





SLFN11 inhibits checkpoint maintenance and homologous recombination repair

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Abstract

High expression levels of SLFN11 correlate with the sensitivity of human cancer cells to DNA-damaging agents. However, little is known about the underlying mechanism. Here, we show that SLFN11 interacts directly with RPA1 and is recruited to sites of DNA damage in an RPA1-dependent manner. Furthermore, we establish that SLFN11 inhibits checkpoint maintenance and homologous recombination repair by promoting the destabilization of the RPA-ssDNA complex, thereby sensitizing cancer cell lines expressing high endogenous levels of SLFN11 to DNA-damaging agents. Finally, we demonstrate that the RPA1-binding ability of SLFN11 is required for its function in the DNA damage response. Our findings not only provide novel insight into the molecular mechanisms underlying the drug sensitivity of cancer cell lines expressing SLFN11 at high levels, but also suggest that SLFN11 expression can serve as a biomarker to predict responses to DNAdamaging therapeutic agents.

Keywords checkpoint initiation; checkpoint maintenance; DNA damage response; homologous recombination repair; RPA

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Introduction

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The integrity of genomic DNA is constantly challenged by a variety of endogenous and exogenous DNA-damaging agents, which can cause many different types of DNA lesions [1–5]. These lesions can block DNA replication and transcription and, if not repaired or repaired incorrectly, produce mutations or large-scale genome aberrations that might threaten the survival of the individual cells and the whole organism [1–5]. To counteract the potentially deleterious effects of DNA damage and preserve genome stability, cells have

evolved the ability to sense DNA damage, activate the DNA damage checkpoint, and initiate DNA repair [1–5]. Defects in any of these functions promote tumor development but also make cancer cells more vulnerable to DNA-damaging agents [1–5]. Thus, increasing the knowledge of DNA damage response is not only important to deepen our basic understanding of how cancer develops, but also to develop drugs exhibiting selective toxicity toward cancer cells.

Schlafen (Slfn) (from the word schlafen, which in German means sleeping) genes originally identified during screening for growth regulatory genes that are differentially expressed during lymphocyte development encode a family of proteins limited to mammalian organisms [6-10]. To date, 10 mouse (Slfn1, 1L, 2, 3, 4, 5, 8, 9, 10, and 14) and 6 human (Slfn5, 11, 12, 12L, 13, and 14) Slfn genes have been identified [6–10]. There is emerging evidence that several SLFN family proteins play critical roles in development, immune response, and cell proliferation [6–10]. Human Slfn11 gene encodes a member of a protein family with structural similarity to RNA helicases [6,7,11-13]. A previous study has shown that SLFN11 binds transfer RNA and can specifically abrogate the production of retroviruses such as human immunodeficiency virus 1 (HIV-1) by selectively blocking the expression of viral proteins in a codonusage-dependent manner [12]. Besides its important antiviral properties, SLFN11 is able to sensitize cancer cells to DNA-damaging agents [11,14,15]. However, mechanistically how this is achieved remains elusive and largely speculative.

Replication protein A (RPA) is a heterotrimeric protein complex composed of three subunits known as RPA1, RPA2, and RPA3 [16,17]. RPA is the main eukaryotic single-stranded DNA (ssDNA) binding protein that is essential for a variety of DNA metabolic pathways including DNA replication, recombination, DNA damage checkpoint, as well as DNA repair [16,17]. The ability of RPA to specifically bind ssDNA is dependent on its four OB (oligonucleotide/oligosaccharide binding) folds commonly referred to as DNA-binding domains DBD-A, DBD-B, DBD-C, and DBD-D [18,19]. The DBD-A, DBD-B, and DBD-C domains are all located in the RPA1 subunit, whereas DBD-D domain residues in the RPA2 subunit [18,19]. A growing body of evidence demonstrates that RPA-bound

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In this study, we report the identification of RPA as a binding partner of SLFN11 by tandem affinity purification and mass spectrometry. We show that SLFN11 is recruited to sites of DNA damage in an RPA-dependent manner. We further demonstrate that SLFN11 is able to promote the destabilization of RPA-ssDNA complex. As a result, cells expressing high levels of SLFN11 display defects in checkpoint maintenance and homologous recombination repair and thus are hypersensitive to DNA-damaging agents. Collectively, our results provide important mechanistic insights into how SLFN11 sensitizes cancer cells to DNA-damaging agents and will shed new light on personalized cancer therapy.

Results

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SLFN11 localizes to sites of DNA damage

Although SLFN11 is capable of sensitizing cancer cells to DNAdamaging agents and has been speculated to play a role in the DNA damage response, exactly how SLFN11 participates in this process remains unclear. To gain insight into the cellular function of SLFN11, we first generated polyclonal anti-SLFN11 antibody and analyzed its expression at the protein level in several human cell lines. As shown in Fig 1A, SLFN11 was only detected in DU145 and SF268 cells, but not in HEK293T, U2OS, HeLa, and HCT116 cells. We next sought to determine whether SLFN11 can be recruited to sites of DNA damage. As shown in Fig 1B, we found that endogenous SLFN11 was recruited to DNA damage sites following laser micro-irradiation and co-localized with singlestranded DNA (ssDNA)-binding protein RPA in both SF268 and DU145 cell lines expressing high endogenous levels of SLFN11, but not in HeLa and U2OS cell lines expressing very low or undetectable levels of SLFN11. Similarly, discrete foci of Flag-tagged SLFN11, which co-localized with RPA, were readily detected in both SF268 and DU145 cell lines following topoisomerase I inhibitor camptothecin (CPT) or IR treatment (Fig 1C and D). Taken together, these results suggest that SLFN11 is a DNA damageresponsive protein and may have an important role in the regulation of DNA damage response.

SLFN11 interacts with RPA

In order to understand how SLFN11 might participate in the DNA damage response, we established a stably transfected HEK293 derivative cell line that expresses N-terminally triple-tagged (S-protein, FLAG, and streptavidin-binding peptide) full-length SLFN11 for the identification of potential SLFN11-binding partners. After a tandem affinity purification (TAP) scheme, proteins associated with SLFN11 were identified by mass spectrometry analysis. As shown in Fig 2A, we reproducibly found that RPA, the primary ssDNA-binding protein in eukaryotes composed of three subunits known as RPA1, RPA2, and RPA3, exists in a complex with SLFN11.

To confirm the interaction between SLFN11 and RPA, we performed transient transfection and co-immunoprecipitation experiments. As shown in Fig 2B, SFB-tagged SLFN11 interacted with

Myc-tagged RPA1, RPA2, and RPA3 but not with the Morc3 control protein. To further examine the interaction between endogenous SLFN11 and the RPA complex, SF268 whole-cell extracts were prepared and subjected to immunoprecipitation assays in the presence of either control IgG or anti-SLFN11 antibody. Western blot analysis revealed that both RPA1 and RPA2 were clearly detected in the immunoprecipitates with the anti-SLFN11 antibody but not with the control IgG (Fig 2C, left panel). We also performed a reciprocal co-immunoprecipitation assay. As shown in Fig 2C (right panel), the endogenous SLFN11 was readily immunoprecipitated with the RPA2-specific antibody, but not with the control IgG. Interestingly, the interaction between SLFN11 and RPA was slightly enhanced following CPT treatment (Fig 2C). We also observed co-immunoprecipitation of SLFN11 and RPA in cell extracts from human DU145 cells (Fig 2D).

In order to determine whether the interaction between SLFN11 and RPA is direct, we expressed and purified recombinant MBP-tagged RPA1, RPA2, RPA3, and GST-tagged SLFN11 (the recombinant SLFN11 protein is very unstable, probably due to proteolysis) (Fig 2E, lower panel). *In vitro* pull-down assays showed that SLFN11 binds with RPA1 but not with RPA2 or RPA3 (Fig 2E, upper panel), suggesting that SLFN11 interacts with the RPA complex through RPA1.

