PICH regulates the abundance and localization of SUMOylated proteins on mitotic chromosomes

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ABSTRACT

Proper chromosome segregation is essential for faithful cell division and if not maintained results in defective cell function caused by the abnormal distribution of genetic information. Polo-like kinase 1–interacting checkpoint helicase (PICH) is a DNA translocase essential for chromosome bridge resolution during mitosis. Its function in resolving chromosome bridges requires both DNA translocase activity and ability to bind chromosomal proteins modified by the small ubiquitin-like modifier (SUMO). However, it is unclear how these activities cooperate to resolve chromosome bridges. Here, we show that PICH specifically disperses SUMO2/3 foci on mitotic chromosomes. This PICH function is apparent toward SUMOylated topoisomerase II\textalpha\ (Topoll\textalpha) after inhibition of Topoll\textalpha by ICRF-193. Conditional depletion of PICH using the auxin-inducible degron (AID) system resulted in the retention of SUMO2/3-modified chromosomal proteins, including Topoll\textalpha, indicating that PICH functions to reduce the association of these proteins with chromosomes. Replacement of PICH with its translocase-deficient mutants led to increased SUMO2/3 foci on chromosomes, suggesting that the reduction of SUMO2/3 foci requires the remodeling activity of PICH. In vitro assays showed that PICH specifically attenuates SUMOylated Topoll\textalpha activity using its SUMO-binding ability. Taking the results together, we propose a novel function of PICH in remodeling SUMOylated proteins to ensure faithful chromosome segregation.

INTRODUCTION

Accurate chromosome segregation is a complex and highly regulated process during mitosis. Sister chromatid cohesion is necessary for proper chromosome alignment and is mediated by both cohesin and catenated DNA at centromeric regions (Michaelis et al., 1997; Losada et al., 1998; Bauer et al., 2012). Compared to the well-described regulation of cohesin (Morales and Losada, 2018), the regulation of catenated DNA cleavage by DNA topoisomerase II\textalpha (Topoll\textalpha) is not fully understood despite its critical role in chromosome segregation. ATP-dependent DNA decatenation by Topoll\textalpha takes place during the metaphase-to-anaphase transition, and this allows for proper chromosome segregation (Shamu and Murray, 1992; Wang et al., 2010; Gomez et al., 2014). Failure in resolution of catenanes by Topoll\textalpha leads to the formation of chromosome bridges and ultrafine DNA bridges (UFBs) to which PICH localizes (Spence et al., 2007). PICH is a SNF2 family DNA translocase (Baumann et al., 2007; Biebricher et al., 2013), and its binding to UFBs recruits other proteins to UFBs (Chan et al., 2007; Hengeveld et al., 2015). In addition to the role in UFB binding during anaphase, PICH has been shown to play a key role in chromosome segregation at the metaphase-to-anaphase transition by optimizing the Os-TIR1 integration locus and creating constructs for aID-mediated degradation of Py-S2. B.T.L. performed initial analysis of immunofluorescent images in Figure 5. V.A., A.A., and M.D. established the AID-mediated degradation system by optimizing the Os-TIR1 integration locus and creating constructs for genome editing by CRISPR/Cas9 for that system. Y.A. designed the study, supervised the project, and wrote the manuscript.

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Abbreviations used: CSF, cytostatic factor; dnUbc9, dominant negative E2 SUMO-conjugating enzyme; PIAS, protein inhibitor of activated STAT; PICH, Polo-like kinase-interacting checkpoint helicase; SENP, sentrin-specific protease; SIM, SUMO-interacting motif; SPR, strand passage reaction; SUMO, small ubiquitin-like kinase-interacting checkpoint helicase; SENP, sentrin-specific protease; SIM, SUMO-interacting motif; Topoll, topoisomerase II\textalpha; XEE, Xenopus egg extract.

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transition (Baumann et al., 2007; Nielsen et al., 2015; Sridharan and Azuma, 2016).

Previously, we demonstrated that PICH binds SUMOylated proteins using its three SUMO-interacting motifs (SIMs) (Sridharan et al., 2015). PICH utilizes its ATPase activity to translocate on DNA, similar to known nucleosome-remodeling enzymes (Whitehouse et al., 2003); thus it is a putative remodeling enzyme for chromosomal proteins. But the nucleosome-remodeling activity of PICH was shown to be limited as compared with established nucleosome-remodeling factors (Ke et al., 2011). Therefore, the target of PICH-remodeling activity has not yet been determined. Importantly, both loss of function PICH mutants in either SUMO-binding activity or translocase activity showed chromosome bridge formation (Sridharan and Azuma, 2016), suggesting that both of these activities cooperate to accomplish proper chromosome segregation, albeit the molecular mechanism linking these two functions is unknown. Previous studies demonstrated that proper regulation of mitotic chromosomal SUMOylation is required for faithful chromosome segregation (Nacerdine et al., 2005; Diaz-Martinez et al., 2006; Cubenas-Potts et al., 2013). Studies using Caenorhabditis elegans demonstrated the dynamic nature of SUMOylated proteins during mitosis and its critical role in chromosome segregation (Pelisch et al., 2014). Several SUMOylated chromosomal proteins were identified for their potential role in chromosome segregation, for example, TopoII alpha, CENP-A, CENP-E, FoxM1, and Orc2 (Bachant et al., 2002; Zhang et al., 2008; Schimmel et al., 2014; Huang et al., 2016; Ohkuni et al., 2018). Because PICH is able to specifically interact with SUMO moieties (Sridharan et al., 2015), these SUMOylated chromosomal proteins could be targets of the SIM-dependent function of PICH in mediating faithful chromosome segregation. Among the known SUMOylated chromosomal proteins, TopoII alpha has been shown to functionally interact with PICH. PICH-knockout cells have increased sensitivity to ICRF-193, a potent TopoII catalytic inhibitor, accompanied by increased incidence of chromosome bridges, binucleation, and micronuclei formation (Wang et al., 2008; Kurasawa and Yu-Lee, 2010; Nielsen et al., 2015). ICRF-193 stalls TopoII alpha at the last step of the strand passage reaction (SPR) in which two DNA strands are trapped within the TopolIIx molecule without DNA strand breaks (Roca et al., 1994; Patel et al., 2000). In addition to that specific mode of inhibition, ICRF-193 has been shown to increase SUMOylation of TopolIIx (Agostinho et al., 2008; Pandey et al., 2020). Because PICH has SUMO-binding ability, it is possible that increased SUMOylation of TopolIIx contributes to interaction with PICH after ICRF-193 treatment. However, no study has shown a link-between SUMOylation of TopolIIx and PICH function.

To elucidate possible functional interactions of PICH with SUMOylated chromosomal proteins, we established the connection between PICH and SUMOylation by utilizing specific TopolII inhibitors and genome edited cell lines. Our results demonstrate that increased SUMOylation during ICRF-193 treatment leads to the recruitment and enrichment of PICH on chromosomes. Depletion of SUMOylation abrogates this enrichment, suggesting that PICH specifically targets SUMOylated chromosomal proteins. Depletion of PICH led to the retention of SUMOylated proteins, including SUMOylated TopolIIx on the chromosomes in ICRF-193-treated cells. Replacing endogenous PICH with a translocase-deficient PICH mutant resulted in increased SUMO2/3 foci on chromosomes where PICH was located, suggesting that PICH utilizes its translocase activity to remodel SUMOylated proteins on the chromosomes. In vitro assays showed that PICH specifically interacts with SUMOylated TopolIIx to attenuate SUMOylation TopolIIx activity in a SIM-dependent manner. Taking the results together, we propose a novel mechanism for PICH in promoting proper chromosome segregation during mitosis by remodeling SUMOylated proteins on mitotic chromosomes including TopolIIx.

