.

The PSO4 core complex is composed of PSO4/PRP19/SNEV, CDC5L, PLRG1, and BCAS2/SPF27. Besides its well defined functions in pre-mRNA splicing, the PSO4 protein complex has been shown recently to play an important role in the DNA damage response.

Conclusion: The PSO4 complex modulates ATR activation through interaction with RPA.

Significance: We reveal a role for RNA processing factor PSO4 in ATR activation.

The PSO4 core complex is composed of PSO4/PRP19/SNEV, CDC5L, PLRG1, and BCAS2/SPF27 (5–11). This complex has been shown to have a role in pre-mRNA splicing from yeast to humans (12–17). The only identified catalytic center in the subunits of this complex is a U-box domain located in the N terminus of PSO4. U-box domains have been shown to have E3 ubiquitin ligase activity (18, 19). Several studies have shown that the E3 ubiquitin ligase activity of PSO4 is crucial for its function in pre-mRNA splicing (20–22). Besides its well defined roles in pre-mRNA splicing, the PSO4 complex has been found recently to play an important role in the DNA damage response (5, 14, 19, 23–26). In fact, cells harboring a mutant PSO4 in budding yeast showed broad hypersensitivity to DNA damage-inducing agents, suggesting that PSO4 is essential for mediating the DNA damage response (14, 26). Human PSO4 has also been shown to be a DNA-binding protein and plays a role in DNA repair through its interaction with terminal deoxynucleotidyl transferase (24). In addition, human PSO4 is required for the recruitment of the DNA repair protein Metnase to DNA damage sites (25). Moreover, CDC5L was found to directly interact with ATR and is required for the S phase cell-cycle checkpoint (27). Although these findings clearly indicate that the PSO4 complex participates in the DNA damage response, the nature of its function in these pathways is still not clear.

In this study, we adopted the tandem affinity purification approach to isolate an RPA-containing protein complex and

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identified the PSO4 complex as a component involved in RPA-mediated DNA damage response. We demonstrate that both the BCA(S2 and PSO4 subunits of the PSO4 complex directly interact with RPA1 and are recruited to DNA damage sites. Depletion of BCA(S2 or PSO4 leads to defects in the recruitment of ATRIP to DNA damage sites, CHK1 activation, and RPA2 phosphorylation. We further show that both the RPA1-binding ability of BCA(S2 and the E3 ligase activity of PSO4 are required for efficient accumulation of ATRIP at DNA damage sites and the subsequent CHK1 and RPA2 phosphorylation. We propose that the PSO4 complex, via its interaction with RPA, modulates the DNA damage response.

EXPERIMENTAL PROCEDURES

Plasmids—All cDNAs were subcloned into pDONR201 (Invitrogen) as entry clones and were subsequently transferred to gateway-compatible destination vectors for the expression of N- or C-terminal-tagged fusion protein. All deletion mutants were generated using the QuikChange site-directed mutagenesis kit (Stratagene) and verified by sequencing.

Cell Cultures, Transfection, and shRNAs—HEK293T and HeLa cells were maintained in DMEM supplemented with 10% fetal bovine serum and 1% penicillin and streptomycin. Sf9 insect cells were maintained in Grace’s medium supplemented with 2% fetal bovine serum. Human cell lines were maintained in a 37 °C incubator with 5% CO2, whereas insect cells were maintained at 27 °C. Cell transfection was performed using Lipofectamine 2000 (Invitrogen) following the protocol of the manufacturer. Lentiviral nonsilencing control shRNA and shRNA targeting sequences were as follows: 1, 5’TATTCTGCTTTG A-3; 2, 5’GCTACAAAGCGTTTCTTTA-3; 3, 5’CCCATAAGAGTAATAATAT-3’; 4, 5’GCTGTGGTAATTCTATTG-3’; 5, 5’GG-GAGTCAGATGGTCTGATA-3’. The nonsilencing control sequence was as follows: 5’-CCCTTAAGAGTATAATAT-3’. The PSO4 targeting sequences were as follows: 1, 5’-GGTAACGCGATGGGAAAGGAAAGAA-3’; 2, 5’-GCTGTCTC- TAATCATGTTTAT-3’. The RPA1 targeting sequences were as follows: 1, 5’-GCCATAAAGGTCTTTTATA-3’; 2, 5’-GAGTCAGATGGTCTGATA-3’. The shRNA-resistant wild-type and mutant PSO4 constructs were generated by changing seven nucleotides in the shRNA 2 targeting region (T57C, C60G, T63C, T66C, T69C, T72G, and T75C substitutions). The shRNAs were packaged into lentiviral vectors and transfected into HEK293T and HeLa cells. Infected cells were selected with medium containing puromycin (2 μg/ml) and confirmed by immunoblotting and immunostaining. For affinity purification, HEK293T cells stably expressing tagged proteins were lysed with NETN buffer (20 mM Tris-HCl (pH 8.0), 100 mM NaCl, 1 mM EDTA, and 0.5% Nonidet P-40) for 20 min. Crude lysates were removed by centrifugation at 14,000 rpm at 4 °C for 10 min, and the pellet was sonicated for 40 s in high-salt solution (20 mM HEPES (pH 7.8), 0.4 M NaCl, 1 mM EDTA, 1 mM EGTA, and protease inhibitor) to extract chromatin-bound protein fractions. The supernatants were cleared at 14,000 rpm to remove debris and then incubated with streptavidin-conjugated beads (Amersham Biosciences) for 2 h at 4 °C. The beads were washed three times with NETN buffer, and then bead-bound proteins were eluted with NETN buffer containing 1 mg/ml biotin (Sigma). The elutes were incubated with S protein beads (Novagen) for 2 h at 4 °C. The beads were then washed three times with NETN buffer and subjected to SDS-PAGE. Protein bands were excised and digested, and the peptides were analyzed by mass spectrometry.