To identify the region of SLFN11 that is required for its interaction with RPA1, we made a series of SLFN11 deletion mutants and performed co-immunoprecipitation experiments (Fig 2F). As shown in Fig 2G, SLFN11 interacts with RPA1 via its very C-terminus, since deletion mutant lacking the C-terminal 161 amino acid (Δ 5) was unable to co-precipitate with RPA1.

RPA is required for SLFN11 localization to sites of DNA damage

Since RPA has a DNA-binding function and exists in a complex with SLFN11, we sought to test whether RPA is important for the assembly of DNA damage-induced SLFN11 nuclear foci. We depleted RPA1 expression using lentiviral shRNAs and confirmed RPA1 depletion by Western blotting analysis. Consistent with the observation that RPA1, among the three subunits of RPA, contains the major ssDNA-binding activity, RPA1 depletion dramatically reduced the accumulation of RPA2 at sites of CPT- or laser-induced DNA damage (Fig 3A-D). Remarkably, SLFN11 accumulation at sites of DNA damage was also severely impaired in RPA1-depleted SF268 cells, but unaffected in mock treated (Fig 3A–D). In addition, the $\Delta 5$ mutant, which does not bind to RPA1, failed to re-localize to sites of DNA damage (Fig 3E). These results, taken together with the observation that RPA1 depletion did not result in any detectable change in the protein level of SLFN11 (Fig 3C), suggest that RPA plays an important role in the recruitment of SLFN11 to sites of DNA damage.

In response to DNA damage induced by CPT, RPA-coated ssDNA is generated by the MRN (Mre11-RAD50-NBS1) complex and CtIP-dependent DNA-end resection [20,21]. The observation that RPA is required for efficient recruitment of SLFN11 to sites of DNA damage led us to propose that the MRN complex and CtIP may also be involved in SLFN11 recruitment. As expected, Mre11 or CtIP depletion significantly decreased CPT-induced recruitment of RPA2 and the downstream SLFN11 to DNA damage sites (Fig 3F–H).

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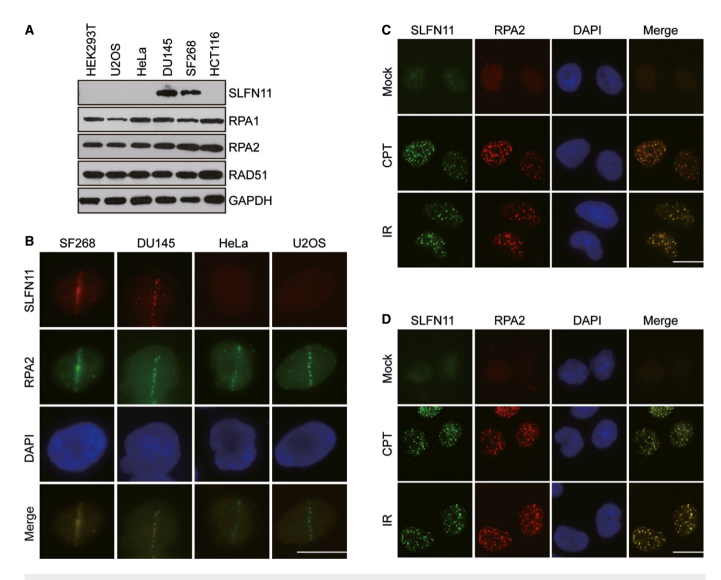


Figure 1. SLFN11 is a DNA damage-responsive protein.

- A Expression analysis of SLFN11 in human cell lines.
- B Recruitment of endogenous SLFN11 to laser-induced DNA damage sites. Forty minutes after laser irradiation, cells were stained with antibodies against SLFN11 and RPA2. Scale bar, 10 μm.
- C, D SLFN11 co-localizes with RPA2 in DNA damage-induced nuclear foci. Flag-tagged SLFN11 was expressed in SF268 (C) or DU145 (D) cells. Foci assembled by this fusion protein and by RPA2 following exposure to CPT (1 µM) for 3 h or IR (10 Gy) for 3 h were detected by immunofluorescence using anti-Flag and anti-RPA2 antibodies, respectively. Flag-SLFN11 foci were detected in green, while RPA2 foci were detected in red. A merged image shows co-localization. Scale bar, 10 µm.

Source data are available online for this figure.

SLFN11 inhibits maintenance but not initiation of DNA damage checkpoint

Replication protein A plays essential roles in DNA replication, repair, and DNA damage checkpoint. The localization of SLFN11 to RPA-coated ssDNA and its ability to sensitize cancer cells to DNA-damaging agents indicate that SLFN11 may function together with RPA in the DNA damage response. To test this possibility, we first examined the kinetics of CHK1 and CHK2 phosphorylation in cell lines expressing SLFN11 at high (SF268 and DU145) and very low or undetectable levels (HeLa and U2OS), respectively (Fig 4A). We treated these cells with CPT for 1 h and subsequently released the

cells into fresh medium lacking CPT and then monitored the CHK1 and CHK2 phosphorylation over time. As shown in Fig 4A, after exposure to CPT for 1 h, both CHK1 and CHK2 were similarly activated in all four cell lines, indicating that SLFN11 is not involved in the initial activation of the checkpoint. In contrast, both cell lines expressing high endogenous levels of SLFN11 (SF268 and DU145) showed a markedly earlier decline of CHK1 and CHK2 phosphorylation when compared to cell lines expressing very low or undetectable levels of SLFN11 (HeLa and U2OS) (Fig 4A). These results suggest that cells expressing high endogenous SLFN11 display a specific defect in the maintenance of CHK1 and CHK2 activation after DNA damage.

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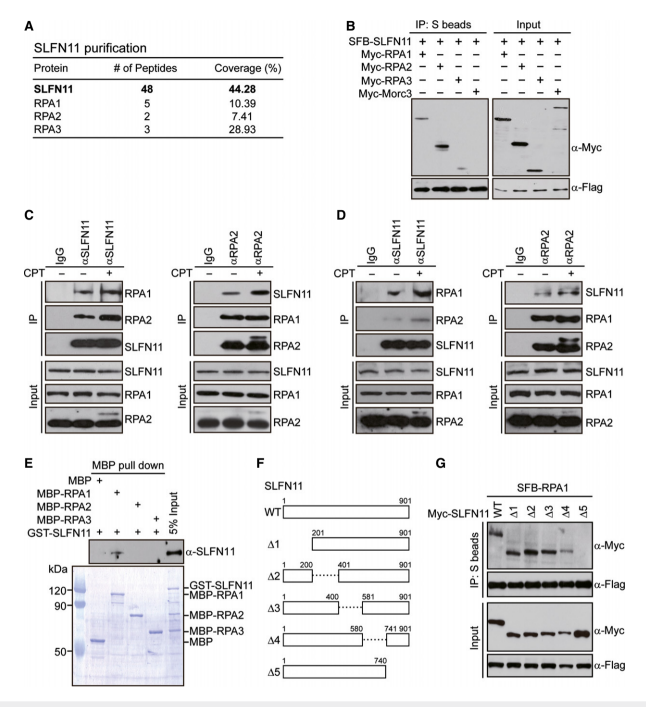


Figure 2. SLFN11 forms a complex with RPA.

