# Cohesin Function in Cohesion, Condensation, and DNA Repair Is Regulated by Wpl1p via a Common Mechanism in Saccharomyces cerevisiae

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**ABSTRACT** Cohesin tethers DNA to mediate sister chromatid cohesion, chromosome condensation, and DNA repair. How the cell regulates cohesin to perform these distinct functions remains to be elucidated. One cohesin regulator, Wpl1p, was characterized in *Saccharomyces cerevisiae* as a promoter of efficient cohesion and an inhibitor of condensation. Wpl1p is also required for resistance to DNA-damaging agents. Here, we provide evidence that Wpl1p promotes the timely repair of DNA damage induced during S-phase. Previous studies have indicated that Wpl1p destabilizes cohesin's binding to DNA by modulating the interface between the cohesin subunits Mcd1p and Smc3p. Our results suggest that Wpl1p likely modulates this interface to regulate all of cohesin's biological functions. Furthermore, we show that Wpl1p regulates cohesion and condensation through the formation of a functional complex with another cohesin-associated factor, Pds5p. In contrast, Wpl1p regulates DNA repair independently of its interaction with Pds5p. Together, these results suggest that Wpl1p regulates distinct biological functions of cohesin by Pds5p-dependent and -independent modulation of the Smc3p/Mcd1p interface.

KEYWORDS sister chromatid cohesion; condensation; DNA repair; cohesin; Wpl1

COHESIN, a member of the SMC family of protein complexes, is comprised of four subunits: Smc1p, Smc3p, Mcd1p (Scc1/Rad21), and Scc3p (SA/STAG). Cohesin mediates nuclear functions essential for both viability and the accurate transmission of genetic information, including sister chromatid cohesion, chromosome condensation, and repair of DNA damage (Onn *et al.* 2008). Cohesin is thought to perform these different functions through the spatial and temporal regulation of its ability to tether two genomic loci (Guacci *et al.* 1997; Michaelis *et al.* 1997; Hartman *et al.* 2000; Ström *et al.* 2007; Unal *et al.* 2007). Cohesin's DNA-binding and -tethering activities are regulated by factors including Eco1p (Ctf7p), Pds5p, and Wpl1p (Rad61p) (Skibbens *et al.* 1999; Tóth *et al.* 1999; Hartman *et al.* 2000; Rolef Ben-Shahar *et al.* 2008; Unal *et al.* 2008). How these regulatory factors interface with each other and with cohesin to promote its biological functions remains poorly understood.

Wpl1p was first implicated as a negative regulator of the cohesin complex, serving to inhibit both cohesion and condensation. Evidence that Wpl1p inhibits condensation stems from findings that the deletion of WPL1 (wpl1 $\Delta$ ) restores both viability and condensation to cells lacking Eco1p function (*eco1* $\Delta$ ) (Guacci and Koshland 2012), and that *wpl1* $\Delta$  cells prematurely condense their DNA (Lopez-Serra *et al.* 2013). Additionally, Wpl1p's role as an inhibitor of cohesion stems from findings that Wpl1p overexpression in human or yeast cells induces a partial cohesion loss (Gandhi *et al.* 2006; Lopez-Serra *et al.* 2013). Wpl1p is thought to inhibit cohesin function by removing it from DNA in a nonproteolytic manner (Gandhi *et al.* 2006; Kueng *et al.* 2006).

Recent biochemical studies suggest that Wpl1p destabilizes the interface between the N-terminus of Mcd1p and the base of the coiled-coil of Smc3p (Buheitel and Stemmann 2013; Beckouët *et al.* 2016). Additionally, mutating an Smc3p residue in the Smc3p/Mcd1p interface abolishes cohesin localization to centromere-proximal regions, providing *in vivo* support for a role for this interface (Gligoris *et al.* 2014). However, the biological function and regulation of

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destabilization of the Smc3p/Mcd1p interface is poorly understood.

To limit Wpl1p inhibition, cohesin is acetylated by Eco1p at two conserved lysine residues on Smc3p (K112 and K113 in the budding yeast, Saccharomyces cerevisiae) (Rolef Ben-Shahar et al. 2008; Unal et al. 2008). Additionally, Pds5p helps to preserve Smc3p acetylation during and after S-phase, suggesting a common molecular mechanism for how Pds5p and Eco1p promote cohesion (Chan et al. 2013). These functions are also thought to promote condensation, as inactivation of either factor results in dramatic defects in both cohesion and condensation (Skibbens et al. 1999; Hartman et al. 2000). Furthermore, overexpression of Pds5p suppresses mutants containing eco1-ts alleles, and vice versa, supporting the idea that Pds5p and Eco1p promote cohesin function through a common molecular mechanism (Noble et al. 2006). Taken together, these data suggest that both Eco1p and Pds5p prevent Wpl1p-mediated antagonization of cohesion and condensation.