RESULTS

TopolII inhibitor ICRF-193 induces increased SUMO2/3 modification and increased PICH foci on mitotic chromosomes

We previously reported that PICH utilized its SIMs for proper chromosome segregation and for its mitotic chromosomal localization (Sridharan and Azuma, 2016). We wished to examine whether modulating mitotic SUMOylation affected PICH localization on mitotic chromosomes. Treatment with ICRF-193, a catalytic inhibitor of TopolII that blocks TopolII at the last stage of its SPR, after DNA decatenation but before DNA release, increases SUMO2/3 modification of TopolIIx on mitotic chromosomes. In contrast, treatment with another catalytic TopolII inhibitor, merbarone, which blocks TopolII before the cleavage step of the SPR, does not affect the level of SUMO2/3 modification of TopolIIx (Agostinho et al., 2008; Pandey et al., 2020). We utilized these two contrasting inhibitors to assess whether TopolIIx inhibition and/or SUMOylation changes PICH distribution on mitotic chromosomes. DLD-1 cells were synchronized in prometaphase, mitotic cells were collected by mitotic shake off, treated with inhibitors for 20 min, and then plated onto fibronectin-coated coverslips. As seen in Western blot analysis, increased intensity of SUMO2/3 modification and increased PICH levels on mitotic chromosomes including TopolII alpha (marked by red asterisks in Figure 1A). Intriguingly, a novel finding showed that PICH levels on mitotic chromosomes were significantly increased in cells treated with ICRF-193. In contrast, merbarone did not increase the level of these proteins on the chromosomes, suggesting that there is a specificity of ICRF-193 inducing increased levels of PICH and SUMOylation of TopolIIx (Figure 1A).

To investigate the localization of PICH on mitotic chromosomes treated with ICRF-193, mitotic cells were subjected to immunofluorescence staining. Synchronized cells were collected by mitotic shake off, treated with inhibitors for 20 min, and then plated onto fibronectin-coated coverslips. As seen in Western blot analysis, increased intensity of SUMO2/3 foci were observed on the chromosomes, where they overlapped with TopolIIx foci upon ICRF-193 treatment (Figure 1B, enlarged images). Although the TopolIIx signal changed during merbarone treatment, showing a less punctate signal, no enrichment of SUMO2/3 foci was observed (Figure 1B). A novel observation showed that treatment with ICRF-193 caused a redistribution of PICH from all over the chromosomes to an enrichment at foci on the chromosomes where they overlapped with the SUMO2/3 foci (Figure 1C, enlarged images). Treatment with merbarone did not affect PICH localization (Figure 1C). By outlining single chromosomes using the DNA signal in multiple images and then placing outlines on SUMO2/3 or TopolIIx channels, the mean intensities of these signals were measured. Both TopolIIx and SUMO2/3 chromosome signal intensities were significantly higher after ICRF-193 treatment, but not in merbarone-treated cells (Figure 1D). PICH foci intensity was measured by using circles equal in size; the PICH foci intensity was found to be significantly increased in ICRF-193-treated cells (Figure 1D, bottom graph). These data show that treatment with ICRF-193, but not merbarone, induces increased TopolIIx SUMOylation and enrichment of PICH and SUMO2/3 foci on the chromosomes.
FIGURE 1: TopoIIα inhibition by ICRF-193 leads to increased PICH, SUMO2/3, and TopoIIα levels on mitotic chromosomes. (A) DLD-1 cells were synchronized and treated with the indicated inhibitors (7 μM ICRF-193: ICRF and 40 μM merbarone: MERB); DMSO was used as a control. Mitotic chromosomes were isolated and subjected to Western blotting with the indicated antibodies. Intensity of signals (arbitrary unit; a.u.) normalized by H2A are shown with mean value and SD. * indicates SUMOylated TopoIIα. p values for comparison from three experiments were calculated using a one-way ANOVA with Tukey multicomparison correction. ns: not significant; *: \( p \leq 0.05 \); **: \( p < 0.01 \); ***: \( p < 0.001 \). (B) Mitotic cells treated with DMSO (control), ICRF-193, and merbarone were stained with antibodies against TopoIIα (green) and SUMO2/3 (red). DNA was stained with DAPI (blue). Scale bar = 11 μm. The white square indicates enlarged area. (C) Mitotic cells were treated as in B and stained with antibodies against PICH (green) and SUMO2/3 (red). DNA was stained with DAPI (blue). Scale bar = 11 μm. The white square indicates enlarged area. (D) Using DAPI signal the mean intensities (a.u.) of each channel of at least five individual chromosomes per experimental replicate were measured. The bar indicates the mean value of the intensities. p values for comparison of all obtained values from three experiments were calculated using a one-way ANOVA with Tukey multicomparison correction ns: not significant; **: \( p < 0.01 \); ****: \( p < 0.0001 \).
SUMOylation is required for PICH enrichment in ICRF-193–treated cells

Although results obtained from inhibiting Topollα suggest that increased SUMOylation plays a critical role in PICH enrichment, the distinct effects of the different inhibitor treatments, for example, differences in Topoll conformation, could also play a role. To determine whether mitotic SUMOylation is critical for PICH enrichment in ICRF-193–treated cells, we developed a novel method to inhibit mitotic SUMOylation in cells. First, we generated a fusion protein, called Py-S2, which consists of the N-terminal region of human PIASy and the SENP2-catalytic domain (required for deSUMOylation) (Reverter and Lima, 2004; Ryu et al., 2015; Sridharan et al., 2015). The N-terminal region of PIASy localizes to mitotic chromosomes, in part via its specific interaction with the RZZ (Rod-Zw10-Zwilch) complex (Ryu and Azuma, 2010). Thus, the fusion protein is expected to bring deSUMOylation activity where mitotic SUMOylation occurs on chromosomes by PIASy. As a negative control, we substituted a cysteine to alanine at position 548 of SENP2 (called Py-S2 Mut) to create a loss of function mutant (Reverter and Lima, 2004, 2006) (Figure 2A). The activity of the recombinant fusion proteins on chromosomal SUMOylation was verified in Xenopus egg extract (XEE) assays (Supplemental Figure S1). As predicted, the addition of Py-S2 protein to XEE completely eliminated mitotic chromosomal SUMOylation. To our surprise, the Py-S2 Mut protein stabilized SUMOylation of chromosomal proteins, thus acting as a dominant negative mutant against endogenous deSUMOylation enzymes. To express the fusion proteins in cells, we created inducible expression cell lines using the tetracycline-inducible system (Supplemental Figure S2) (Natsume et al., 2016). We utilized CRISPR/Cas9 genome editing to integrate each of the fusion genes into the human H11 (hH11) safe harbor locus (Zhu et al., 2014; Ruan et al., 2015) in DLD-1 cells.

To test whether the Py-S2 fusion protein worked as expected, cells were synchronized and doxycycline was added after release from a thymidine block. After treatment with ICRF-193, chromosomes were isolated and subjected to Western blot analysis. The Py-S2–expressing cells had nearly undetectable levels of chromosomal SUMOylation as well as SUMOylated Topollα (Figure 2B). Intriguingly in Py-S2–expressing cells, PICH levels on chromosomes no longer showed a significant increase under ICRF-193 treatment, suggesting that the response of PICH to ICRF-193 depends on the cell’s ability to SUMOylate chromosomal proteins (Figure 2B, quantification). The role of SUMOylation in the enrichment of PICH on mitotic chromosomes is further supported by the Py-S2 Mut–expressing cells. Western blot analysis of mitotic chromosomes expressing Py-S2 Mut revealed a slight increase in overall SUMOylation levels in the absence of ICRF-193 (Figure 2C, lane 1 vs. lane 3, quantification). But SUMOylated Topollα did not show a significant increase in either control or ICRF-193–treated conditions. This suggests that a similar stabilization of SUMOylation occurs in cells as was observed in the XEE assays, albeit with less penetrance. PICH levels were also slightly increased in the Py-S2 Mut–expressing cells in the absence of ICRF-193 (Figure 2C, lane 1 vs. lane 3, quantification).