- A HEK293 cells stably expressing SFB-tagged (S-tag, Flag epitope tag, and streptavidin-binding peptide tag) SLFN11 were used for TAP of protein complexes. Tables are summaries of proteins identified by mass spectrometry analysis. Letters in bold indicate the bait proteins.
- B SLFN11 interacts with RPA, but not with the Morc3 control protein. HEK293 cells were transiently transfected with plasmids encoding SFB-tagged SLFN11 together with plasmids encoding Myc-tagged RPA or Morc3. Cell lysates were immunoprecipitated with S beads, and Western blot analysis was performed with anti-Flag and anti-Myc antibodies.
- C, D Association of endogenous SLFN11 with RPA in SF268 (C) or DU145 (D) cells was performed by co-immunoprecipitation using anti-SLFN11 or anti-RPA2 antibody. Nuclease-treated cell lysates were incubated with protein A agarose beads conjugated with indicated antibodies, and Western blot analysis was performed according to standard procedures.
- E Direct binding between recombinant GST-tagged SLFN11 and MBP-tagged RPA1. Upper panel: SLFN11 was detected by immunoblotting. Lower panel: Purified proteins visualized by Coomassie staining.
- F Schematic representation of wild-type and deletion mutants of SLFN11 used in this study.
- G HEK293 cells were transfected with indicated plasmids. Cell lysates were immunoprecipitated with S beads, and Western blot analysis was performed with anti-Flag and anti-Myc antibodies.

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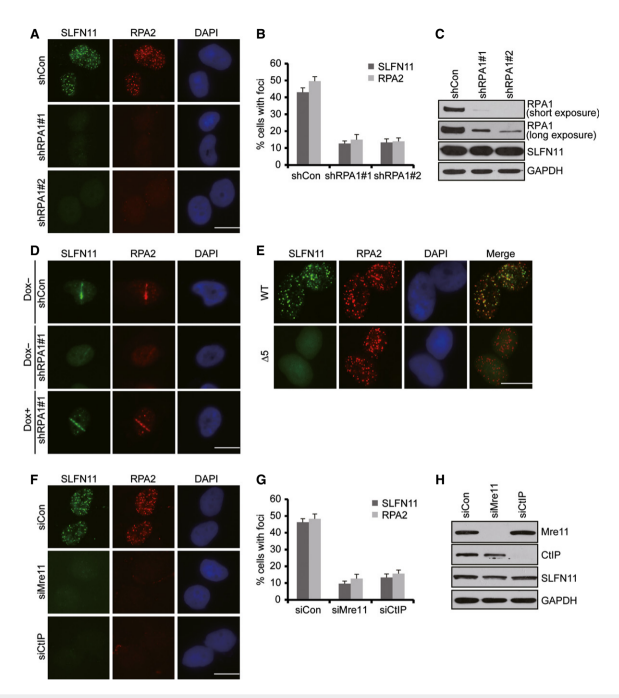


Figure 3. RPA recruits SLFN11 to sites of DNA damage.

- A–C SF268 cells stably expressing Flag-tagged SLFN11 were infected with non-target or RPA1-specific lentiviral shRNAs. Forty-eight hours after infection, cells were treated with CPT (1 μM). Three hours later, cells were subjected to immunostaining using anti-RPA2 and anti-Flag antibodies. Representative SLFN11 and RPA2 foci were shown (A). Scale bar, 10 μm. Quantification of RPA2 and SLFN11 foci formation using NIH ImageJ software (B). Error bars represent SD; n = 3. Knockdown efficiency of RPA1 in SF268 cells was confirmed by immunoblotting (C).
- D RPA1 is required for the accumulation of SLFN11 at laser-induced DNA damage tracks. A SF268 cell line stably expressing wild-type RPA1 under the control of a tetracycline-inducible promoter was generated. The resulting cell line infected with RPA1-specific lentiviral shRNA targeting the 3′ UTR of RPA1 transcript was induced by doxycycline addition for 48 h and then laser micro-irradiated. Thirty minutes later, cells were fixed and stained with anti-RPA2 and anti-SLFN11 antibodies. Scale bar, 10 μm.
- E The C-terminus of SLFN11 is required for its foci formation. SF268 cells were transfected with plasmids encoding Flag-tagged wild-type or mutant SLFN11. Immunostaining experiments were performed 3 h after CPT (1 μM) treatment using the indicated antibodies. Scale bar, 10 μm.
- F–H SF268 cells stably expressing Flag-tagged SLFN11 were transfected twice with the indicated siRNAs. Forty-eight hours after second transfection, cells were treated with CPT (1 μM). Three hours later, cells were subjected to immunostaining using anti-RPA2 and anti-Flag antibodies. Representative SLFN11 and RPA2 foci were shown (F). Scale bar, 10 μm. Quantification of RPA2 and SLFN11 foci formation using NIH ImageJ software (G). Error bars represent SD; n = 3. Knockdown efficiency of Mre11 and CtIP in SF268 cells was confirmed by immunoblotting (H).

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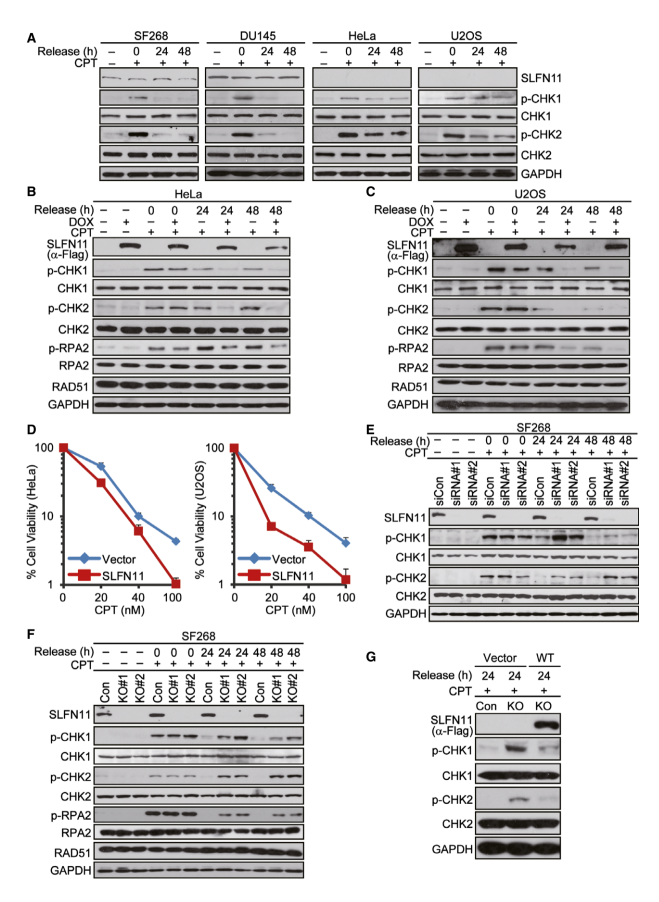


Figure 4.

Figure 4. SLFN11 inhibits maintenance but not initiation of DNA damage checkpoint.

- A Analysis of the kinetics of CHK1 and CHK2 phosphorylation in cell lines expressing SLFN11 at high (SF268 and DU145) and very low or undetectable levels (HeLa and U2OS), respectively. The indicated cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and harvested at the indicated time points for immunoblotting with the indicated antibodies.
- B, C HeLa (B) or U2OS (C) cells overexpressing exogenous SLFN11 display a specific defect in checkpoint maintenance. A HeLa or U2OS cell line to express Flag-tagged SLFN11 under the control of a tetracycline-inducible promoter was generated. The resulting cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and harvested at the indicated time points for immunoblotting with the indicated antibodies.
- D Overexpression of exogenous SLFN11 rendered both HeLa and U2OS cells more sensitive to CPT treatment. A HeLa or U2OS cell line stably expressing Flag-tagged SLFN11 was generated. The resulting cells were treated with various doses of CPT for 24 h, then shifted to fresh medium and permitted to grow for 14 days before staining. Error bars represent SD; *n* = 3.
- E Downregulation of SLFN11 delays the decline of phosphorylated CHK1 and CHK2 in SF268 cells. SF268 cells were transfected twice with control siRNA or siRNAs specific for SLFN11. Forty-eight hours after transfection, cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and harvested at the indicated time points for immunoblotting with the indicated antibodies.
- F SLFN11 knockout SF268 cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and harvested at the indicated time points for immunoblotting with the indicated antibodies.
- G A SLFN11-deficient SF268 cell line stably expressing Flag-tagged SLFN11 was generated. The resulting cell line was treated with CPT (1 μM) for 1 h. Cells were then washed and shifted to fresh medium. Twenty-four hours later, cells were harvested for immunoblotting with the indicated antibodies.