However, the function of Wpl1p and Pds5p in regulating cohesin is more complicated. In budding yeast,  $wpl1\Delta$  cells display a partial cohesion defect, implicating Wpl1p as a positive factor required for the efficient establishment of cohesion (Rowland et al. 2009; Sutani et al. 2009; Guacci and Koshland 2012). However, the molecular differences between Wpl1p's positive and negative functions remain a mystery. Furthermore, Wpl1p and Pds5p form a complex that is capable of unloading of cohesin from DNA in vitro (Kueng et al. 2006; Murayama and Uhlmann 2015). This finding suggests that Pds5p inhibits cohesin in addition to its wellestablished role in promoting cohesin function. Consistent with this idea, in Schizosaccharomyces pombe, deletion of PDS5 suppresses a deletion of the ECO1 homolog, Eso1 (Tanaka et al. 2001). Moreover, in budding yeast, certain pds5 alleles suppress the inviability of the eco1-1 temperature-sensitive mutant, which has reduced cohesin acetylation (Rowland et al. 2009; Sutani et al. 2009). This suppression suggests that these pds5 mutations inactivate an inhibitory activity of Pds5p. Together, these results suggest that Wpl1p and Pds5p can act both positively and negatively to regulate cohesin functions.

The complex regulation of Wpl1p on cohesin function raises important questions that we address in this study. First, are there additional roles of Wpl1p in regulating cohesin function? Does Wpl1p regulate all cohesin's biological functions through a common molecular mechanism? Finally, is Wpl1p's ability to form a complex with Pds5p important for any or all of Wpl1p's regulatory functions? The answers to these questions provide important new insights into cohesin regulation by Wpl1p and its interplay with Pds5p.

#### **Materials and Methods**

#### Yeast strains, media, and reagents

Yeast strains used in this study had an A364A background and their genotypes are listed in Supplemental Material, Table S1

in File S1. YPD liquid media was prepared containing 1% yeast extract, 2% peptone, and 2% dextrose, 0.01 mg/ml adenine. YPD solid media was prepared the same way as liquid media and contained 2% agar. Camptothecin (CPT) (Sigma [Sigma Chemical], St. Louis, MO) was made as a 10 mg/ml stock in dimethyl sulfoxide (DMSO) and added to a final concentration of 20 µg/ml in YPD solid or liquid media containing 25 mM pH 7.4 [4-(2-hydroxyethyl)-1-pipera-zineethanesulfonic acid (HEPES; Fisher Scientific, Fair Lawn, NJ)]. Next, 99% pure methyl methanesulfonate (MMS) (Sigma) was added to a final concentration of 0.01% in YPD solid media. MMS was diluted 1:10 in DMSO and added to YPD liquid media to a final concentration of 0.01%. Agar plates containing MMS were made within 2 days of use to prevent degradation. 5-FOA (US Biological Life Sciences, Salem, MA) was used at a final concentration of 1 g/liter in URA dropout plates supplemented with 50 mg/liter uracil (Sigma).

#### Dilution plating

Cells were grown to saturation in YPD liquid media at  $30^{\circ}$  ( $23^{\circ}$  for temperature-sensitive strains) then plated in 10-fold serial dilutions. Cells were incubated on plates at relevant temperatures or containing drugs as described. For plasmid shuffle assays, cells were grown to saturation in YPD liquid media to allow loss of covering plasmid, then plated in 10-fold serial dilutions on YPD or 5-FOA media.

#### Cohesin and condensation time course

Cells were inoculated into 5 ml YPD starter cultures and incubated overnight at 23°, then starter cultures were used to inoculate into larger volumes of YPD and grown overnight to midlog phase (~0.2–0.3 OD).  $\alpha$ -factor (Sigma) was added to midlog cultures (10<sup>-8</sup> M final) and incubated for 3 hr to arrest cells in G1. Cells were released from G1 by washing 3× in YPD containing 0.2 µg/ml Pronase E, once in YPD, and then resuspended in YPD containing 15 µg/ml nocodazole (Sigma) and incubated at 23° for 3 hr to allow cell cycle progression until arrest in mid-M. To assess cohesion through separation of LacI-GFP foci, cells were fixed for 15–30 min in 4% paraformaldehyde (w/v) and 3.4% sucrose (w/v) solution, washed, and resuspended in 0.1 KPO<sub>4</sub> 1.2 M sorbitol buffer, then stored at 4°.