Coimmunoprecipitation and Western Blotting—For whole-cell extracts, the cells were solubilized in NETN lysis buffer supplemented with 50 units/μl benzonase (Novagen), protease inhibitors, and phosphatase inhibitors. After removal of cell debris by centrifugation, the soluble fractions were collected. For FLAG immunoprecipitations, 2.0 ml aliquots of lysate were incubated with 1 μg of the FLAG monoclonal antibody and 25 μl of a 1:1 slurry of protein A-Sepharose for 2 h at 4 °C. For endogenous immunoprecipitations, 1 mg of the whole-cell extract was incubated with 25 μl of a 1:1 slurry of protein A-Sepharose coupled with 2 μl of the indicated antibodies for 2 h at 4 °C. The Sepharose beads were washed three times with NETN buffer, boiled in 2× SDS loading buffer, and resolved on SDS-PAGE. Membranes were blocked in 5% milk in TBST (50 mM Tris, 150 mM NaCl, and 0.05% Tween 20) buffer and then probed with antibodies as indicated.

Immunofluorescence Staining—Indirect immunofluorescence was carried out as described previously (28, 29). HEK293T or HeLa cells cultured on coverslips were treated with CPT (1 μM) for 1 h or IR (10 gray) for 1 h, respectively. Cells were then washed with PBS, pre-extracted with buffer containing 0.5% Triton X-100 for 5 min, and fixed with 3% paraformaldehyde for 10 min at room temperature. Cells were incubated in primary antibody for 20 min at room temperature. Following three 5-min washes with PBS, secondary antibody was added at room temperature for 20 min. Cells were then stained with DAPI to visualize nuclear DNA. The coverslips were mounted onto glass slides with antifade solution and visualized using a Nikon Eclipse i80 fluorescence microscope with a Nikon Plan Fluor ×60 oil objective lens.
Protein Purification—Full-length RPA1 was cloned into MBP-His-tagged vector for the expression of MBP-RPA1-His protein in insect cells. Transposition occurred in DH10Bac-competent cells, and correct bacmids confirmed by PCR were transfected into Sf9 cells for baculovirus production. After viral amplification, Sf9 cells were infected with baculovirus stocks expressing MBP-RPA1-His for 48 h. Cells were harvested and washed with 1× PBS and resuspended in lysis buffer (20 mM Tris-HCl; 300 mM NaCl; 1% Triton X-100; and 1 μg/ml each of leupeptin, aprotinin, and pepstatin). The extract was centrifuged at 18,000 rpm for 40 min. The supernatant was collected and loaded onto a pre-equilibrated nickel-nitrilotriacetic acid-agarose and washed with lysis buffer plus 20 mM imidazole and protease inhibitor. The bound protein was then eluted with lysis buffer containing 200 mM imidazole and protease inhibitor. Peak fractions were pooled and incubated with amylose resin for 2 h at 4 °C. After washing the beads with 100 ml of washing buffer (20 mM Tris-HCl; 500 mM NaCl; 0.5% Nonidet P-40; 1 mM DTT; and 1 μg/ml each of leupeptin, aprotinin, and pepstatin), the bound protein was used for a pulldown assay. Full-length PSO4 and BCAS2 were cloned into the PGEX-6P-1 vector for the expression of GST-tagged fusion proteins in E. coli. Cells were grown at 37 °C until log phase and induced with 0.2 mM isopropyl 1-thio-β-D-galactopyranoside at 17 °C for 16 h. Cells were then harvested and resuspended in lysis buffer (20 mM Tris-HCl; 300 mM NaCl; 1% Triton X-100; 1 mM DTT; and 1 μg/ml each of leupeptin, aprotinin, and pepstatin). After sonication, the extract was centrifuged at 18,000 rpm for 40 min. The supernatant was collected and incubated with glutathione-Sepharose resin for 4 h at 4 °C. After washing the beads with washing buffer (20 mM Tris-HCl; 500 mM NaCl; 0.5% Nonidet P-40; 1 mM DTT; and 1 μg/ml each of leupeptin, aprotinin, and pepstatin), the bound proteins were eluted with washing buffer containing 20 mM glutathione and used for pull-down assays.