To determine how deSUMOylation affects PICH and SUMO2/3 distribution on chromosomes, Py-S2–expressing mitotic cells were subjected to immunofluorescence staining. Immunofluorescence analysis of Py-S2–expressing cells reiterated what was observed in Western blot analysis. By utilizing the same quantification methodology as in Figure 1, measuring individual chromosome mean intensities, we saw that even under ICRF-193 treatment, Py-S2–expressing cells displayed significantly lower levels of SUMO2/3 signals on chromosomes (Figure 3A, quantification). Then the distribution of the SUMO2/3 signal was assessed by calculating the ratio of SUMO2/3 foci intensity divided by the total intensity (hereafter this analysis will be referred to as “granularity” as marked on each graph). The results showed that expression of Py-S2 completely diminishes the enrichment of SUMO2/3 foci even after ICRF-193 treatment. PICH chromosomal intensity was also reduced in Py-S2–expressing cells, where no significant change in intensity or distribution occurred upon ICRF-193 treatment (Figure 3B, quantification). But total Topollα signals remained unaffected by inhibition of SUMOylation, agreeing with our previous observations in XEE assays (Azuma et al., 2005) (Figure 3C, quantification). By measuring Topollα granularity, the enrichment of Topollα foci with ICRF-193 treatment was still observed, suggesting that ICRF-193 treatment affects the distribution of Topollα independent of SUMOylation. Next the Py-S2 Mut–expressing cells were analyzed to examine the effect of stabilizing SUMOylation. The increase of both PICH and SUMO2/3 signals observed in Western blot quantification was also apparent with immunofluorescence analysis. In control cells (dimethyl sulfoxide [DMSO]) expressing Py-S2 Mut, both total SUMO2/3 intensity and granularity on chromosomes showed a significant increase. No additive effects were observed with ICRF-193 treatment in either total intensity or granularity (Figure 4A, quantifications). The discrepancy between Western blot data (Figure 2C) and quantification of immunofluorescence images implies that these increased SUMO2/3-modified proteins could be lost from chromosomal fractions during the isolation process for Western blotting. Intriguingly, in control cells (DMSO) expressing Py-S2 Mut, we observed increased chromosomal PICH intensity as compared with ICRF-193–treated Py-S2 Mut–expressing cells (Figure 4B, quantification), which mirrors the SUMO2/3 total intensity result. PICH granularity was not clearly increased by ICRF-193 treatment in Py-S2 Mut–expressing cells, and this also mirrors the SUMO2/3 granularity results. Similar to Figure 3C, Topollα signal intensity did not change upon Py-S2 Mut expression (Figure 4C, quantification). However, Topollα granularity is increased by expression of Py-S2 Mut without ICRF-193, suggesting that increased SUMO2/3 modification on chromosomes affects the chromosomal distribution of Topollα. In all, these data reinforce the indication that chromosomal SUMOylation plays a critical role in the enrichment of chromosomal PICH localization, including PICH foci formation in ICRF-193–treated cells.

Increased PICH levels observed after ICRF-193 treatment are lost upon Topollα depletion

ICRF-193 treatment led to a distinct enrichment of SUMO2/3 and PICH foci unlike simply stabilizing SUMO2/3 modification on chromosomal proteins by Py-S2 Mut expression. Therefore, we tested whether the PICH response to ICRF-193 is due to Topollα SUMOylation. To accomplish this, we generated an mAID-Topollα cell line that enables rapid and complete elimination of Topollα in the presence of auxin (Nishimura et al., 2009; Natsume et al., 2016). First, we established a cell line that has an integration of an auxin–dependent ubiquitin E3 ligase, OsTIR1 gene, at the promoter of a housekeeping (RCC1) gene (Supplemental Figure S3, A–C) using CRISPR/Cas9 editing technology (Yau et al., 2020). The integration of the OsTIR1 gene under the RCC1 promoter achieved stable and low-level expression of the protein and thus minimized the nonspecific degradation of AID-tagged proteins without auxin. Using the established OsTIR1-expressing DLD-1 cell line, DNA encoding a mAID-Flag tag was inserted into both Topollα loci (Supplemental Figure S4, A–C). After 6-h treatment with auxin, Topollα was degraded to undetectable levels in all cells analyzed (Supplemental Figure S4, D and E). This rapid elimination allowed us to examine the effect of Topollα depletion in a single cell cycle.
To deplete Topollα, the cells were treated with auxin after release from a thymidine block. After mitotic shake off and treatment with ICRF-193, isolated chromosomes were subjected to Western blotting with anti-SUMO2/3, anti-Topollα, and anti-PICH antibodies, and anti-H2A was used as a loading control. A slight overall increase of global SUMOylation was still observed in ΔTopollα cells treated with ICRF-193. This suggests that ICRF-193 affects SUMOylation of other chromosomal proteins, such as Topollβ (Figure 5A). Notably,
FIGURE 3: Decreased mitotic SUMOylation eliminates PICH response to ICRF-193. (A) Mitotic cells were fixed and stained with antibodies against SUMO2/3 (red) and mNeon (green). DNA was stained by DAPI (blue). Scale bar = 11 μm. The mean intensities of SUMO2/3 signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; *: p ≤ 0.05; ****: p < 0.0001. (B) Mitotic cells were fixed and stained with antibodies against PICH (red) and mNeon (green). DNA was stained by DAPI (blue). Scale bar = 11 μm. The mean intensities of PICH signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; *: p ≤ 0.05; **: p < 0.01; ****: p < 0.0001. (C) Mitotic cells were fixed and stained with antibodies against TopoIIα (red) and mNeon (green). DNA was stained by DAPI (blue). Scale bar = 11 μm. The mean intensities of TopoIIα signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; *: p ≤ 0.05; ****: p < 0.001.
FIGURE 4: Mutant form of deSUMOylation enzyme promotes PICH and SUMO2/3 foci in both control and ICRF-193–treated cells. (A) Mitotic cells were fixed and stained with antibodies against SUMO2/3 (red) and mNeon (green). DNA was stained by DAPI (blue). Scale bar = 11 μm. The mean intensities of SUMO2/3 signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; *: p ≤ 0.05; **: p < 0.01; ****: p < 0.0001. (B) Mitotic cells were fixed and stained with antibodies against PICH (red) and mNeon (green). DNA was stained by DAPI (blue). Scale bar = 11 μm. The mean intensities of PICH signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; *: p ≤ 0.05; **: p < 0.01; ****: p < 0.0001. (C) Mitotic cells were fixed and stained with antibodies against TopoIIα (red) and mNeon (green). DNA was stained by DAPI (blue). Scale bar = 11 μm. The mean intensities of TopoIIα signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; **: p < 0.01.
FIGURE 5: Depletion of Topolıa attenuates SUMO2/3 modification and eliminates PICH response in ICRF-193–treated cells. (A) DLD-1 cells with endogenous Topolıa tagged with a mAID were synchronized in mitosis and treated with DMSO (control) and ICRF-193. Auxin was added to the cells after release from thymidine for 6 h. Mitotic chromosomes were isolated and subjected to Western blotting with the indicated antibodies. * indicates SUMOylated Topolıa.

Intensities of signals (a.u.) normalized by H3 are shown with mean values and SD. p values for comparison from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; *: p ≤ 0.05; **: p < 0.01.

(B) Mitotic cells were fixed and stained with antibodies against PICH (green) and CENP-C (red). DNA was stained with DAPI (blue). Scale bar = 11 μm. The white square indicates enlarged area. The mean intensities of PICH signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; **: p < 0.01; ****: p < 0.0001.