For auxin treatment, time courses were performed as above, except that 3-indoleacetic acid (auxin; Sigma) was added to final concentration of 500  $\mu$ M to  $\alpha$ -factor-arrested cells from a 1 M stock solution in DMSO. Cells were then incubated for an additional 1 hr. Auxin (500  $\mu$ M) was present in all YPD washes and the releasing media containing nocodazole.

#### CPT- and MMS-treatment time course

Cells were grown, arrested in G1, and released as described above. Upon release from  $\alpha$ -factor, cells were split and resuspended into YPD containing either DMSO, 20  $\mu$ g/ml CPT and 25 mM HEPES pH 7.4, or 0.01% MMS, and incubated at 23° to allow cell cycle progression. Ninety minutes after release,

 $\alpha$ -factor was readded to cultures at 10<sup>-8</sup> M to arrest in subsequent G1. Cells were harvested every 30 min and fixed in 70% ethanol. To assess chromosome segregation, fixed cells were washed and resuspended in 1× PBS containing DAPI.

For assessment of chromosome segregation when treated with MMS or CPT in nocodazole (G2/M) cells, cells were grown, arrested, and released from  $\alpha$ -factor arrest into nocodazole in the absence of drugs, as described above. Cells were then released from nocodazole arrest by washing 3× in YPD. Cells were then split and resuspended into media containing  $\alpha$ -factor at 10<sup>-8</sup> M and either DMSO, 20 µg/ml CPT and 25 mM HEPES pH 7.4, or 0.01% MMS.

#### Fluorescence in situ hybridization (FISH)

Nocodazole-arrested cells were fixed and processed for FISH as previously described (Guacci and Koshland 1994; Guacci *et al.* 1997), except cells were stained with ProLong Gold Antifade Mountant with DAPI (Life Technologies, Carlsbad, CA) instead of propidium iodide.

# Flow cytometry

To assess DNA content, cells were fixed in 70% ethanol. Fixed cells were washed twice in 50 mM sodium citrate (pH 7.2) and then treated with RNase A [50 mM sodium citrate (pH 7.2), 0.25 mg/ml RNase A, and 1% Tween-20 (v/v)] overnight at 37°. Proteinase K was then added to a final concentration of 0.2 mg/ml and samples were incubated at 50° for 2 hr. Samples were sonicated for 30 sec or until cells were adequately disaggregated. SYBR Green DNA I dye (Life Technologies) was then added at 1:20,000 dilution and samples were run on a Guava easyCyte flow cytometer (Millipore, Billerica, MA). For each time point, 20,000 events were captured. Quantification was performed using FlowJo analysis software.

# Microscopy

Images were acquired with an Axioplan2 microscope  $[100 \times$  objective, numerical aperture (NA) 1.40; Zeiss [Carl Zeiss], Thornwood, NY] equipped with a Quantix charge-coupled device camera (Photometrics, Tucson, AZ).

# Preparation of cells for immunoprecipitation

Cells were inoculated into 5 ml starter cultures and grown overnight at 23°. Strains were then inoculated into 60 ml cultures and grown to a final OD of 0.8. Twenty ODs were then harvested, washed in  $1 \times PBS$ , spun down, liquid aspirated, and cells were flash frozen in liquid N<sub>2</sub>.

For CPT, untreated cells were grown to a final  $OD_{600}$  of 0.4. Next, 1 M HEPES pH 7.4 was added to cultures to a final concentration of 25 mM, 10 mg/ml CPT stock was added to cells to a final concentration of 20 µg/ml, and cells were incubated for 3 hr. Twenty ODs were then harvested and prepared as described above.

# Immunoprecipitation

Cell lysates were prepared by bead beating for 30 sec with 1 min rest,  $4\times$  at 4° in GNK100 buffer (100 mM KCl, 20 mM

HEPES pH 7.5, 0.2% NP40, 10% glycerol, and 2.5 mM MgCl<sub>2</sub>) containing complete mini EDTA-free protease inhibitor (Roche), 5 mM sodium butyrate, 5 mM  $\beta$ -mercaptoethanol, 1 mM PMSF, and 20 mM b-glycerophosphate. Lysates were cleared of insoluble cell debris and then incubated with anti-FLAG antibody (Sigma) and Protein A dynabeads for 1 hr at 4°. Dynabeads were then washed 4× with GNK100 buffer with additives, as described above, containing 100  $\mu$ M MG132. Samples were then run on SDS-PAGE gels and analyzed through western blot analysis.