Source data are available online for this figure.

To further explore the role of SLFN11 in the regulation of checkpoint maintenance, we examined the effects of exogenous overexpression of SLFN11 in cells expressing SLFN11 at very low or undetectable levels. We engineered a HeLa cell line to express Flagtagged SLFN11 under the control of a tetracycline-inducible promoter and examined the kinetics of CHK1 and CHK2 phosphorylation after CPT treatment. As shown in Fig 4B, the expression of wild-type SLFN11 was induced in HeLa cells when the cells were treated with doxycycline. Interestingly, HeLa cells overexpressing exogenous SLFN11 displayed a specific defect in the maintenance of CHK1 and CHK2 phosphorylation, but not the initial CHK1 and CHK2 phosphorylation. As RPA2 is hyper-phosphorylated upon DNA damage or replication stress [22], we further determined whether SLFN11 is also able to affect the phosphorylation of RPA. As shown in Fig 4B, overexpression of exogenous SLFN11 in HeLa cells inhibited the maintenance but not initiation of RPA2 phosphorylation on Ser4/Ser8. Similar results were observed in U2OS cells (Fig 4C). Consistently, overexpression of exogenous SLFN11 rendered both HeLa and U2OS cells more sensitive to CPT treatment (Fig 4D). Taken together, our results suggest that SLFN11 may specifically abrogate the DNA damage-induced checkpoint maintenance in the presence of DNA damage and then forces cells to undergo a lethal mitosis.

To verify and extend the above conclusions, we depleted SLFN11 expression in SF268 cells using two independent siRNAs specifically targeting SLFN11. As shown in Fig 4E, SLFN11 depletion did not result in any detectable change in initial CHK1 and CHK2 phosphorylation upon CPT treatment. By contrast, SLFN11 depletion blocked the rapid decline of both CHK1 and CHK2 phosphorylation after CPT withdrawal (Fig 4E). Similar results were obtained using DU145 cells (Fig EV1A). To rule out the possible off-target effects of the siRNA duplexes and confirm the aforementioned results, we took advantage of the newly described CRISPR/Cas9 gene-editing approach to generate a SF268 cell line that is specifically defective in SLFN11 (Fig 4F). Like siRNA-treated SF268 cells, SLFN11 knockout SF268 cells were proficient in both checkpoint initiation and maintenance (Figs 4F and EV1B). Consistently, SLFN11 silencing did not affect DNA replication under normal conditions, but resulted in the arrest of cells in G2 phase for prolonged periods after CPT withdrawal (Fig EV2). In order to further ensure that the observed phenotype was directly related to the loss of SLFN11, we introduced Flag-tagged SLFN11 into SLFN11-deficient SF268 cells and performed recovery experiments. Notably, re-expression of SLFN11 rendered SLFN11-deficient SF268 cells unable to maintain DNA damage checkpoint, indicating that the phenotype caused by the SLFN11 deficiency is SLFN11-dependent (Fig 4G).

SLFN11 inhibits homologous recombination repair

Homologous recombination (HR) repair is essential for the maintenance of genome stability, since it allows efficient and precise repair of double-strand breaks and is crucial for restarting collapsed DNA replication forks [3,23,24]. Cells defective in HR are often hypersensitive to DNA-damaging agents and exhibit increased chromosome instability [3,23,24]. For instance, BRCA1- or BRCA2-deficient cells are hypersensitive to a wide variety of anticancer drugs, such as CPT, etoposide, and cisplatin [3,23,24]. HR repair is initiated by nuclease-mediated DNA-end resection to generate 3' ssDNA overhangs that are initially bound and stabilized by the ssDNA-binding protein complex RPA [3,23,24]. As the interaction between SLFN11 and RPA serves to recruit SLFN11 to regions of ssDNA that are formed during DNA damage, it will be of interest to determine whether SLFN11 also plays a possible role in HR. We employed the sister chromatid exchange (SCE) assay in SLFN11-deficient SF268 cells to investigate whether SLFN11 is involved in SCE suppression. Strikingly, the SCE level in SLFN11-deficient SF268 cells was significantly elevated when compared to wild-type cells (Fig 5A and B). These results suggest that SLFN11 is able to suppress SCE in SF268

In order to further confirm the role of SLFN11 in HR, we engineered a DR-GFP U2OS cell line stably expressing exogenous SLFN11 and then performed a gene conversion assay to examine HR efficiency using the DR-GFP reporter system [25,26] (Fig 5C). Consistent with the notion that SLFN11 suppresses SCE, overexpressing exogenous SLFN11 in U2OS cells decreased the frequency of HR-mediated repair by approximately threefold (Fig 5D and E).

SLFN11 promotes the destabilization of RPA-ssDNA complex

Since SLFN11 forms a complex with RPA and contributes to the defects in checkpoint maintenance and HR repair in cells expressing

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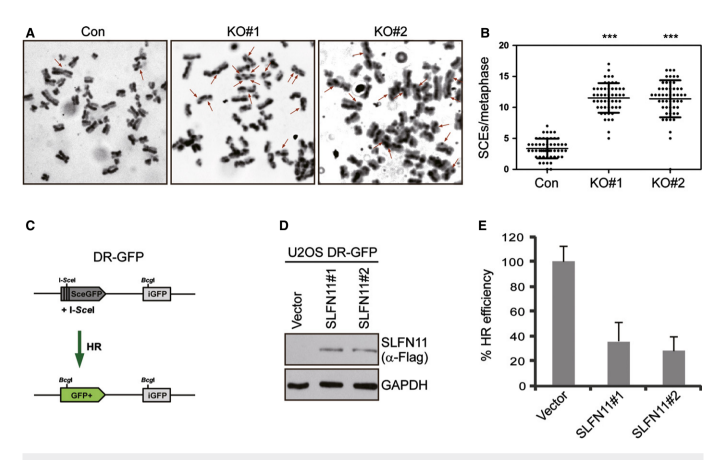


Figure 5. SLFN11 inhibits homologous recombination repair.

- A, B SLFN11 suppresses SCE. Representative metaphase spreads showing SCEs from wild-type and SLFN11-deficient SF268 cells (red arrows) (A). SCEs were scored for 50 metaphase spreads for each cell line (B). Each point represents the total number of SCEs in a single metaphase spread, and black bars indicate the average value for all spreads. Statistical significance was determined using the Mann–Whitney U-test (***P < 0.0001).
- C—E Overexpression of exogenous SLFN11 in U2OS cells results in decreased HR-directed DNA repair. A U2OS DR-GFP cell line stably expressing Flag-tagged SLFN11 was generated (C, D). The resulting cells were electroporated with an I-Scel expression plasmid. Forty-eight hours after electroporation, cells were harvested and assayed for GFP expression by FACS analysis. Error bars represent SD; n = 3 (E).

Source data are available online for this figure.

high endogenous SLFN11, we speculate that SLFN11 might be able to promote the destabilization of RPA-ssDNA complex. To test this hypothesis, we conducted an immunostaining experiment and examined the RPA2 focus formation in wild-type versus SLFN11deficient SF268 cells at different time points post-CPT removal. As shown in Fig 6A and B, we failed to observe any obvious difference in RPA2 focus formation between wild-type and SLFN11deficient SF268 cells at 1 h after CPT treatment, indicating that SLFN11 has no effect on the initial recruitment of RPA to ssDNA after onset of DNA damage. By contrast, the decline in RPA2 foci occurred much later in SLFN11-deficient SF268 cells, as compared to wild-type cells (Fig 6A and B). Similar results were observed when cells were treated with IR (Fig EV3). We also examined the effect of SLFN11 silencing on RPA2 focus formation in DU145 cells. As shown in Fig 6C and D, depletion of SLFN11 rendered the DU145 cells able to retain RPA2 foci after CPT withdrawal. Moreover, compared with wild-type SF268 cells, SLFN11-deficient SF268 cells exhibited a substantial increase in chromatin-bound RPA after CPT withdrawal (Fig 6E).