Lentivirus Packaging and Infection—The Tet-On-inducible, SFB-tagged lentiviral vector and packaging plasmids (pMD2G and pSPAX2) were provided by Prof. Songyang Zhou (Baylor College of Medicine). BCAS2 and PSO4 entry constructs were transferred into the Gateway-compatible, SFB-tagged lentiviral vector. Virus supernatant was collected 48 h after the cotransfection of lentiviral vectors and packaging plasmids (pMD2G and pSPAX2) into HEK293T cells. Cells were infected with viral supernatants with the addition of 8 μg/ml Polybrene (Sigma), and stable pools were selected with medium containing 500 μg/ml G418 (Calbiochem). The expression of the indicated genes in the stable pools was induced by the addition of 1 μg/ml doxycycline (Sigma) for 48 h for the experiments presented in this work.

RESULTS AND DISCUSSION

Both BCAS2 and PSO4 Interact and Colocalize with RPA—RPA plays a central role in DNA replication, repair, and recombination (1, 2). In an attempt to search for previously undetected proteins present in the RPA-containing complex, we performed tandem affinity purification using HEK293T cells stably expressing triple epitope (S-peptide, FLAG, and streptavidin-binding peptide)-tagged wild-type RPA1 or RPA3 for the identification of RPA-interacting proteins. Mass spectrometry analysis revealed a number of known RPA-associated proteins, including hPrImpol1/PrImpol/CCDC111, BLM, and RAD52 (supplemental Tables 1 and 2) (30–35). Interestingly, we also identified that all four subunits of PSO4 exist in a complex with RPA (supplemental Tables 1 and 2).

To determine whether the PSO4 complex indeed interacts with RPA, we performed transient transfection and coimmunoprecipitation experiments. The results demonstrated that both the PSO4 and BCAS2 subunits could interact with RPA1 (Fig. 1A). In the same experiments, a weak interaction between PSO4 and RPA2 and RPA3 was also found to take place (Fig. 1A). These results indicate that the PSO4 complex associates with the RPA complex mainly through RPA1.

We next tested the possibility that there could be a direct protein-protein interaction between RPA1 and PSO4 and BCAS2. Pulldown assays using recombinant MBP-tagged RPA1 and GST-tagged PSO4 or BCAS2 purified from E. coli demonstrated that RPA1 interacts strongly with BCAS2 and weakly with PSO4 in vitro (Fig. 1B).

To examine the interaction between endogenous RPA and the PSO4 complex, HeLa cell extracts were prepared and subjected to immunoprecipitation assays in the presence of either control IgG or anti-RPA2 antibody (Fig. 1C). Western blot analysis revealed that both PSO4 and BCAS2 were clearly detected in the immunoprecipitations obtained with the anti-RPA2 antibody but not with the control IgG (Fig. 1C). We also performed a reciprocal coimmunoprecipitation assay. As shown in Fig. 1D, the endogenous RPA1 and RPA2 were readily immunoprecipitated with the BCAS2-specific antibody but not with the control IgG. Moreover, treatment with the topoisomerase I inhibitor CPT slightly enhanced the interaction between BCAS2/PSO4 and RPA (Fig. 1, C and D). In these experiments, benzonase was included in the lysis buffer to exclude the possibility that the interaction occurs indirectly via DNA bridging (Fig. 1, C and D).