# Preparation of bulk chromatin pellets

Cells were inoculated into 5 ml YPD starter culture and incubated overnight at 23°. Cells were inoculated from starter cultures into fresh YPD to grow overnight at 23° to midlog phase, then 15  $\mu$ g/ml nocodazole was added and cells were incubated for 3 hr to arrest cultures in mid-M-phase. Cells were processed for bulk chromatin pelleting as described in Ciosk *et al.* (2000), with the modification that spheroplasted cells were washed three times in 0.4 M sorbitol, 50 mM HEPES/KOH pH 7.5, 100 mM KCl, and 2.5 mM MgCl<sub>2</sub> before lysis.

# Data availability

Strains are available on request.

# Results

# Wpl1p is necessary to mitigate CPT- and MMS-induced cell cycle delay

Wpl1p has been implicated in regulating cohesin function in both cohesion and condensation. However, Wpl1p function in another cohesin-regulated process, DNA repair, has not been well characterized. The wpl1/rad61 mutant was originally identified in a budding yeast screen due to its weak sensitivity to ionizing radiation (Game et al. 2003). Subsequent work showed that  $wpl1\Delta$  cells have reduced viability when grown on plates containing either the topoisomerase I inhibitor CPT (10–15  $\mu$ g/ml) or the alkylating agent MMS (Sutani et al. 2009; Guacci and Koshland 2012; Guacci et al. 2015). In contrast,  $eco1\Delta$  wpl1 $\Delta$  cells exhibited severe sensitivity to both CPT and MMS (Sutani et al. 2009; Guacci and Koshland 2012). *wpl1* $\Delta$  cells have a modest cohesion defect, whereas  $eco1\Delta$  wpl1 $\Delta$  cells have a dramatic cohesion defect (Sutani et al. 2009; Guacci and Koshland 2012). The hypersensitivity of  $eco1\Delta wpl1\Delta$  cells to CPT and MMS is likely due to the severe cohesion defect impairing use of the sister chromatid as a template for DNA repair. These results clearly indicate that Wpl1p promotes resistance to DNA-damaging agents, but how it does so is unknown. To gain insight into this Wpl1p function, we analyzed how loss of Wpl1p affected viability and chromosome segregation under DNA-damaging conditions.

We first revisited the sensitivity of  $wpl1\Delta$  cells by analyzing their growth on media containing higher concentrations of

CPT than had been previously tested. We compared the growth of wild-type,  $wpl1\Delta$ , and  $eco1\Delta wpl1\Delta$  cells on media containing 20  $\mu$ g/ml CPT. As expected,  $eco1\Delta$  wpl1 $\Delta$  cells were unable to grow even after 5 days, confirming that Eco1p plays a critical role in surviving DNA damage (Figure 1A). Interestingly, after 3 days of growth on CPT,  $wpl1\Delta$  cells appeared to have much lower viability than wild-type cells. However, by 5 days,  $wpl1\Delta$  cells exhibited similar viability to wild-type cells, but formed smaller colonies, indicating slower growth (Figure 1A). Given this delayed growth phenotype, we then examined  $wpl1\Delta$  cell growth on MMScontaining plates. Consistent with our findings for CPT,  $wpl1\Delta$  cell growth on MMS was significantly delayed compared to wild-type cells, taking several days to form colonies, but the overall viability was similar to wild-type cells (Figure 1B). The similar viability but slower growth seen in  $wpl1\Delta$  cells subjected to DNA damage suggests that Wpl1p may promote efficient DNA repair.

We further characterized the kinetics of DNA-damage repair, by analyzing how CPT and MMS treatment affected cell cycle progression of wild-type and *wpl1* $\Delta$  cells. We utilized the extent and duration of a drug-induced cell cycle delay as an indirect measure of DNA damage and repair. We synchronized wild-type and *wpl1* $\Delta$  cells in G1 with  $\alpha$ -factor, and then released cells into media either containing no drugs (DMSO), CPT (20 µg/ml), or MMS (0.01%) (*Materials and Methods*). Once cells budded, we reintroduced  $\alpha$ -factor into the media to enable cells to progress through the cell cycle and then rearrest in the following G1. Aliquots of cells were collected every 30 min after G1 release and analyzed for bud morphology, DNA content, and chromosome segregation (Figure 2A).