We further examined the direct readout of ssDNA production at DNA damage sites by using a method to detect BrdU-substituted

cellular DNA under non-denaturing conditions [20,27]. In agreement with the observation that SLFN11 does not affect the initial recruitment of RPA2 to DNA damage sites (Fig 6A–D), we failed to observe any obvious difference in BrdU-labeled ssDNA between wild-type and SLFN11-deficient SF268 cells at 1 h after CPT treatment (Fig 6F and G). However, while BrdU-labeled ssDNA was persistent in SLFN11-deficient SF268 cells even after 48 h of CPT withdrawal, it was significantly degraded in SLFN11-proficient SF268 cells after a withdrawal period of 24 h (Fig 6F and G). Taken together, these results indicated that SLFN11 might inhibit checkpoint maintenance and HR repair by promoting the destabilization of RPA–ssDNA complex.

The RPA1-binding ability of SLFN11 is required for its function in DNA damage response

SLFN11 is a member of a protein family with structural similarity to RNA helicases and has been shown to bind transfer RNA and selectively inhibit the expression of viral proteins [12,13]. To elucidate whether this putative RNA helicase activity of SLFN11 is important for its function in inhibiting checkpoint maintenance, we generated

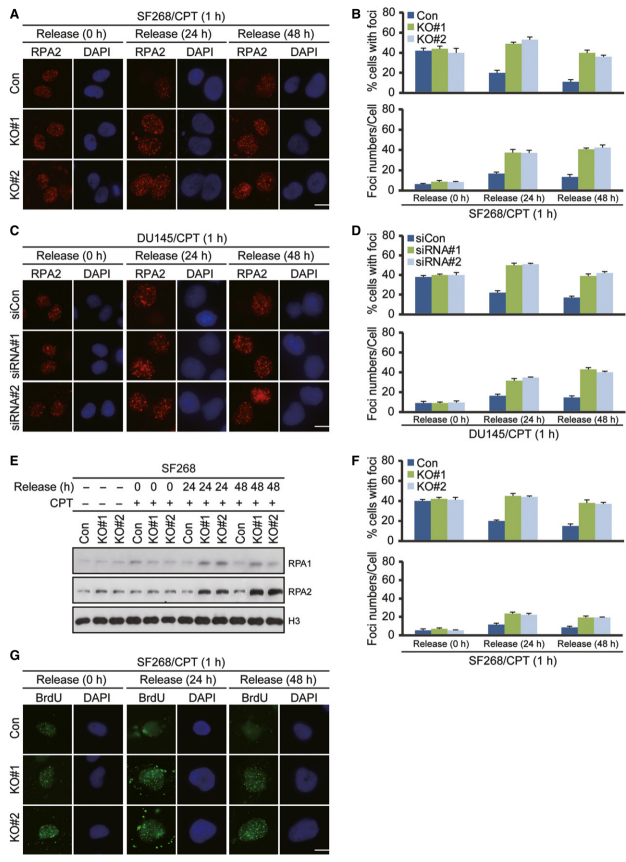


Figure 6.

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Figure 6. SLFN11 promotes the destabilization of RPA-ssDNA complex.

- A, B Loss of SLFN11 in SF268 cells delays the decline of RPA foci. Wild-type or SLFN11-deficient SF268 cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and processed at the indicated time points for immunofluorescence by using anti-RPA2 antibody. Representative RPA2 foci were shown (A). Scale bar, 10 μm. Quantification of RPA2 foci formation using NIH Image| software (B). Error bars represent SD; n = 3.
- C, D SLFN11 depletion in DU145 cells delays the decline of RPA foci. Wild-type or SLFN11-depleted DU145 cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and processed at the indicated time points for immunofluorescence by using anti-RPA2 antibody. Representative RPA2 foci were shown (C). Scale bar, 10 μm. Quantification of RPA2 foci formation using NIH ImageJ software (D). Error bars represent SD; n = 3.
- E SLFN11-deficient SF268 cells exhibited a substantial increase in chromatin-bound RPA after CPT withdrawal. Wild-type or SLFN11-deficient SF268 cells were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and harvested at the indicated time points for immunoblotting with the indicated antibodies.
- F, G Loss of SLFN11 in SF268 cells delays the decline of BrdU foci. SLFN11 wild-type or SLFN11-deficient SF268 cells were labeled with BrdU and 24 h later were treated with CPT (1 μM) for 1 h. Cells were then washed, shifted to fresh medium (time 0), and processed at the indicated time points for immunofluorescence by using anti-BrdU antibody under non-denaturing conditions. Representative BrdU foci were shown (G). Scale bar, 10 μm. Quantification of BrdU foci formation using NIH ImageJ software (F). Error bars represent SD; n = 3.

Source data are available online for this figure.

a double-point mutant (K605M/D668A) (Lys 605 and Asp 668 are the conserved residues within the Walker A and Walker B motifs of SLFN11, and mutations of the analogous residues in other helicases typically produce helicase-deficient enzymes). As expected, this mutation abolished the ATPase activity of SLFN11 (Fig EV4A and B). Notably, re-introduction of both wild-type SLFN11 and the K605M/D668A mutant into SLFN11-deficient SF268 cells inhibited checkpoint maintenance and the retention of RPA2 and RAD51 at sites of DNA damage after CPT withdrawal (Fig 7A–D). Likewise, re-introduction of both wild-type SLFN11 and the K605M/D668A mutant re-sensitized the SLFN11-deficient SF268 cells to DNA-damaging agents (Figs 7E and EV4C and D). These results suggest that the putative RNA helicase activity of SLFN11 is not essential for its role in the regulation of DNA damage response.

To investigate the biological significance of the direct interaction between SLFN11 and RPA1, we performed rescue experiments similar to those described above. As shown in Fig 7A–E, the phenotypes observed in SLFN11-deficient SF268 cells cannot be reversed by the re-expression of the SLFN11 deletion mutant defective in RPA1 binding (Δ 5). These results indicated that the role of SLFN11 in regulating DNA damage response depends on a physical protein–protein interaction between SLFN11 and RPA1.

Discussion

Defects in DNA damage repair and checkpoint control can result in gene mutations, chromosomal instability, and aneuploidy all of which can contribute to tumorigenesis [1–5]. On the other hand, defective DNA repair and checkpoint control also make tumor cells more vulnerable to chemotherapy regimens that damage DNA

[1-5]. SLFN11 is a nuclear protein and its expression is causally associated with the activity of DNA-damaging agents in human cancer cells [11,14]. In addition, the expression levels of SLFN11 can be used as a predictive marker for ovarian cancer patients treated with cisplatin-based chemotherapy [14]. Understanding the molecular mechanisms behind SLFN11 involvement in DNA damage response will help clarify its clinical importance and may pave the way for human cancer therapies. In this study, we have provided several lines of evidence to show that SLFN11 is a novel endogenous inhibitor of DNA damage response in cancer cells expressing high levels of SLFN11. First, cells expressing SLFN11 at high levels (SF268 and DU145) had a markedly earlier decline of CHK1 and CHK2 phosphorylation when compared with cells expressing SLFN11 at very low or undetectable levels (HeLa and U2OS) (Fig 4A). Second, in contrast to their respective wild-type counterparts, SLFN11-deficient SF268 and DU145 cells were proficient in checkpoint maintenance (Figs 4E and F, and EV1) and were resistant not only to Top1/2 inhibitors, DNA synthesis inhibitors, and alkylating agents [14], but also to IR and UV radiation (Fig EV4C and D). Third, both HeLa and U2OS cells overexpressing exogenous SLFN11 displayed a specific defect in checkpoint maintenance (Fig 4B and C). Consistently, overexpression of exogenous SLFN11 rendered both HeLa and U2OS cells more sensitive to CPT treatment (Fig 4D). Fourth, while the SCE level in SLFN11-deficient SF268 cells was significantly elevated when compared to wild-type SF268 cells, U2OS cells stably expressing exogenous SLFN11 showed a dramatic reduction in the frequency of HR repair (Fig 5). Finally, SLFN11 was capable of promoting the destabilization of RPA-ssDNA complex (Fig 6). Based on these results, we propose that, upon DNA damage, ssDNA generated by nucleolytic processing of stalled or collapsed replication forks or by processing of DNA

Figure 7. The RPA1-binding ability of SLFN11 is required for its function in DNA damage response.