(C) Mitotic cells were fixed and stained with antibodies against SUMO2/3 (green) and CENP-C (red). DNA was stained with DAPI (blue). Scale bar = 11 μm. The white square indicates enlarged area. The mean intensities of SUMO2/3 signals are shown. p values for comparison of all obtained values from three experiments were calculated using a two-way ANOVA with Tukey multicomparison correction; ns: not significant; **: p < 0.01; ***: p < 0.001; ****: p < 0.0001.
ΔTopoIIα cells treated with ICRF-193 showed no changes in PICH levels on the chromosomes. This suggests that increased levels of PICH seen in ICRF-193 treatment is a SUMOylated TopoIIα-dependent response (Figure 5A). Immunofluorescence analysis of PICH showed fewer foci on the chromosomes after ICRF-193 treatment in ΔTopoIIα cells as compared with cells with intact TopoIIα treated with ICRF-193. This observation was quantified by measuring the PICH chromosomal signal intensity, which showed a statistically significant decrease in mean PICH intensity in ICRF-193–treated ΔTopoIIα cells (Figure 5B, quantification). Further, measuring the centromeric PICH foci intensity and calculating the ratio of centromeric PICH among total PICH on chromosomes (Figure 5B, granularity quantification) revealed that when TopoIIα is depleted in cells, ICRF-193 treatment did not increase PICH. Similarly, ΔTopoIIα cells showed a significant decrease in SUMO2/3 chromosome signal intensity even under ICRF-193 treatment (Figure 5C, quantification). The discrepancy from Western blotting analysis (Figure 5A) could originate from the extraction conditions of the chromosome samples as indicated in the Py-S2 Mut section (Figures 2C and 4B). However, centromeric SUMO2/3 granularity was still significantly increased in ΔTopoIIα cells with ICRF-193 (SUMO2/3 granularity quantification), indicating that other proteins located at the centromere were still modified under ICRF-193 treatment and these were not significantly contributing to PICH enrichment at centromeres. These results suggest that TopoIIα SUMOylation critically contributes to the centromeric enrichment of PICH on chromosomes under ICRF-193 treatment.

Loss of PICH leads to enrichment of chromosomal SUMOylated proteins

So far, the results indicate that PICH is recruited to SUMOylated chromosomal proteins, in particular SUMOylated centromeric TopoIIα in ICRF-193–treated cells. Because the ability of PICH to interact with SUMO via its SUMO-interacting motifs is required for proper chromosome segregation, we wished to determine whether PICH is required for regulating the association of SUMOylated proteins with chromosomes. To examine this, mAID-PICH cells were generated as described above for TopoIIα. After auxin was added to the cells for 6 h, PICH levels became undetectable by Western blot and immunofluorescence analysis (Supplemental Figure S5, A–E). To deplete PICH, auxin was added to the cells after release from a thymidine block, and then mitotic cells were collected by mitotic shake off. Isolated chromosomes were then subjected to Western blot analysis. Intriguingly, compared with cells treated with DMSO (control) and no auxin, ΔPICH control cells showed an increase in SUMOylated TopoIIα, revealed by the appearance of a second upshifted band marked by an asterisk (Figure 6A). In addition, Western blots showed that cells depleted for PICH and treated with DMSO (control) showed an overall increase in SUMO2/3 signal, although not statistically significant (Figure 6A, compare lanes 1 and 3). Immunofluorescence staining further supported this novel role of PICH on SUMOylated chromosomal proteins. ΔPICH cells stained for SUMO2/3 showed a significant increase in mean chromosomal intensity as compared with control cells with intact PICH (Figure 6B, quantification). In ΔPICH cells treated with ICRF-193, the intensity of chromosomal SUMO2/3 was not significantly higher than in cells with intact PICH. In addition, centromeric SUMO2/3 granularity showed that depletion of PICH increased centromeric SUMO2/3 foci intensity on chromosomes, which was further increased under ICRF-193 treatment (Figure 6B, granularity quantification). This suggests that PICH acts on SUMO2/3-modified chromosomal proteins to attenuate their binding to chromosomes, including centromeric enrichment in both control and ICRF-193–treated conditions. The discrepancy between Western blot and immunofluorescence image quantification suggests that these stabilized SUMO2/3-modified proteins on chromosomes after PICH depletion are rather weakly associated with chromosomes and likely dissociate during biochemical isolation similar to Py-S2 Mut–expressing cells. As compared with control cells (DMSO) with intact PICH, ΔPICH cells treated with DMSO showed a significant increase of mean TopoIIα chromosomal intensity and granularity (Figure 6C, quantification). ICRF-193 treatment did not further increase total TopoIIα intensity. The granularity of centromeric TopoIIα in ΔPICH cells after DMSO and ICRF-193 treatment was lower than that of cells with PICH. Together, the results suggest that PICH functions in the regulation and proper localization of SUMOylated chromosomal proteins and TopoIIα at the centromere.

ATP-dependent translocase activity and SIMs of PICH are required for regulating SUMOylated chromosomal proteins

To identify which function of PICH is required for the chromosomal distribution of SUMOylated proteins, we created a PICH-replacement cell line by combining mAID-mediated PICH depletion and inducible expression of exogenous PICH mutants. The mAID-PICH cells had CRISPR/Cas9-targeted integration of either Tet-inducible WT PICH-mCherry, an ATPase dead mutant (K128A-mCherry), or a non–SUMO-interacting form of PICH (Δ3SIM-mCherry) into the CCRI5 safe harbor locus (Papapetrou and Schambach, 2016). After clonal isolation and validation (Supplemental Figure S6, A–C), PICH-mCherry expression was tested in asynchronous cells by treatment with auxin and doxycycline for 14 h, and the whole cell lysates were used for Western blot analysis. Although the expression level of the exogenous proteins was variable, we were able to replace endogenous PICH with exogenous PICH (Figure 7A). We did observe variation of mCherry expression within each clonal isolate (Supplemental Figure S6D), and this may explain the variation in expression levels observed in Western blot analysis. The PICH replacement for mitotic cell analysis was achieved by incubating cells with auxin or auxin and doxycycline for 22 h before mitotic shake off. The mitotic cells were treated with DMSO (control) or ICRF-193, and then mitotic chromosomes were isolated. Western blot analysis was performed to determine how translocase activity and SIMs contribute to PICH-binding mitotic chromosomes (Figure 7B). The PICH WT-mCherry was observed to have a response to ICRF-193 similar to that of endogenous PICH, showing increased binding with ICRF-193 treatment. The K128A mutant also showed increased binding after ICRF-193 treatment. In contrast, the Δ3SIM mutant could not bind to chromosomes. This suggests that PICH SIMs are required for chromosomal association, which is consistent with our previous observations (Sridharan and Azuma, 2016).

To further examine how the PICH mutants affect chromosomal localization of SUMO2/3 and TopoIIα, immunofluorescence analysis of prometaphase cells was performed. PICH WT-mCherry showed the same staining patterns as endogenous PICH, and its response to ICRF-193 was similar to that in Figure 1. Both SUMO2/3 and TopoIIα total intensities increased in response to ICRF-193 treatment. Distribution analysis also showed increased granularity for both SUMO2/3 and TopoIIα under ICRF-193 treatment. But TopoIIα granularity was not statistically significantly increased in PICH WT-mCherry–replaced cells. This discrepancy likely originates from the limitations of the system in terms of control of expression levels of transgenes. Loss of PICH showed increased TopoIIα granularity (Figure 6C); thus insufficient expression of PICH WT-mCherry could affect the PICH ability to form foci.
Despite the limitation of expression-level control, the results support the feasibility of this system to assess PICH function by replacing the endogenous PICH with mutants.