We first compared cell cycle progression of wild-type and *wpl1* $\Delta$  cells when treated with DMSO (control) and either CPT or MMS by analyzing DNA content using flow cytometry. Progression through S-phase (transitioning from 1C to 2C) was the same for drug-treated cells and DMSO control cells (Figure S1, A and B in File S1). However, in CPT-treated *wpl1* $\Delta$  cells, the 2C DNA peak persisted longer than in either CPT-treated wild-type cells or DMSO-treated cells of either genotype (Figure S1A in File S1; 180-240 min). MMS-treated wild-type and  $wpl1\Delta$  cells both exhibited 2C DNA peaks that persisted longer than DMSO controls, but the delay was more pronounced in  $wpl1\Delta$  cells (Figure S1B in File S1; 150–240 min). These results suggest that  $wpl1\Delta$ cells delay in mitosis because of persisting DNA damage generated by either CPT or MMS, which activates the G2/M DNA-damage checkpoint.

We further characterized the CPT- and MMS-generated G2/M delays by examining bud and DNA morphologies of these cells. During an unperturbed cell cycle in yeast, DNA replication is completed when the bud is small and a single nuclear DNA mass bearing all the chromosomes is present. As the cell cycle progresses, the bud grows to medium size while mitosis quickly ensues. As chromosomes segregate, two separated DNA masses of equal size can be distinguished, one in the mother cell and one in the bud (telophase cells). If cells



**Figure 1** Wpl1p promotes proper growth on CPT and MMS media. (A)  $wp/1\Delta$  cells grow slowly on media containing camptothecin (CPT). Haploid wild-type (WT) (VG3349-1B),  $wp/1\Delta$  (VG3360-3D), and  $eco1\Delta$   $wp/1\Delta$  (VG3503 #4) cells were grown to saturation in YPD at 23°, then plated in 10-fold serial dilutions on YPD alone or containing 20 µg/ml CPT. Plates were incubated at 23° and assessed at 3 and 5 days postplating. (B)  $wp/1\Delta$  cells grow slowly on media containing MMS. Haploid WT (VG3349-1B) and  $wp/1\Delta$  (VG3360-3D) cells were grown to saturation in YPD at 23°, then plated in 10-fold serial dilution on YPD alone or containing 0.01% MMS. Plates were incubated at 23° and assessed at 3 and 5 days post plating.

stall prior to anaphase, the undivided nucleus remains at the bud neck while the bud continues to grow, giving rise to largebudded cells with unsegregated chromosomes, seen as a single DNA mass (G2/M cells) (Figure 2A; Hartwell 1974). As expected, control DMSO-treated wild-type and *wpl1* $\Delta$  cells had few large-budded G2/M cells, as most cells entered telophase when buds were midsized, consistent with no cell cycle delay (Figure 2, B and C, left panels). By 150 min post-release, most cells were in telophase (large-budded with divided nuclei). The number of telophase cells declined as cells underwent cytokinesis, and most cells were arrested in G1 by 210 min (Figure 2, B and C, left panels).

When treated with CPT, both wild-type and  $wpl1\Delta$  cultures exhibited a large increase in the amount of large-budded G2/M cells ( $\sim$ 40% of cells), and few telophase cells were seen at 120 min postrelease compared to their DMSO-treated counterparts (Figure 2B, right panels). The similar G2/M cell cycle delays of both wild-type and  $wpl1\Delta$  cells were consistent with both initially experiencing the same level of DNA damage when treated with CPT. Wild-type cells quickly overcame this delay, as seen by the high level of telophase cells at 150 and 180 min. Additionally, most wild-type cells had exited mitosis by 240 min (Figure 2B, top-right panel). In contrast, the amount of  $wpl1\Delta$  cells stalled in G2/M increased until 150 min, with a significant amount remaining stalled through 240 min (Figure 2B, bottom-right panel). Eventually, most  $wpl1\Delta$  cells entered telophase and exited mitosis, indicating that the CPT-induced damage was repaired.

Similar to CPT-treated cells, wild-type and *wpl1* $\Delta$  cells treated with MMS exhibited a G2/M delay to similar degrees, as ~50% of cells were large-budded with undivided nuclei 120 min after release (Figure 2C, right panels). However, ~50% of wild-type cells entered telophase by 150–180 min and few G2/M cells remained. In contrast, *wpl1* $\Delta$  cells were