- A–C A SLFN11-deficient SF268 cell line to express Flag-tagged wild-type SLFN11, the K605M/D668A mutant, or the Δ5 mutant was generated. The resulting cell line was treated with CPT (1 μM) for 1 h. Cells were then washed and shifted to fresh medium. Twenty-four hours later, cells were fixed and processed for RPA2 or RAD51 immunofluorescence. Scale bar, 10 μm. Error bars represent SD; n = 3.
- D The ability of SLFN11-deficient SF268 cells to maintain DNA damage checkpoint was disrupted by the re-expression of wild-type SLFN11 and the K605M/D668A mutant, but not the Δ5 mutant. Cells were treated with CPT (1 μM) for 1 h, then washed and shifted to fresh medium. Twenty-four hours later, cells were harvested for immunoblotting with the indicated antibodies.
- E Re-introduction of wild-type SLFN11 and the K605M/D668A mutant, but not the Δ5 mutant, re-sensitized the SLFN11-deficient SF268 cells to DNA-damaging agents. Cells were treated with various doses of CPT for 24 h, then shifted to fresh medium and permitted to grow for 14 days before staining. Error bars represent SD: n = 3.
- F A proposed model for the role of SLFN11 in the regulation of DNA damage response. Please refer to the main text for details.

Source data are available online for this figure.

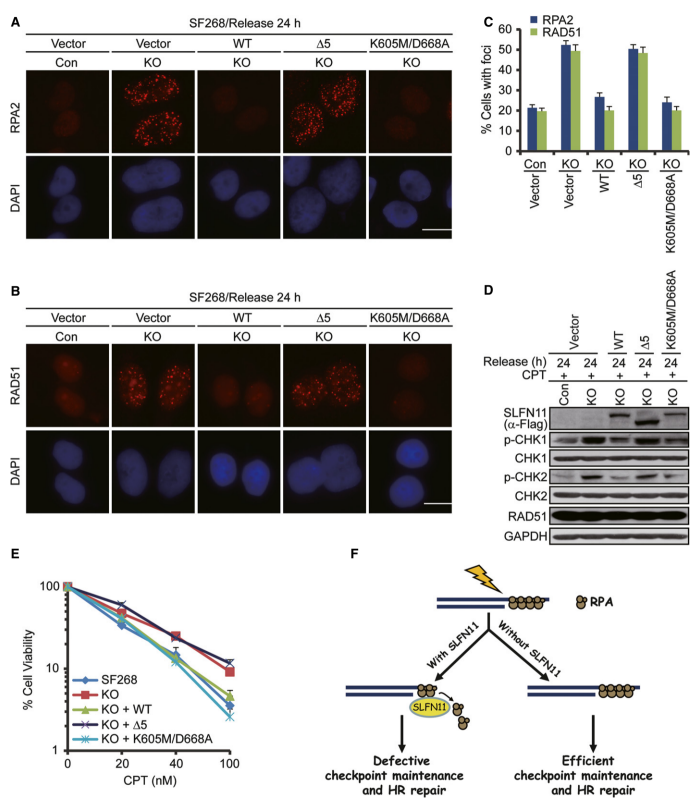


Figure 7.

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nicks and breaks is bound by RPA. Through the direct interaction with RPA1, SLFN11 is recruited to sites of DNA damage and promotes the destabilization of RPA–ssDNA complex and thus to inhibit checkpoint maintenance and HR repair (Fig 7F).

Besides SLFN11, RPA interacts with a large number of proteins involved in DNA replication, cell cycle regulation, and DNA repair and has been considered as a platform that promotes critical biochemical reactions that occur at ssDNA. For example, we

and several other groups have demonstrated that hPrimpol1/CCDC111/Primpol [18,28–31], the PSO4/PRP19 complex [19,32], HARP/SMARCAL1 [33–37], and several other proteins possessing enzymatic activities can be recruited to sites of DNA damage through the direct interaction with RPA. While SLFN11 inhibits checkpoint maintenance and HR repair, almost all the known RPA-binding proteins positively regulate DNA damage response. Whether and how SLFN11 competes with these proteins to modulate the DNA damage response remains to be investigated in the further studies.

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SLFN11 contains a putative RNA helicase domain and has been shown to bind transfer RNA [12,13]. While it is unclear whether the transfer RNA-binding ability of SLFN11 has any role in promoting the destabilization of RPA-ssDNA complex, there is evidence that the putative RNA helicase activity is not involved in this process (Figs 7A-E and EV4A and B). By contrast, the RPA1-binding ability is essential for its function in destabilizing RPA-ssDNA complex (Fig 7A-E). We propose that the direct interaction between SLFN11 and RPA1 may induce a conformational change of RPA and then favors its release from ssDNA, leading to ssDNA degradation as well as defective checkpoint maintenance and HR repair. This hypothesis is reminiscent of a previous study demonstrating that PARI acts in a non-enzymatic stoichiometric mode to promote the removal of RAD51 from ssDNA [38]. Similar to the situation with PARI, we cannot rule out the possibility that some sort of enzymatic activity (e.g., nuclease activity) is provided in trans by a vet-to-be-identified SLFN11-associated protein. The exact mechanism by which SLFN11 promotes the destabilization of RPA-ssDNA complex requires considerable future analysis.

In summary, we have demonstrated that human SLFN11 interacts with RPA and negatively regulates the DNA damage response by inhibiting checkpoint maintenance and HR repair (Fig 7F). Our findings, together with the previous observation that high expression levels of SLFN11 are strongly correlated with longer overall survival in a group of ovarian cancer patients treated with cisplatin-based chemotherapy regimens [11,14], raise the possibility of a strategy to increase the anti-neoplastic effects of the chemotherapeutic agents by developing drugs that specifically enhance SLFN11 expression.

Materials and Methods

Antibodies

Rabbit polyclonal anti-SLFN11 and anti-RPA2 antibodies were generated by immunizing rabbits with GST-SLFN11 (residues 1–300) or MBP-RPA2 fusion proteins expressed and purified from *E. coli*, respectively. Antisera were affinity-purified using Amino-Link plus Immobilization and purification kit (Pierce). Monoclonal anti-RPA2 antibody and polyclonal anti-phospho-RPA2 (S4/S8) were purchased from Abcam and Bethyl Laboratories, respectively. Anti-GAPDH and anti-H3 antibodies were purchased from Millipore. Anti-CHK1 and anti-CHK2 antibodies were purchased from Santa Cruz. Anti-CHK1 pS317 and anti-CHK2 pT68 antibodies were purchased from Cell Signaling Technology. Anti-Mre11 and anti-CtIP antibodies were purchased from Gene Tex and Bethyl Laboratories, respectively. Anti-Flag and anti-Myc antibodies were purchased from Sigma and Covance, respectively. Antibody specifically recognizing RAD15 was previously described [39].