When the K128A mutant, which cannot translocate DNA, was expressed, strong mCherry foci were observed on the chromosomes regardless of ICRF-193 treatment. Importantly, these foci...
overlap with SUMO2/3 foci in all K128A cells observed, regardless of treatment (Figure 8A). SUMO2/3 mean intensity was observed to significantly increase after ICRF-193 treatment (Figure 8A, left, quantification), suggesting that the response to ICRF-193 remains intact. Intriguingly, the SUMO2/3 granularity was not statistically different between control and ICRF-193–treated cells, presumably due to increased granularity caused by expression of the PICH K128A compared with PICH WT (Figure 7C). This suggests that the PICH K128A mutant interacts with SUMOylated targets but due to its inability to translocate DNA, PICH K128A remains stably associated with the chromosomes where the SUMOylated proteins are located. TopoII\(\alpha\) signals in PICH K128A cells showed a clear increase in intensity compared with PICH WT and more diffuse localization on chromosomes. Both total chromosomal intensity and granularity of TopoII\(\alpha\) showed no significant increase with ICRF-193 treatment (Figure 8A, right, quantifications). This is consistent with \(\Delta\)PICH conditions (Figure 6C). This suggests that the increased chromosomal association of TopoII\(\alpha\) after loss of PICH is responsible for the ICRF-193 ineffectiveness in this analysis. Notably, PICH foci did not show any apparent overlap with TopoII\(\alpha\) foci after ICRF-193 treatment, as was observed in parental DLD-1 cells (Figure 1B) and PICH WT–replaced cells (Figure 7C). This could indicate that stable binding to other SUMO2/3-modified chromosomal proteins hinders PICH from interacting with foci-forming TopoII\(\alpha\) after ICRF-193 treatment. As observed by Western blot analysis, the PICH d3SIM-mCherry mutant did not show any chromosomal signal, but rather a diffuse signal was observed throughout the cell. SUMO2/3 signals were observed on chromosomes in the PICH d3SIM-expressing cells. The total intensity of SUMO2/3 signal on chromosomes was not significantly increased with ICRF-193 treatment. This might originate from the already increased level of SUMO2/3-modified proteins on chromosomes in the PICH
d3SIM (Figure 8B, quantification) compared with PICH WT (Figure 7C). However, the granularity of SUMO2/3 showed that the ICRF-193–dependent increase of SUMO2/3 foci was retained. Similar to PICH K128A, the Topollα signal on the chromosome was increased compared with PICH WT, and Topollα localization on chromosomes in PICH d3SIM-expressing cells was more diffused. No significant difference was observed after ICRF-193 treatment in terms of total chromosomal intensity (Figure 8B, quantification). Again, consistent with ΔPICH (Figure 6C), ICRF-193 treatment did not result in an increase in Topollα foci formation in the PICH d3SIM-replaced cells. This suggests that the SIM-dependent chromosomal association of PICH, or regulation of SUMO2/3-modified chromosomal proteins by PICH, is required for proper distribution of Topollα on mitotic chromosomes. Taking the results together, using its SIMs and translocase activity, PICH attenuates the association of SUMO2/3-modified proteins with chromosomes. Thus, these activities of PICH affect chromosome organization, including Topollα association with chromosomes.

FIGURE 8: Translocase activity and SIMs on PICH are necessary for redistribution of SUMOylated proteins and SUMOylated Topollα on mitotic chromosomes. (A) K128A PICH mCherry mitotic cells were fixed and stained with antibodies against SUMO2/3 (green), Topollα (green), and mCherry (red). DNA was stained with DAPI (blue). Scale bar = 11 μm. The mean intensities of SUMO2/3 signals are shown. p values for comparison of all values from three experiments were calculated using a Student’s t test ANOVA; ns: not significant; **: p < 0.01; ***: p < 0.001. (B) d3SIM PICH mCherry mitotic cells were fixed and stained with antibodies against: SUMO2/3 (green), Topollα (green), and mCherry (red). DNA was stained with DAPI (blue). Scale bar = 11 μm. The mean intensities of Topollα signals are shown. p values for comparison of all values from three experiments were calculated using a Student’s t test ANOVA; ns: not significant; ***: p < 0.001.
PICH attenuates decatenation activity of SUMOylated TopoII dependent on its SIMs

The cell-based assays suggest that PICH is required for proper organization of chromosomal SUMOylated proteins and SUMOylated TopoII is one of the targets. To determine the potential role of PICH interaction with SUMOylated TopoII, we performed an in vitro DNA decatenation assay to compare the effect of PICH on non-SUMOylated and SUMOylated TopoII (Figure 9A). Using the same conditions established in our previous study, recombinant *Xenopus laevis* TopoII was SUMOylated in vitro, and then its DNA decatenation activity was analyzed by using catedenated kDNA as the substrate (Ryu et al., 2010b). The decatenation activity was measured by calculating the percentage of decatenated kinetoplast DNA (kDNA) separated by gel electrophoresis. On average, 70% of kDNA is decatenated at the 5- and 10-min time points when non-SUMOylated TopoII is present in the reaction (Figure 9B, lanes marked by i). As we have previously shown, the decatenation activity of SUMOylated TopoII was reduced compared with non-SUMOylated TopoII (Figure 9B, lanes marked by ii). Importantly, when we added PICH to each of the reactions at concentrations equimolar to that of TopoII (200 nM), the decatenation activity of SUMOylated TopoII was further attenuated (Figure 9, B, marked by iii, and C). The reduction in decatenation activity of SUMOylated TopoII was statistically significant at both the 5- and 10-min time points (Figure 9C, light gray gray bars). A dose-dependent effect of PICH on SUMOylated TopoII decatenation activity was observed, but that was not the case for non-SUMOylated TopoII. The concentration of TopoII in the reaction was 200 nM, and PICH significantly reduced decatenation activity of SUMOylated TopoII ranging from 200 nM (equimolar) up to 400 nM (Figure 9, D and E). Only SUMOylated TopoII was inhibited by PICH dose-dependently, which is distinct from the PICH/non-SUMOylated TopoII interaction. PICH binding to TopoII has been shown to increase TopoII activity in vitro (Nielsen et al., 2015). In our assay, however, that increase was not clearly detected. This discrepancy might originate from the assay conditions, such as the existence of SUMO in the reaction. It is possible that excess SUMO in the reaction interacts with PICH and that affects PICH interaction with unmodified TopoII.

To determine which activity of PICH is required for inhibiting SUMOylated TopoII decatenation activity, we utilized the PICH mutants that have defects in the SUMO-binding ability (PICH-d3SIM) or in translocase activity (PICH-K128A) (Figure 10A) (Sridharan and Azuma, 2016). If PICH/SUMO interaction is critical for inhibiting the decatenation activity of SUMOylated TopoII, the PICH-d3SIM mutant would lose its inhibitory function. In addition, we also expect that the PICH translocase activity-deficient (PICH-K128A) mutant would lose its inhibitory function on SUMOylated TopoII, if the translocase-deficient mutant could not remove SUMOylated TopoII from kDNA. Supporting our hypothesis, PICH-d3SIM lost its inhibitory function and SUMOylated TopoII decatenation activity returned to levels similar to that of no PICH addition (Figure 10C, compare ST to ST + PICH d3SIM). This suggests that direct SUMO/SIM interactions between PICH and SUMOylated TopoII have a key role in this inhibition. In contrast, the translocase-deficient PICH mutant was able to attenuate SUMOylated TopoII...
and regulates their distribution on mitotic chromosomes. Increasing SUMOylation, whether by a modulating deSUMOylation enzyme or using a specific inhibitor of Topolll, led to the enrichment of PICH foci on mitotic chromosomes. Mutant PICH cell lines demonstrated that both DNA translocase activity and SUMO-binding abilities of PICH are required for regulating the distribution of SUMO2/3-modified proteins on chromosomes: without these activities of PICH, increased SUMO2/3 foci formation was observed.

PICH targets and regulates chromosomal SUMOylated proteins using its SUMO-binding ability and translocase activity

SUMOylation has been shown to play a role in complex assembly by mediating SUMO/SIM interactions (Lin et al., 2006; Guzzo et al., 2012; Pelisch et al., 2017; Matmati et al., 2018). It has been demonstrated that numerous proteins are SUMOylated on mitotic chromosomes (Schimmel et al., 2014; Cubenas-Potts et al., 2015; Huang et al., 2016). Proper regulation of SUMOylation on chromosomal proteins is apparently key to promoting faithful chromosome segregation, as revealed by modulating enzymes that control SUMOylation (Hari et al., 2001; Diaz-Martinez et al., 2006; Cubenas-Potts et al., 2013; Pelisch et al., 2014). Our current study demonstrates that SUMOylated chromosomal proteins are targeted by PICH through itsSIMs. Increased SUMO2/3 modification, either by treating cells with ICRF-193 (Figure 1) or by targeting a dominant negative deSUMOylation enzyme mutant to chromosomes (Figure 4), promotes focal enrichment of PICH and SUMO2/3 on mitotic chromosomes. This suggests that PICH efficiently targets SUMOylated chromosomal proteins, including Topolll, and either redistributes them away from foci or evicts them from chromosomes. Given the fact that PICH can interact with SUMO moieties (Sridharan et al., 2015) using its three SIMs, this enrichment of PICH foci and SUMO2/3 foci suggests that PICH can target multiple SUMOylated chromosomal proteins. More importantly, the translocase-deficient mutant of PICH showed an enrichment of SUMO2/3 and PICH foci that overlapped on chromosomes without ICRF-193 treatment (Figure 8A). This suggests that loss of translocase activity of PICH stabilizes SUMOylated protein(s) forming a stable complex on chromosomes. The PICH primary structure suggests that it acts as a nucleosome-remodeling enzyme. However, PICH has not been shown to have robust nucleosome-remodeling activity toward nucleosomes composed of canonical histones (Ke et al., 2011). Our observations imply that PICH could utilize its translocase activity for remodeling chromosomal SUMOylated proteins. Because both translocase-deficient PICH and SIM-deficient PICH mutants showed defective Topolll localization, specifically that Topolll signals became diffuse and lost their response to ICRF-193 treatment (shown in Figure 8), the remodeling of SUMO2/3-modified chromosomal proteins by