Figure 2 Wpl1p promotes recovery from G2/M delays generated by exposure to the DNA-damageinducing agents camptothecin (CPT) and MMS. (A) Schematic of time course and analysis of cell cycle progression for untreated and CPT- or MMS-treated cells in (B and C). Wild-type (WT) (VG3349-1B) and wpl1 $\Delta$  (VG3360-3D) cells were grown to midlog phase in YPD at 23°, arrested in G1 by addition of  $\alpha$ -factor, then released from G1 into YPD (Materials and Methods). At the time of release from G1, cells were split into two aliquots: either CPT (final 20 µg/ ml) or MMS (final 0.01%) was added to one and DMSO was added to the other. Once most cells had entered S-phase (90 min after release from G1),  $\alpha$ -factor was added to ensure cells would progress through one cell cycle and rearrest in G1. Aliguots were taken every 30 min and fixed in 70% ethanol. Fixed cells were stained with DAPI to detect chromosomal DNA for scoring. Cells were scored for bud morphology (unbudded, small-medium bud, or large bud) and whether they contained a single DAPI chromosomal mass or two DAPI masses. (B) wpl1 $\Delta$  cells grown in the presence of CPT from G1 onward exhibit a prolonged mitotic delay. WT (VG3349-1B) and wpl1 $\Delta$  (VG3360-3D) cells were synchronously released from G1, as described (A), in YPD media buffered with 25 mM HEPES pH 7.4 containing DMSO alone or with 20 µg/ml CPT (Materials and Methods). Graphs show the percentage of large-budded cells with a single DNA mass (G2/ M; black) or two DNA masses (telophase; gray). (C) wpl1 $\Delta$  cells grown in the presence of MMS from G1 onward exhibit a prolonged mitotic delay. WT (VG3349-1B) and wpl1 $\Delta$  (VG3360-3D) cells were synchronously released from G1, as described in (B), except that YPD media (unbuffered) contained DMSO alone or 0.01% MMS. Cells were collected, processed, and scored as described in (A). Graphs show the percentage of large-budded cells with a single DNA mass (G2/M; black) or two DNA masses (telophase; gray).

slow to enter telophase, and the fraction of G2/M cells increased by 150 min and remained elevated through 240 min (Figure 2C, right panels). Eventually, almost all *wpl1* $\Delta$  cells completed mitosis. Together, these results show that both CPT and MMS treatment cause an initial delay in segregation in both wild-type and *wpl1* $\Delta$  cells, from which they eventually recover. However, *wpl1* $\Delta$  cells experience a prolonged delay, which most likely results from defects in the timely repair of DNA damage.

CPT-mediated damage is thought to cause double-strand breaks (DSBs) when toposiomerase I-induced single-strand nicks encounter replication machinery during S-phase (Avemann *et al.* 1988; Strumberg *et al.* 2000; Saleh-Gohari *et al.* 2005). Additionally, MMS is thought to cause DNA damage through stalling replication forks during S-phase. Thus, our results implicate Wpl1p as being important in G2/M-phase for the repair of damage generated during S-phase. To test whether the exacerbated delay observed in  $wpl1\Delta$  cells was due to the DNA damage generated during S-phase, we allowed cultures of wild-type and  $wpl1\Delta$  cells to progress synchronously through S-phase in the absence of either CPT or MMS, and arrest in G2/M by the addition of the microtubule poison nocodazole. Upon release from nocodazole, cultures were split, and either CPT or MMS was added to one aliquot while DMSO was added to the other. We then monitored progression every 30 min through mitosis and cytokinesis in either the presence or absence of CPT or MMS. α-factor was added to the cultures to prevent progression of cells beyond the ensuing G1 (Figures S2A and S3A in File S1). Upon release from nocodazole, wild-type and  $wpl1\Delta$ cells segregated their chromosomes with similar kinetics when treated with either DMSO or CPT (Figure S2, B and C in File S1). Similarly, MMS addition to nocodazole-arrested cultures failed to induce a cell cycle delay (Figure S3, B and C in File S1). Thus, the drug-induced delay in the initiation of chromosome segregation in  $wpl1\Delta$  cells required the presence of either CPT or MMS prior to M-phase. Taken together, these results suggest that Wpl1p is important for the efficient repair of multiple types of DNA damage induced during S-phase.

# The Pds5p N-terminus and the Smc3p/Mcd1p interface regulate the inhibition of condensation

Our findings, along with those of previous studies, suggest that Wpl1p's regulation of diverse cohesin functions is complicated, that Wpl1p forms a complex with Pds5p, and that Wpl1p can destabilize the interface between the N-terminus of Mcd1p and the base of the coiled-coil of Smc3p (Shintomi and Hirano 2009; Chan *et al.* 2012; Beckouët *et al.* 2016). These results provided insights into the molecular mechanisms of Wpl1p function that led us to use two approaches to parse how Wpl1p distinguishes its functions in the regulation of cohesin. The first approach was to understand the relationship between Pds5p and Wpl1p. The second utilized an *SMC3-MCD1* fusion to assess the consequences of blocking the destabilization of the Smc3p/Mcd1p interface.