Constructs

All cDNAs were subcloned into pDONR201 or pDONR221 (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 QuickChange site-directed mutagenesis kit (Stratagene) and verified by sequencing.

Cell culture and transfection

SF268 and HeLa Cells were maintained in DMEM supplemented with 10% fetal bovine serum and 1% penicillin and streptomycin. DU145 cells were maintained in MEM supplemented with 10% fetal bovine serum and 1% penicillin and streptomycin. U2OS cells with DR-GFP integration were kindly provided by Maria Jasin at Memorial Sloan-Kettering Cancer Center (New York). Human cell lines were maintained in 37°C incubator with 5% $\rm CO_2$. Cell transfection was performed using Lipofectamine 2000 (Invitrogen), following the manufacturer's protocol.

RNA interference

All siRNAs were synthesized by Dharmacon, Inc. The siRNAs were 21 base pairs and sequences are as follows: Con: UUCAAUAAAUU CUUGAGGUUU; SLFN11#1: CAGGGAACCUUACGAAUUAdTdT; SLFN11#2: GGUAUUUCCUGAAGCCGAAdT dT; Mre11: GGAGGUA CGUCGUUUCAGAdTdT; and CtIP: GCUAAAACAGGAACG AAUC dTdT. The siRNA transfection was performed with 100 nM siRNA duplexes using Lipofectamine RNAiMAX (Invitrogen) following the manufacturer's instruction. Transfection was repeated twice with an interval of 24 h to achieve maximal RNAi effect. Lentiviral nonsilencing control shRNA and shRNA target sets were purchased from Open Biosystems. The RPA1 targeting sequences are as follows: #1, 5'-GCTACAAAGCGTTTCTTTA-3'; #2, 5'-GAGTCAGAT GGGTCTGATA-3'. The non-silencing control sequence is as follows: 5'-CCCATAAGAGTAATAT-3'. The shRNAs were packaged into lentiviruses by co-transfecting with packaging plasmids pMD2G and pSPAX2 (kindly provided by Songyang Zhou, Baylor College of Medicine) into HEK293T cells. Forty-eight hours after transfection, the supernatant was collected for infection of SF268 cells. Infection was repeated twice with an interval of 24 h to achieve maximal infection efficiency.

Laser micro-irradiation

DNA double-strand breaks were introduced in the nuclei of cultured cells by micro-irradiation with a pulsed nitrogen laser (365 nm, 20 Hz pulse). Briefly, cells cultured on 35-mm glass bottom dishes were maintained with 10 μM BrdU for 24 h before being visualized with a Nikon Eclipse Ti-E inverted microscope equipped with a computer-controlled MicroPoint laser Ablation System (Photonics Instruments). The output of laser power was set to 7 pulses and 30% of transmission. Forty minutes later, cells were 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 then incubated in primary antibodies against SLFN11 and RPA2 for 30 min at room temperature. Following three 5-min washed with

PBS, cells were incubated with the appropriate secondary antibodies for 30 min. The cells were further counterstained with DAPI to visualize nuclear DNA. Images were captured using MetaMorph microscope automation and image analysis software and analyzed using Adobe Photoshop CS5.

Immunofluorescence staining

Indirect immunofluorescence was carried out as described [40,41]. Cells cultured on coverslips were treated with CPT or IR and recovered for the indicated times. 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 then incubated in primary antibody for 30 min at room temperature. Following three 5-min washed with PBS, secondary antibody was added at room temperature for 30 min. Cells were then stained with DAPI to visualize nuclear DNA. Images were captured with use of a fluorescence microscope (Eclipse 80i; Nikon) equipped with a Plan Fluor 60× oil objective lens (NA 0.5-1.25; Nikon) and a camera (CoolSNAP HQ2; PHOTO-METRICS). Images were captured using NIS-Elements basic research imaging software (Nikon) and analyzed using Adobe Photoshop CS5. For the quantification of RPA2 foci formation using the ImageJ software, images were converted into 8-bit grayscale pictures and fluorescence intensity threshold was set for images based on a black and white intensity scale ranging between 50 and 255.

Detecting BrdU incorporation in ssDNA

The experiment was performed as described previously [20,27]. Briefly, SF268 cells were grown in culture medium containing 10 μM BrdU (Sigma) for 24 h prior to treatment with CPT (1 μM). One hour later, cells were pre-extracted with buffer containing 0.5% Triton X-100 for 5 min and fixed with 3% paraformaldehyde solution for 10 min at room temperature without any DNA denaturation treatment. Cells were incubated in anti-BrdU antibody for 30 min at room temperature. Following three 5-min washed with PBS, secondary antibody was added at room temperature for 30 min. Cells were then stained with DAPI to visualize nuclear DNA.

Co-immunoprecipitation and Western blotting

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For transient transfection and co-immunoprecipitation assays, constructs encoding SFB-tagged and Myc-tagged proteins were transiently co-transfected into HEK293 cells. The transfected cells were lysed with NETN buffer (20 mM Tris–HCl, pH 8.0, 100 mM NaCl, 1 mM EDTA, 0.5% Nonidet P-40) containing 20 mM NaF, and 1 µg/ml of pepstatin A and aprotinin on ice for 20 min. After removal of cell debris by centrifugation, the soluble fractions were collected and incubated with S-protein beads for 4 h at 4°C. Beads were washed three times with NTEN buffer, boiled in 2× SDS loading buffer, and resolved on SDS–PAGE. For endogenous immunoprecipitations, the cells were solubilized in NETN lysis buffer supplemented with 500 U/ml benzonase (Novagen), protease inhibitors, and phosphatase inhibitors. After removal of cell debris by centrifugation, the soluble fractions were collected. One milligram of the whole-cell extract was then incubated with 25 µl of a 1:1

slurry of protein A–Sepharose coupled with 2 μ g of the indicated antibodies for 2 h at 4°C. The Sepharose beads were washed three times with NTEN buffer, boiled in 2× SDS loading buffer, and resolved on SDS–PAGE. Membranes were blocked in 5% milk in TBST buffer and then probed with antibodies as indicated.

Tandem affinity purification of SFB-tagged SLFN11

HEK293 cells were transfected with plasmids encoding SFB-tagged SLFN11. Cell lines stably expressing SLFN11 were selected by culturing in medium containing puromycin (2 µg/ml) and confirmed by immunoblotting and immunostaining. For affinity purification, cells stably expressing SFB-tagged SLFN11 were lysed with NETN buffer (20 mM Tris-HCl [pH 8.0], 100 mM NaCl, 1 mM EDTA, and 0.5% Nonidet P-40) containing benzonase (Novagen) for 20 min. The supernatants were cleared at 20,817 g to remove debris and then incubated with streptavidin-conjugated beads (Amersham Biosciences) for 4 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 again 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.

CRISPR-Cas9 approach to generate SLFN11 knockout SF268 cells

The guide RNA (gRNA) sequences designed to target the human SLFN11 gene are as follows: SLFN11#1: GTTGAGCATCCCGTGG AGAT; SLFN11#2: GGTCAGCAGGA TCCGAGTTA. gRNA sequences were purchased as DNA oligonucleotides and cloned into the pX330-U6-Chimeric_BB-CBh-hSpCas9 gRNA cloning vector (a gift from Dr. Feng Zhang) according to the standard protocol [42,43]. Twenty-four hours after transfection with the gRNA/Cas9 expression construct, SF268 cells were cultured in the presence of puromycin (2 μ g/ml). Approximately 2–3 weeks later, puromycin-resistant clones were then picked and expanded. Lack of SLFN11 protein expression in SF268 knockout cells was confirmed by Western blotting with SLFN11-specific antibody.

Cell survival assays

Cells (2×10^2) were seeded onto 35-mm dish in triplicates. At 24 h after seeding, cells were treated with CPT. The medium was replaced 24 h later, and cells were then incubated for 14 days. Resulting colonies were fixed and stained with Coomassie blue, and numbers of colonies were counted.