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**DISCUSSION**

We previously demonstrated that both DNA translocase activity and SUMO-interacting ability are required for the essential functions of PICH in proper chromosome segregation (Sridharan and Azuma, 2016). The results presented in this report provide the link between these two functions of PICH during mitosis. Collectively, the results indicate that PICH interacts with SUMOylated chromosomal proteins and indicates that PICH recognizes the SUMO moieties on Topolllα through its SIM interactions (Lin et al., 2001). Our observations imply that PICH could utilize its translocase activity for remodeling chromosomal SUMOylated proteins. Because both translocase-deficient PICH and SIM-deficient PICH mutants showed defective Topolll localization, specifically that Topolll signals became diffuse and lost their response to ICRF-193 treatment (shown in Figure 8), the remodeling of SUMO2/3-modified chromosomal proteins by
SUMOylated Topollα could explain our results that show Topollα mislocalization on chromosomes in PICH mutant–replaced cells. Further study for identification of which SUMOylated chromosomal proteins are targeted by PICH and a more detailed analysis of the chromosome structures in PICH-replaced cells will address our hypothesis.

SUMOylated Topollα is a target of PICH
ICRF-193 inhibits Topollα SPR by stalling Topollα in a closed-clamp conformation that holds two decatenated DNA strands within Topollα (Roca et al., 1994; Morris et al., 2000). Therefore, increased Topollα foci after ICRF-193 could represent an SPR active population of Topollα stalled on genomic DNA. Our results demonstrated that these stalled Topollα enzymes subsequently undergo SUMOylation and are then recognized by PICH (Figures 3 and 4), mainly observed at the centromere (Figures 5 and 6). These findings provide a novel mechanistic understanding of the interaction between SUMOylated Topollα and PICH and provide insight into why PICH knockout cells were found to be sensitive to ICRF-193. Perhaps more importantly, depletion of PICH led to increased abundance of Topollα on mitotic chromosomes even in the absence of ICRF-193 treatment. Therefore, the data provide evidence that PICH has a specific function under physiological conditions, to regulate the abundance of Topollα on chromosomes in mitosis. Though speculative, it may be the case that PICH evicts Topollα that becomes stalled during the SPR even in the absence of a Topollα catalytic inhibitor.

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**FIGURE 11:** Model for demonstrating the role of PICH in regulating chromosomal SUMOylated proteins for proper chromosome segregation. SUMOylation plays a critical role in chromosome regulation and mitotic timing; this is due in part to regulating the activity and mediating the binding of critical proteins. During mitosis, proteins become SUMOylated and PICH recognizes and binds these proteins using its three SIMs and then using its translocase activity redistributes or removes SUMOylated proteins from the chromosomes, and this enables proper chromosome segregation.

PICH is required for proper organization of chromosomes. This is consistent with abnormal Topollα localization on chromosomes in PICH knockdown/knockout cells, as has been shown in previous studies (Biebricher et al., 2013; Nielsen et al., 2015). Loss of PICH results in chromosome bridge formation in anaphase (Nielsen et al., 2015), and both translocase activity and SIMs are essential for preventing chromosome bridge formation (Sridharan and Azuma, 2016). SUMOylated protein remodeling by PICH may affect the overall chromosome structure and organization. PICH-d3SIM could not localize to chromosomes, but endogenous PICH could still associate with mitotic chromosomes without Topollα (Figure 5). In addition to Topollα, other important chromosomal structural proteins are SUMOylated, including the SMC complex proteins (Stephan et al., 2011; Wagner et al., 2019; Xu and Yanagida, 2019), which are likely targets of PICH function. Therefore, chromosomal structural anomalies in PICH mutants could explain the anaphase defects described above. Alternatively, recent studies indicate that the translocase activity of PICH can be used to control the supercoiling status of DNA together with topoisomerase IIα (TOP3A) (Bizard et al., 2019). This PICH/TOP3A function could control Topollα activity/binding to genomic DNA by providing the preferable DNA topology for Topollα. If the SIM-deficient mutant has a defect in this PICH/TOP3A function, this DNA topology–dependent regulation of Topollα could help resolve tangled DNA during anaphase (Nielsen et al., 2015). In addition, a recent study of PICH/TOP3A in controlling the supercoiling status of DNA for optimal Topollα activity supports the specific role of PICH in controlling Topollα activity on genomic DNA (Bizard et al., 2019). These mechanisms certainly help Topollα perform the SPR to prevent or resolve the formation of UBFs, which are created by tangled genomic DNA. In this study we presented evidence that depleting Topollα abrogates the enrichment of PICH foci at centromeres even in the presence of ICRF-193 and increased SUMO2/3 foci at centromeres (Figure 5), suggesting that PICH specifically targets SUMOylated Topollα in ICRF-193-treated cells. Further, PICH depletion increased chromosome-associated SUMOylated Topollα (Figure 6A), suggesting PICH attenuates SUMOylated Topollα association with chromosomes. The SIM-dependent interaction of PICH with SUMOylated Topollα was shown in the in vitro assays. PICH binding with SUMOylated Topollα has different consequences, that is, inhibition of decatenation activity (Figure 9). The inhibition of activity requires SIMs, suggesting that direct interaction of PICH and SUMO moieties on Topollα is critical (Figure 10). The mechanism of how both WT PICH and translocase-deficient mutants similarly inhibit the decatenation activity of SUMOylated Topollα is currently unclear. If we apply our model of PICH as a
SUMOylated protein remodeler to this context, the mechanism of inhibition might be by removing SUMOylated TopolⅡⅡ from DNA. If that is the case, the translocase-deficient mutant could inhibit decatenation activity by forming a stable complex with SUMOylated TopolⅡⅡ on DNA, which can be predicted by the observation of stabilized SUMOylated protein on chromosomes in PICH-K128A-replaced cells. Further analysis of the complex formation of PICH and SUMOylated TopolⅡⅡ in vitro or in cells is our next goal in elucidating the mechanism of this inhibition.

**Broader implications of the novel function of PICH as a SUMOylated protein remodeler**

Our data suggest a novel function of PICH on mitotic chromosomes in remodeling SUMO2/3-modified chromosomal proteins for promoting faithful chromosome segregation. The defect in PICH SIM or translocase activity mutants causes structural abnormalities revealed by aberrant localization of TopolⅡⅡ and lack of response to ICRF-193. The remodeling activity of PICH toward SUMOylated chromosomal proteins is predicted to control their mobility on chromosomes. This can be tested by live-imaging analysis after tagging endogenous SUMO2 or 3 and visualizing localization in cells. Identification of SUMO2/3-modified proteins targeted by PICH will be critical in advancing our understanding of how temporal SUMO2/3 modification on mitotic chromosomes contributes to chromosome segregation. We expect that knowing these targets will reveal how and why loss of PICH and misregulation of mitotic SUMOylation affect chromosome structure and faithful segregation of chromosomes. One clear target of PICH is SUMOylated TopolⅡⅡ after ICRF-193 treatment. Recently, we demonstrated that ICRF-193 treatment resulted in a mitotic arrest in cells that requires SUMOylated TopolⅡⅡ and subsequent Aurora B activation (Pandey et al., 2020). It is possible that PICH can control this stalled TopolⅡⅡ-dependent mitotic checkpoint by evicting SUMOylated TopolⅡⅡ from chromosomes. This can be tested using PICH depletion or replacement cell lines as well as modulating PICH activity in TopolⅡⅡ-replaced cell lines with a non-SUMOylatable mutant.