The ability of  $wpl1\Delta$  to restore viability to cells lacking Eco1p function ( $eco1\Delta$ ) was previously shown (Rowland et al. 2009; Sutani et al. 2009; Feytout et al. 2011). Our further analysis showed that  $wpl1\Delta$  suppressed the condensation defect of an  $eco1\Delta$  mutant, but not the cohesion defect (Guacci and Koshland 2012). This result is consistent with the idea that abrogating Wpl1p-mediated inhibition of condensation is essential in Eco1p-deficient cells. Given that Wpl1p and Pds5p form a complex, we wondered whether they cooperate to inhibit condensation. Functional cooperation between Pds5p and Wpl1p is suggested by the fact that  $wpl1\Delta$  and specific N-terminal pds5 mutant alleles restore viability to eco1-ts cells that have reduced acetylase activity (Rowland et al. 2009; Chan et al. 2012). We further explored this relationship by assessing whether three of these N-terminal pds5 alleles (pds5-S81R, pds5-P89L, and pds5-*E181K*) shared all the phenotypes characteristic of a  $wpl1\Delta$ .

We first tested whether these pds5 alleles could suppress  $eco1\Delta$  inviability as a  $wpl1\Delta$  does. For this purpose, we constructed strains where pds5-S81R, pds5-P89L, or pds5-E181K was the sole pds5 allele in cells. We then generated ECO1 shuffle strains in these pds5 mutants and in a wild-type strain by introducing a centromere plasmid containing ECO1 and URA3 (ECO1 CEN URA3), then deleting ECO1 from its endogenous locus ( $eco1\Delta$ ). Counterselection against cells containing the ECO1 URA3 CEN plasmid by plating on media containing 5-FOA revealed whether any of the pds5  $eco1\Delta$  double-mutants were viable.

As expected, cells containing wild-type *PDS5* and *eco1* $\Delta$  could not grow on 5-FOA, as *ECO1* is an essential gene. In contrast, both the *pds5-S81R* and *pds5-P89L* alleles enabled robust growth on 5-FOA, indicating suppression of *eco1* $\Delta$  (Figure 3A). Consistent with our findings, previous results also showed that *pds5-P89L* restored viability to *eco1* $\Delta$  (Sutani *et al.* 

2009). In contrast, pds5-E181K did not support viability to  $eco1\Delta$  (Figure 3A). To determine whether the inability of pds5-E181K to restore viability to  $eco1\Delta$  was due to weak suppressor activity, we rebuilt pds5-E181K into a strain containing the eco1-203 temperature-sensitive allele. At the restrictive temperature,  $34^\circ$ , pds5-E181K eco1-203 cells grew (Figure S4A in File S1), consistent with a previous report in which pds5-E181K suppressed the inviability of the eco1-1 temperature-sensitive allele (Rowland et al. 2009). Thus, pds5-S81R and pds5-P89L are akin to a  $wpl1\Delta$  as they suppress an  $eco1\Delta$ , whereas pds5-E181K can only suppress inviability when Eco1p function is reduced but not abolished.

A second phenotype characteristic of  $wpl1\Delta$  cells is restoration of condensation but not cohesion to  $eco1\Delta$  cells (Guacci and Koshland 2012). This pattern of suppression distinguishes the inactivation of Wpl1p function from smc1 and smc3 suppressor mutants, which partially restore cohesion in eco1 mutants (Çamdere et al. 2015; Guacci et al. 2015). To test whether these pds5 N-terminal alleles mimicked the  $wpl1\Delta$  phenotype, we assessed chromosome condensation in pds5-S81R eco1 $\Delta$  and pds5-P89L eco1 $\Delta$  cells arrested in mid-M-phase. We utilized a standard method for assessing yeast chromosome condensation by monitoring the repetitive rDNA locus (Guacci and Koshland 1994; Guacci et al. 1997). The rDNA locus is located in the nucleolus and protrudes from the bulk chromosomal mass, making it easy to monitor its condensation state. A condensed rDNA locus forms a distinct loop structure, while decondensed rDNA locus form a "puff" morphology (Figure 3B; Materials and Methods) (Guacci et al. 1993; Guacci and Koshland 1994). We analyzed the morphology of the *rDNA* in cells synchronously arrested in mid-M-phase using nocodazole (Figure 3B). Since  $eco1\Delta$  cells are not viable, we used an ECO1-AID strain as a control for decondensation in cells depleted for Eco1p activity. The ECO1-AID strain was treated the same as other strains, except that 500 µM auxin was added to the media to induce ECO1-AID depletion from G1 through mid-M-phase arrest. Most wild-type (PDS5) cells that arrested in mid-M had condensed rDNA loops, so few had decondensed rDNA, whereas > 80% of ECO1-AID cells had decondensed rDNA (Figure 3C). Most pds5-S81R eco1 $\Delta$  and pds5-P89L eco1 $\Delta$ cells had condensed rDNA loops, so only  $\sim$ 20–30% of cells exhibited decondensed rDNA loci (Figure 3C). This result is similar to what was previously reported for  $eco1\Delta$  wpl1 $\Delta$  cells (Guacci and Koshland 2012). Thus, pds5-S81R and pds5-*P89L*, like wpl1 $\Delta$ , suppress the condensation defect engendered by loss of Eco1p activity.