Upon occurrence of DNA damage, RPA and several other proteins involved in the DNA damage response could form large nuclear foci. A physical interaction between RPA and BCAS2 and PSO4, as demonstrated above, raises the possibility that BCAS2 and PSO4 may colocalize with RPA at DNA damage sites. Indeed, discrete foci of BCAS2 and PSO4 were readily detected in cells following CPT or IR treatment (Fig. 1, E and F). Moreover, these foci colocalize with RPA2 foci (more than 90%), indicating that the localization of BCAS2 and PSO4, like that of RPA, is regulated in response to DNA damage (Fig. 1, E and F).

The N Terminus of BCAS2 Binds to the C Terminus of RPA1—BCAS2 is a small subunit of the PSO4 complex that contains two coiled coil motifs (10, 11). Because BCAS2 directly interacts with RPA1, we next mapped the RPA1-binding domain on BCAS2. A series of deletion mutants that span the entire coding region of BCAS2 were generated (Fig. 2A). Coimmunoprecipitation experiments revealed that BCAS2 associated with RPA1 via its N terminus because the deletion mutant lacking the N-terminal 60 amino acids (D1) failed to coprecipitate with RPA1 (Fig. 2B). Interestingly, the D1 mutant, which does not bind to RPA1, failed to relocalize to DNA damage sites, suggesting that the binding of BCAS2 to RPA1 is important for its proper localization in response to DNA damage (Fig. 2C). In
support of this conclusion, RPA1 depletion abolished BCAS2 focus formation after DNA damage (Fig. 2, D–F).

To understand how RPA1 interacts with BCAS2 and facilitates its recruitment to sites of DNA damage, we used a panel of RPA1 constructs (Fig. 2G). The results indicated that RPA1 associated with BCAS2 via its DBD-C domain because the mutant that lacks the DBD-C domain (D5) failed to interact with BCAS2 (Fig. 2H).

**FIGURE 1.** The PSO4 complex associates and colocalizes with RPA. A, RPA1 interacts with PSO4 and BCAS2. HEK293T cells were transiently transfected with plasmids encoding SFB-tagged RPA1, RPA2, or RPA3 together with plasmids encoding Myc-tagged PSO4, PLRG1, CDC5L, or BCAS2. Cell lysates were immunoprecipitated (IP) with anti-FLAG antibody, and Western blot analysis was performed with anti-FLAG and anti-Myc antibodies. B, direct binding between recombinant MBP-tagged-RPA1 and GST-tagged-BCAS2 or GST-tagged-PSO4. Top panel, BCAS2 and PSO4 were detected by immunoblotting. Bottom panel, purified proteins visualized by Coomassie staining. C and D, association of endogenous PSO4 complex with RPA in HeLa cells was performed by coimmunoprecipitation using anti-RPA2 (C) or anti-BCAS2 (D) antibody. HeLa cells treated with CPT (1 μM) for 1 h were lysed in the presence of benzonase. Cell lysates were then incubated with protein A-agarose beads conjugated with the indicated antibodies, and Western blot analysis was carried out as indicated. E, BCAS2 colocalizes with RPA2. 293T cells were either mock-treated or treated with CPT (1 μM) or IR (10 gray) for 1 h before fixing and processing for BCAS2 and RPA2 immunofluorescence. A merged image shows colocalization. F, PSO4 colocalizes with RPA2. SFB-tagged PSO4 was expressed in HEK293T cells. Foci assembled by this fusion protein and by RPA2 following exposure to CPT (1 μM) or IR (10 gray) for 1 h were detected by immunofluorescence using anti-FLAG and anti-RPA2 antibodies, respectively. SFB-PSO4 foci were detected in green, whereas RPA2 foci were detected in red. A merged image shows colocalization.
To further test whether the N-terminal region of BCAS2 is sufficient to bind RPA1, we generated a construct encoding the N-terminal region alone (N1–60, residues 1–60). As shown in Fig. 2, the N-terminal region alone failed to interact with RPA1. In contrast, the RPA1 DBD-C domain (residues 423–616) is not only essential but also sufficient for its interaction with BCAS2 (Fig. 2).