**MATERIALS AND METHODS**

**Plasmids, constructs, and site-directed mutagenesis**

The Py·S2 fusion DNA construct of human PIASy-NTD (amino acids 1–135) and SENP2-CD (amino acids 363–589) was created by the fusion PCR method using a Glycine-Alanine linker between the two fragments. Then the Py·S2 fusion DNA fragment was subcloned into a recombinant expression PET28a plasmid at the BamHI/Xhol sites. To generate the Py·S2 Mut fusion DNA construct, substitution of cysteine to alanine at 548 in Py·S2 was introduced using a site-directed mutagenesis QuikChangeII kit (Agilent) according to the manufacturer's instructions.

**Recombinant protein expression and purification, and preparation of antibodies**

Recombinant TopolⅡⅡ and PICH proteins were prepared as previously described (Ryu et al., 2010b; Sridharan and Azuma, 2016). In brief, the pPIC 3.5K plasmids carrying TopolⅡⅡ or PICH cDNA fused with calmodulin-binding protein tag were transformed into the GS115 strain of Pichia pastoris yeast and expressed by following the manufacturer's instructions (Thermo/Fisher). Yeast cells expressing recombinant proteins were frozen and ground with a coffee grinder that contains dry ice, suspended with lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 2 mM CaCl₂, 1 mM MgCl₂, 0.1% Triton X-100, 5% glycerol, 1 mM dithiothreitol [DTT], complete EDTA-free Protease inhibitor tablet [Roche], and 10 mM phenylmethylsulfonyl fluoride). The lysed samples were centrifuged at 25,000 × g for 40 min. To capture the CBP-tagged proteins, the supernatant was mixed with calmodulin-Sepharose resin (GE Healthcare) for 90 min at 4°C. The resin was then washed with lysis buffer, and proteins were eluted with buffer containing 10 mM ethylene glycol-bis(β-aminoethyl ether)-N,N,N′,N′-tetraacetic acid. In the case of PICH, the elution was concentrated by centrifugal concentrator (Amicon ultra with a 100-kDa-molecular-weight cutoff). In the case of TopolⅡⅡ, the elution was further purified by Hi-trap Q anion-exchange chromatography (GE Healthcare). Recombinant Py·S2 proteins fused to the hexa-histidine tag were expressed in Rosetta2 (DE3) (EMD Millipore/Novagen) and purified with hexa-histidine affinity resin (Talon beads from Takara/Clontech). Fractions by imidazole-elution
were subjected to Hi-trap SP cation-exchange chromatography. The peak fractions were pooled and then concentrated by centrifugal concentrator (Amicon ultra with a 30-kDa-molecular-weight cutoff). The E1 complex (Aos1/Uba2 heterodimer), PIASy, Ubc9, dnUbc9, and SUMO paralogues were expressed in Rosetta2(DE3) and purified as described previously (Ryu et al., 2010a).

To generate the antibody for human PICH, the 3’end (coding for amino acids 947–1250) was amplified from PICH cDNA by PCR. The amplified fragment was subcloned into the pET28a vector (EMD Millipore/Novagen), and then the sequence was verified by DNA sequencing. The recombinant protein was expressed in Rosetta2(DE3) strain (EMD Millipore/Novagen). Expressed protein was found in the inclusion body; thus the proteins were solubilized by 8 M urea-containing buffer (20 mM HEPES, pH 7.8, 300 mM NaCl, 1 mM MgCl₂, 0.5 mM tris(2-carboxyethyl)phosphine). The solubilized protein was purified by Talon-resin (Clontech/Takara) using the hexa-histidine-tag fused at the N-terminus of the protein. The purified protein was separated by SDS–PAGE, and protein was excised after InstantBlue (Sigma-Aldrich) staining. The gel slice was used as an antigen, and immunization of rabbits was done by Pacific Immunology, USA. To generate the primary antibody for human TopoIIα, the 3’end of TopoIIα (coding for amino acids 1359–1589) was amplified from TopoIIα cDNA by PCR. The amplified fragment was subcloned into pET28a and pGEX-4T vectors (GE Healthcare), and then the sequence was verified by DNA sequencing. The recombinant protein was expressed in Rosetta2(DE3). The expressed protein was purified using hexa-histidine-tag and GST-tag by Talon-resin (Clontech/Takara) or glutathione-Sepharose (GE Healthcare) following the manufacturer’s protocol. The purified proteins were further separated by cation-exchange column. Purified hexa-histidine-tagged TopoIIα protein as used as an antigen, and immunization of rabbits was done by Pacific Immunology, USA. For both PICH and TopoIIα antigens, antigen affinity columns were prepared by conjugating purified antigens (hexa-histidine-tagged PICH C-terminus fragment) to the NHS(N-hydroxysuccinimide)-activated Sepharose resin following the manufacturer’s protocol (GE Healthcare). The rabbit antisera were subjected to affinity purification using antigen affinity columns. Secondary antibodies used for this study and their dilution rates were, for Western blotting, goat anti-rabbit (IRDye 680RD, 1/20,000, Li-COR) and goat anti-mouse (IRDye 800CW, 1/20,000, Li-COR), and for immunofluorescence staining, goat anti-mouse immunoglobulin G [IgG] Alexa Fluor 568 (#A11031, 1:500, Invitrogen), goat anti-rabbit IgG Alexa Fluor 568 (#A11036, 1:500, Thermo/Fisher), goat anti-rabbit IgG Alexa Fluor 488 (#A11041, 1:500, Thermo/Fisher), and goat anti-guinea pig IgG Alexa Fluor 568 (#A21450, 1:500, Thermo/Fisher). Unless otherwise stated, all chemicals were obtained from Sigma-Aldrich.

In vitro SUMOylation assays and decatenation assays
The SUMOylation reactions were performed in the reaction buffer (20 mM HEPES, pH 7.8, 100 mM NaCl, 5 mM MgCl₂, 0.05% Tween 20, 5% glycerol, 2.5 mM ATP, and 1 mM DTT) by adding 15 nM E1, 15 nM Ubc9, 45 nM PIASy, 500 nM T7-tagged Topolox, and 5 µM SUMO2-GG. For the non-SUMOylated Topolox control, 5 µM SUMO2-G mutant was used instead of SUMO2-GG. After the reaction with the incubation for 1 h at 25°C, it was stopped with the addition of EDTA at a final concentration of 10 mM. For the analysis of the SUMOylation profile of Topolox 3X SDS–PAGE sample buffer was added to the reaction, and the samples were resolved on 8–16% Tris–HCl gradient gels (#XP08165BOX, Thermo/Fisher) by SDS–PAGE and then analyzed by Western blotting with horseradish peroxidase–conjugated anti-T7 monoclonal antibody (#T3699, EMD Millipore/Novagen).

Decatenation assays were performed in the decatenation buffer (50 mM Tris–HCl, pH 8.0, 120 mM NaCl, 5 mM MgCl₂, 0.5 mM DTT, 30 µg bovine serum albumin [BSA]/ml, and 2 mM ATP) with SUMOylated Topolox and non-SUMOylated Topolox and with 6.2 ng/µl of kDNA (TopoGEN). The reaction was performed at 25°C with the conditions indicated in each of the figures. The reactions were stopped by adding one-third volume of 6X DNA dye (30% glycerol, 0.1% SDS, 10 mM EDTA, and 0.2 µg/µl bromophenol blue). The samples were loaded on a 1% agarose gel containing SYBR Safe DNA Gel stain (#S33102, Invitrogen) with 1 kb ladder (#N32325, NEB) and electrophoresed at 100 V in TAE buffer (Tris-acetate-EDTA) until the marker dye reached the middle of the gel. The amount of kDNA remaining in the wells was measured using ImageStudio, and the percentage of decatenated DNA was calculated as (intensity of initial kDNA [at 0 min incubation] – intensity of remaining catenated DNA/intensity of initial kDNA). Obtained percentages of catenated DNA were plotted and analyzed for the statistics by using GraphPad Prism 8 Software.