Lentivirus packaging and infection

Tet-On inducible Flag-tagged lentiviral vector and packaging plasmids (pMD2G and pSPAX2) were kindly provided by Professor Songyang Zhou (Baylor College of Medicine). SLFN11 entry constructs were transferred into the Gateway-compatible Flagtagged lentiviral vector. Virus supernatant was collected 48 h after the co-transfection of lentiviral vectors and packaging plasmids (pMD2G and pSPAX2) into HEK293T cells. HeLa or U2OS cells were infected with viral supernatants with the addition of 8 µg/ml

polybrene (Sigma), and stable pools were selected with medium containing 500 $\mu g/ml$ G418 (Calbiochem). The expression of the indicated genes in the stable pools was induced by the addition of 1 $\mu g/ml$ doxycycline (Sigma) for 24 h for the experiments presented in this report.

Chromatin fractionation

Preparation of chromatin fractions was as described previously [40,41] with modifications. Briefly, 1 h after treatment with CPT, cells were washed with PBS for 3 times and maintained with fresh medium lacking CPT for 24 or 48 h. Cells were then collected, and cell pellets were subsequently re-suspended in NETN buffer (20 mM Tris–HCl [pH 8.0], 100 mM NaCl, 1 mM EDTA, and 0.5% Nonidet P-40) and incubated on ice for 15 min. Nuclei were then recovered and re-suspended in HCl (0.2 M). The soluble fraction was neutralized with Tris–HCl (1 M at pH 8.0) for further analysis.

Sister chromatid exchange assay

Sister chromatid exchange staining was performed as described [39,44]. Briefly, SF268 cells were grown through two cell cycles to achieve preferential labeling of sister chromatid in the presence of 100 μM of 5-bromodeoxyuridine (BrdU). Colcemid (Sigma) was added at a final concentration of 0.2 $\mu\text{g/ml}$ to accumulate mitotic cells 4 h prior to harvesting cells. Harvested cells were swollen in hypotonic solution (75 mM KCl) for 20 min at room temperature and then fixed with 3:1 (vol/vol) methanol/acetic acid. Fixed cell suspension was dropped onto a glass slide and air-dried. Slides at least 1 day old were heated at 88°C for 20 min in buffer (1 M NaH₂PO₄ [pH 8.0]), rinsed in distilled water, and stained with 5% Giemsa for 5 min. Slides were then rinsed and air-dried. More than 2000 chromosomes were scored for each cell group. Statistically analyses of the SCE data were subjected to Student's two-tailed t-test.

Gene conversion assay

A U2OS cell clone stably expressing HR reporter DR-GFP was described previously [25]. This reporter consists of two differentially mutated GFP genes oriented as direct repeats. Expression of I-SceI endonuclease will generate a site-specific DSB between the mutated GFP genes, which when repaired by gene conversion, results in a functional GFP gene. Briefly, 1×10^6 U2OS DR-GFP cells were electroporated with 12 μg of pCBASce plasmid (an I-SceI expression vector) at 270 V, 975 μF using a Bio-Rad Genepulsar II. Cells were plated onto 10-cm dishes and harvested 2 days after electroporation and subjected to flow cytometry analysis to determine percentages of GFP-positive cells, which result from HR repair induced by DNA DSBs. Means were obtained from three independent experiments.

Protein purification

Full-length SLFN11 or RPA1 was cloned into GST-tagged or MBP-His-tagged vector for the expression of GST-tagged SLFN11 or MBP-His-tagged-RPA1 protein in insect cells. Transposition occurred in DH10 Bac-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 GST-SLFN11 or MBP-RPA1-His for 48 h. Cells were harvested and washed with 1× PBS and re-suspended in lysis buffer (20 mM Tris-HCl, 300 mM NaCl, 1% Triton X-100, 1 mM EDTA, 10 mM 2-mercaptoethanol, and 1 μg/ml each of leupeptin, aprotinin, and pepstatin). After sonicating, the extract was centrifuged at 18,300 g for 40 min. The supernatant was collected and incubated with glutathione-Sepharose resins or Amylose resins for 2 h at 4°C. The bound RPA1 protein was used for pull-down assays, and the bound SLFN11 protein was eluted with lysis buffer containing 20 mM of reduced glutathione. Full-length RPA2 and RPA3 were cloned into pDONR201 as entry clones and were then transferred to destination vector for the expression of MBP-tagged fusion protein in E. coli. Cells were harvested and re-suspended in lysis buffer (20 mM Tris-HCl, 300 mM NaCl, 1% Triton X-100, and 1 µg/ml each of leupeptin, aprotinin, and pepstatin). After sonicating, the extract was centrifuged at 18,000 rpm for 40 min. The supernatant was collected and incubated with Amylose resins for 2 h at 4°C. After washing the beads with washing buffer (20 mM Tris-HCl, 500 mM NaCl, 0.5% NP-40, 1 mM DTT, and 1 $\mu g/ml$ each of leupeptin, aprotinin, and pepstatin), the bound protein was used for pull-down assays.

BrdU incorporation assays

BrdU (100 μ M) was added into the medium for 1 h. Cells were then harvested, washed with PBS, and fixed with ice-cold 70% ethanol overnight. Once centrifuged, cells were washed with PBS. DNA was denatured with 2.5 M HCl for 1 h at room temperature. After washing 3 times with PBS, cells were incubated in mouse anti-BrdU antibody (Roche) diluted 1:100 in blocking buffer (PBS + 0.1% Triton X-100 + 5% BSA) for 12 h followed by 3 times wash with blocking buffer containing 500 mM NaCl. FITC conjugated goat anti-mouse IgG (1:100, Jackson immunoresearch) was added and incubated for 4 h. Following wash with blocking buffer containing 500 mM NaCl, cells were re-suspended in PBS containing propidium iodide (20 μ g/ml) and RNase A (200 μ g/ml) at 37°C for 20 min. Cell cycle distribution was analyzed on a FACScan flow cytometer (Beckman).

Cell cycle analysis

Cells were harvested, washed with PBS, and fixed with ice-cold 70% ethanol overnight. Cells were then washed in PBS and treated for 20 min at 37°C with RNase A (200 μ g/ml), followed with propidium iodide (25 μ g/ml), and analyzed on a FACScan flow cytometer (Beckman).

ATPase assays

Wild-type SLFN11 and the K605M/D668A mutant were cloned into pCold vector for the expression of MBP-tagged fusion protein in *E. coli*. Cells were harvested and re-suspended in NETN buffer containing 500 mM NaCl and 10 mM MgCl₂ by sonication. The supernatant was collected and incubated with Amylose resins. After washing the beads with washing buffer, the bound protein was eluted using 10 mM maltose. The released protein was loaded on UNOsphere Anion Exchange column and then eluted using stepwise

gradient of NaCl in $1\times$ NETN buffer. For determining ATPase activity, Malachite green reaction-based colorimetric assay system was used. Assay was performed as per manufacture's instructions (Sigma-Aldrich). Briefly, Equimolar (300 nM) amount of proteins was incubated in $1\times$ assay buffer containing 40 mM Tris, 80 mM NaCl, 8 mM MgAc₂, 1 mM EDTA, pH 7.5 along with 1 mM ATP at 25°C in a final volume of 40 µl for the indicated time. Reaction was stopped by addition of 200 µl Malachite green solution and incubated for additional 30 min. Absorbance was recorded at 630 nM. Concentration of free phosphate [Pi] (µM) was calculated from the phosphate standard curve and plotted.

Expanded View for this article is available online.

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Author contributions

YM, JL, and MS performed the experiments; JH, JC, TL, BZ, and XF designed the experiments; JH, JC, and TL analyzed the data and wrote the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

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