**Figure 2.** BCAS2 localizes to DNA damage sites via an interaction with RPA1. A, schematic representation of wild-type and deletion mutants of BCAS2 used in this study. Their ability to bind to RPA1 is indicated. CC, coiled coil domain. B, the 60 amino acids at the N terminus of BCAS2 are required for RPA1 binding. HEK293T cells were transfected with plasmids encoding Myc-tagged RPA1 together with plasmids encoding SFB-tagged wild-type BCAS2 or the mutants. Cell lysates were immunoprecipitated (IP) with anti-FLAG antibody, and Western blot analysis was performed with the indicated antibodies. C, the N terminus of BCAS2 is required for its focus formation. HEK cells were infected with lentivirus expressing SFB-tagged wild-type BCAS2 or the D1 mutant. 48 h later, cells were treated with CPT (1 μM) for 1 h before fixing and processing for immunofluorescence. D–F, RPA1 deletion impairs BCAS2 focus formation. HEK293T cells infected with non-target or RPA1-specific lentiviral shRNAs for 48 h were treated with CPT (1 μM) for 1 h before fixing and processing for BCAS2 immunofluorescence. Representative BCAS2 foci are shown (D). Quantification results were the average of three independent experiments and are presented as mean ± S.E. sh-Con, control shRNA. E, more than 100 cells were counted in each experiment. The knockdown efficiency of RPA1 using specific shRNAs was confirmed by immunoblotting of lysates prepared from HEK293T cells expressing the indicated shRNA (F). G, schematic of full-length RPA1 and the mutants used in this study. The ability to bind to BCAS2 is indicated. H, RPA1 with the DBD-C domain deletion could not bind to BCAS2. HEK293T cells were transfected with plasmids encoding Myc-tagged wild-type or mutant RPA1 together with a plasmid encoding SFB-tagged BCAS2. Cell lysates were immunoprecipitated with anti-FLAG antibody, and Western blot analysis was performed with the indicated antibodies. I, the N-terminal region of BCAS2 (N1–60, residues 1–60) is not sufficient for its interaction with RPA1. Coimmunoprecipitation experiments were carried out as indicated. J, the RPA1 DBD-C domain (residues 423–616) is not only essential but also sufficient for its interaction with BCAS2. Coimmunoprecipitation experiments were carried out as indicated.
whether BCAS2 and PSO4 also play important roles in the DNA damage response. As shown in Fig. 3, A and B, BCAS2 or PSO4 depletion led to impaired CHK1 and RPA2 phosphorylation after DNA damage. HeLa cells infected with the indicated shRNAs were either mock-treated or treated with CPT (1 μM) or IR (10 gray) for 1 h. Cell lysates were immunoblotted with the indicated antibodies. The asterisk indicates a nonspecific band. sh-Con, control shRNA. C and D, BCAS2 or PSO4 depletion impairs CPT-induced phospho-RPA2 focus formation. BCAS2- or PSO4-depleted HeLa cells were treated with CPT (1 μM) for 1 h before fixing and processing for phospho-RPA2 immunofluorescence. Representative phospho-RPA2 foci are shown (C). Quantification results were the average of three independent experiments and are presented as mean ± S.E. (D). More than one hundred cells were counted in each experiment. E and F, BCAS2 or PSO4 depletion impairs CPT-induced ATRIP focus formation. BCAS2- or PSO4-depleted HeLa cells were treated with CPT (1 μM) for 1 h before fixing and processing for ATRIP immunofluorescence. Representative ATRIP foci are shown (E). Quantification results were the average of three independent experiments and are presented as mean ± S.E. (F). More than 100 cells were counted in each experiment.

The Ability of BCAS2 to Function in the DNA Damage Response Correlates with Its Association with RPA1—To further decipher the biological significance of the BCAS2-RPA1 interaction, we performed rescue experiments to test whether the RPA1-binding region on BCAS2 is required for efficient accumulation of ATRIP at DNA damage sites and the subsequent CHK1 activation and RPA2 phosphorylation. We generated HeLa cell lines to express wild-type BCAS2 or its deletion mutant defective in RPA1 binding under the control of a tetracycline-inducible promoter. By treating these cell lines with shRNA targeting the 3′/H11032 UTR of the BCAS2 transcript (shRNA 2), we can specifically knock down the endogenous, but not exogenous, BCAS2. The expression of wild-type and mutated BCAS2 was induced in BCAS2 knockdown cells when the cells were treated with doxycycline (Fig. 4, A and B). Interestingly, whereas wild-type BCAS2 successfully restored CHK1 activation and RPA2 phosphorylation to levels comparable with that of control cells, BCAS2 deletion mutant defective in RPA1 binding failed to do so (Fig. 4B). Consistently, the defects in ATRIP focus formation after DNA damage could be reversed by the expression of wild-type BCAS2 but not the D1 mutant (Fig. 4, C and D).