Cell culture, transfection, and colony isolation
Targeted insertion using the CRISPR/Cas9 system was used for all integration of exogenous sequences into the genome. DLD-1 cells were transfected with guide plasmids and donor plasmid using ViaFect (#E4981, Promega) on 3.5 cm dishes. The cells were split and replated on 10 cm dishes at ~20% confluency; 2 d later, the cells were subjected to a selection process by being maintained in the medium in the presence of a desired selection reagent (1 µg/ml blasticidin [#ant-bl, Invivogen], 400 µg/ml zeocin [#ant-zn, Invivogen], 200 µg/ml hygromycin B gold [#ant-hg, Invivogen]). The cells were cultured for 10–14 d with a selection medium, the colonies were isolated and grown in 48-well plates, and Western blotting and genomic DNA samples were prepared to verify the insertion of the transgene. Specifically, for the Western blotting analysis, the cells were pelleted and 1X SDS–PAGE sample buffer was added and boiled/vortexed. Samples were separated on an 8–16% gel and then blocked with casein and probed using the antibody described in each figure legend. Signals were acquired using the Li-COR Odyssey Fc imager. To perform genomic PCR, the cells were pelleted, and genomic DNA was extracted using lysis buffer (100 mM Tris–HCl, pH 8.0, 200 mM NaCl, 5 mM EDTA, 1% SDS, and 0.6 mg/ml protease K [#RP81075, NEB]) and purified by ethanol precipitation followed by resuspension with TE buffer containing 50 µg/ml RNase A (#EN0531, ThermoFisher). Primers used for confirming the proper integrations are listed in the Supplemental Information.

To establish AID cell lines, as an initial step, the Oryza sativa E3 ligase (OsTIR1) gene was inserted into the 3’ end of a housekeeping gene, RCC1, using the CRISPR/Cas9 system in the DLD-1 cell line. The RCC1 locus was an appropriate locus to accomplish the modest but sufficient expression level of the OsTIR1 protein so that it would not induce a nonspecific degradation without the addition of auxin (Supplemental Figure S3) (Yau et al., 2020). We then introduced DNA coding for AID-3xFlag tag into the Topolox or PICH locus using CRISPR/Cas9 editing into the OsTIR1-expressing parental line (Supplemental Figures S4 and S5). The isolated candidate clones were subjected to genomic PCR and Western blotting analysis to validate integration of the transgene. Once clones were established and the transgene integration was validated, the depletion of the protein in the auxin-treated cells was confirmed by Western blotting and immunostaining.
Introducing DNA encoding Tet-inducible PICH mCherry into the CCR5 locus or inducible Py-S2 into hH11 was done by CRISPR/Cas9 editing into the desired locus (Supplemental Figures S2 and S6). The OsTIR1-expressing mAID PICH parental cell line was used for introduction of the PICH mCherry mutants targeted to the CCR5 locus. The isolated candidate clones were subjected to genomic PCR and Western blotting analysis to validate integration of the transgene. Once clones were established and the transgene integration was validated, the expression of the transgenes was confirmed by the addition of doxycycline.

**Xenopus egg extract assay for mitotic chromosomal SUMOylation analysis**

Low-speed cytostatic factor (CSF)-arrested XEEs and demembranated sperm nuclei were prepared following standard protocols (Murray, 1991; Powers et al., 2001). To prepare the mitotic replicated chromosome, CSF extracts were driven into interphase by adding 0.6 M CaCl2. Demembranated sperm nuclei were added to interphase extract at 4000 sperm nuclei/μl and then incubated for ~60 min to complete DNA replication confirmed by the morphology of nuclei. Then an equal volume of CSF XEE was added to the reactions to induce mitosis. To confirm the activities of Py-S2 proteins on mitotic SUMOylation, the Py-S2 proteins or dnUbC9 were added to XEEs at final concentrations of 30 nM or 5 μM, respectively, at the onset of mitosis induction. After mitotic chromosome formation was confirmed by microscopic analysis of condensed mitotic chromosomes, chromosomes were isolated by centrifugation using a 40% glycerol cushion as previously described (Yoshida et al., 2016), and then the isolated mitotic chromosomes were boiled in SDS–PAGE sample buffer. Samples were resolved on 8–16% gradient gels and subjected to Western blotting with the indicated antibodies. Signals were acquired using a Li-COR Odyssey Fc machine.

**Preparation of mitotic cells and chromosome isolation**

DLD-1 cells were grown in McCoy’s 5A 1× L-glutamine 10% fetal bovine serum (FBS) media for no more than 10 passages. To analyze mitotic chromosomes, cells were synchronized by the thymidine/blocking arrest protocol. In brief, cells were arrested with 2 mM thymidine for 17 h, released from the thymidine block by washing three times with 1X PBS containing 10 mM Tris-HCl to quench PFA. Following the fixation, the cells were permeabilized using 100% ice cold methanol in a −20°C freezer for 5 min. Cells were then blocked using 2.5% hydroxylated gelatin in 30 min at room temperature. Following washing, the cells were stained with primary antibodies for 1 h at room temperature, washed three times with 1X PBS containing 0.1% Tween 20, and incubated with secondary antibodies for 1 h at room temperature. Following secondary incubation, cells were washed three times with 1X PBS-Tween20 and mounted onto slide glass using VECTASHIELD Antifade Mounting Medium with 4′,6-diamido-2-phenylindole (#H-1200, Vector Laborotory) and sealed with nail polish. Images were acquired using an UltraView VoX spinning-disk confocal system (PerkinElmer) mounted on an Olympus IX71 inverted microscope. It was equipped with a software-controlled piezo-electric stage for rapid Z-axis movement. Images were controlled using a 60 × 1.42 NA planapochromatic objective (Olympus) and an ORCA ERAG camera (Hamamatsu Photonics). Solid state 405-, 488-, and 561-nm lasers were used for excitation. Fluorochrome-specific emission filters were used to prevent emission bleed through between fluorochromes. This system was controlled by Volocity software (PerkinElmer). Minimum- and maximum-intensity cutoffs (black and white levels) for each channel were chosen in Volocity before images were exported. Images are presented as extended focus. No other adjustments were made to the images. The cell images of Supplemental figures were acquired using the Plan Apo 100x/1.4 objective lens on a Nikon Ti Eclipse microscope-equipped Exi Aqua CCD camera (Q Imaging) or a Nikon TE2000-U equipped PRIME-BSI CMOS camera (Photometrics) with MetaMorph imaging software. Figures were prepared from exported images in Adobe illustrator.

**Statistical analysis of immunofluorescence images**

Quantification was performed by measuring at least five chromosomes per treatment across three individual experiments. This was done by outlining the chromosome and superimposing that drawing onto the other channels before measuring. All data provided are mean intensities. PICH foci intensity were measured in Figure 1.
by creating a 10-pixel-diameter circle on chromosome ends, superimposing circles onto PICH channels, and measuring mean intensities. The granularity measurements were performed using a 10-pixel-diameter circle to select foci to measure the intensity of foci. The mean foci intensity was then divided by the mean intensity of the total chromosomal signal measured by outlining chromosomes as described above. Individual chromosomes form five cells/treatment across three individual experiments were measured to obtain at least 20 data points. Mean plus SD are plotted on all graphs.

Statistical analysis

All statistical analyses were performed with either one- or two-way analysis of variance (ANOVA), followed by the appropriate post-hoc analyses using GraphPad Prism 8 software. Graphs are presented as mean with SD.

Animal use

For the XEE assay, frog eggs were collected from a mature female X. laevis, and sperm was obtained from mature male X. laevis. The animal use protocol for the X. laevis studies was approved by the University of Kansas Institutional Animal Care and Use Committee.

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