The E3 Ligase Activity of PSO4 Is Indispensable for the Recruitment of ATRIP to DNA Damage Sites and the Subsequent CHK1 and RPA2 Phosphorylation—PSO4 is a U-box-containing E3 ubiquitin ligase (20–22). To explore the physio-
logical relevance of this E3 ligase activity of PSO4 in ATRIP recruitment, CHK1 activation, and RPA2 phosphorylation, we took advantage of the inducible expression system to express wild-type BCAS2 (WT) or its deletion mutant defective in RPA1 binding (D1) under the control of a tetracycline-inducible promoter were generated. The resulting cell lines were then infected with shRNA targeting the 3' UTR of BCAS2 transcript (shRNA#2). 48 h after infection, cells were induced by doxycycline addition for 24 h prior to CPT (1 μM) treatment. 1 h later, cells were collected, and lysates were immunoblotted with the indicated antibodies (A and B) or subjected to immunostaining using ATRIP antibody (C and D). Representative ATRIP foci are shown (C). Quantification results were the average of three independent experiments and are presented as mean ± S.E. (D). More than 100 cells were counted in each experiment. The asterisk indicates a nonspecific band. sh-Con, control shRNA. E–H, the U-box deletion mutant of PSO4 could not restore PSO4 function in vivo. HeLa cell lines to express shRNA 2-resistant wild-type PSO4 (WT) or its U-box deletion mutant (ΔU) under the control of a tetracycline-inducible promoter were generated. The resulting cell lines were then infected with PSO4 shRNA 2. 48 h after infection, cells were induced by doxycycline addition for 24 h prior to CPT (1 μM) treatment. 1 h later, cells were collected, and lysates were immunoblotted with the indicated antibodies (E and F) or subjected to immunostaining using ATRIP antibody (G and H). Representative ATRIP foci are shown (G). Quantification results were the average of three independent experiments and are presented as mean ± S.E. (H). More than 100 cells were counted in each experiment. The asterisk indicates a nonspecific band.

FIGURE 4. Both the RPA1-binding ability of BCAS2 and the E3 ligase activity of PSO4 are required for the recruitment of ATRIP to DNA damage sites and the subsequent CHK1 and RPA2 phosphorylation. A–D, the mutant defective in RPA1 binding could not restore BCAS2 function in vivo. HeLa cell lines to express wild-type BCAS2 (WT) or its deletion mutant defective in RPA1 binding (D1) under the control of a tetracycline-inducible promoter were generated. The resulting cell lines were then infected with shRNA targeting the 3' UTR of BCAS2 transcript (shRNA#2). 48 h after infection, cells were induced by doxycycline addition for 24 h prior to CPT (1 μM) treatment. 1 h later, cells were collected, and lysates were immunoblotted with the indicated antibodies (A and B) or subjected to immunostaining using ATRIP antibody (C and D). Representative ATRIP foci are shown (C). Quantification results were the average of three independent experiments and are presented as mean ± S.E. (D). More than 100 cells were counted in each experiment. The asterisk indicates a nonspecific band. sh-Con, control shRNA. E–H, the U-box deletion mutant of PSO4 could not restore PSO4 function in vivo. HeLa cell lines to express shRNA 2-resistant wild-type PSO4 (WT) or its U-box deletion mutant (ΔU) under the control of a tetracycline-inducible promoter were generated. The resulting cell lines were then infected with PSO4 shRNA 2. 48 h after infection, cells were induced by doxycycline addition for 24 h prior to CPT (1 μM) treatment. 1 h later, cells were collected, and lysates were immunoblotted with the indicated antibodies (E and F) or subjected to immunostaining using ATRIP antibody (G and H). Representative ATRIP foci are shown (G). Quantification results were the average of three independent experiments and are presented as mean ± S.E. (H). More than 100 cells were counted in each experiment. The asterisk indicates a nonspecific band.

Note that PSO4 may act as an E3 ubiquitin ligase and regulate certain unknown substrates at DNA damage sites that are critical for efficient recruitment of ATRIP to sites of DNA damage and the subsequent CHK1 activation and RPA2 phosphorylation. Interestingly, while our manuscript was in preparation, an independent study reported that PSO4 promotes RPA ubiquitination and facilitates the accumulation of ATRIP at sites of DNA damage (36).
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In summary, we discovered a functional interaction between the PSO4 complex and RPA. We propose that the PSO4 complex, through its interaction with RPA, is recruited to sites of DNA damage and exerts its E3 ligase activity to ubiquitinate RPA and, thus, to modulate the cellular response to DNA damage.

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