We next assessed sister chromatid cohesion in pds5-S81R  $eco1\Delta$  and pds5-P89L  $eco1\Delta$  double-mutant cells synchronously arrested in mid-M. We monitored cohesion at either a chromosome IV CEN-distal or CEN-proximal locus by integration of LacO repeats at either the LYS4 or the TRP1 locus, respectively, in cells containing LacI-GFP (Materials and Methods). Cells containing a single LacI-GFP focus indicated cohesion, whereas cells containing two LacI-GFP foci indicated a loss of cohesion (Figure 3B). As expected from our



Figure 3 pds5 N-terminal mutants and SMC3-MCD1 fusion suppress inviability of  $eco1\Delta$  through restoration of condensation. (A) Plasmid shuffle assay to assess viability of pds5 N-terminal mutants in the  $eco1\Delta$  background. Plasmid pBS1030 (ECO1 CEN URA3) is present in haploid wild-type (WT) (VG3349-1B), eco1 $\Delta$  (VG3499-1B), eco1 $\Delta$  wpl1 $\Delta$ (VG3503 #4), eco1 pds5-S81R (MSB138-1K),  $eco1\Delta$  pds5-P89L (MSB139-2J), and  $eco1\Delta$  pds5-E181K (MSB147-1A) strains. Cells were grown to saturation in YPD media at 23°, plated at 10-fold serial dilutions on YPD or 5-FOA media, and then incubated for 3 days at 23°. 5-FOA selects for loss of pBS1030 (ECO1 CEN URA3). (B) Schematic of time course and analysis of cohesion and condensation. Cells were synchronously arrested in mid-M-phase as described in the Materials and Methods. Cells were processed for cohesion analysis of LacI-GFP at the CEN-distal LYS4 locus and CEN-proximal TRP1, and for condensation by FISH methodology (Materials and Methods). (C) pds5-S81R and pds5-P89L restore condensation in  $eco1\Delta$  cells. PDS5 (VG3349-1B), eco1-AID (VG3633-2D), eco1Δ pds5-S81R (MSB138-1K), and  $eco1\Delta$  pds5-P89L (MSB139-2J) were arrested in G1 using  $\alpha$ -factor, then synchronously arrested in mid-M-phase using nocodazole as described in the Materials and Methods. From G1 through mid-M-phase, 500 µM auxin was present in the media of the eco1-AID strain. Cells were fixed and processed for FISH (Materials and Methods). Chromosome condensation was assessed by morphology of the rDNA locus and cells were scored for condensed rDNA (loops) and defective condensation (puffs). The percentage of cells with defective rDNA condensation (decondensed) is plotted. (D) pds5-S81R eco1 $\Delta$  and pds5-P89L $eco1\Delta$  double-mutants have a dramatic defect on cohesion. PDS5 (VG3349-1B), eco1 wpl1 (VG3503 #4),  $eco1\Delta$  pds5-S81R (MSB138-1K), and  $eco1\Delta$  pds5-P89L (MSB139-2J) cells were synchronously arrested in mid-M-phase using nocodazole (Materials and Methods). Cells were scored for cohesion (one GFP focus) and loss of cohesion (two GFP foci; sister separation) at

the CEN-distal LYS4 locus. The percentage of cells lacking cohesion (sister separation) is shown. (E) SMC3-MCD1 fusion promotes condensation in eco1 $\Delta$  cells. WT (VG3349-1B), eco1-AID (VG3633-2D), eco1 $\Delta$  wp/1 $\Delta$  (VG3502 #A), and eco1 $\Delta$  SMC3-MCD1 (MSB249-3A) were synchronously arrested in mid-M-phase (Materials and Methods). From G1 through mid-M-phase, 500  $\mu$ M auxin was present in the media of the eco1-AID strain. Cells were fixed and processed for FISH to assess rDNA condensation (loops) and defective condensation (puffs), as described in (C). The percentage of cells with defective rDNA condensation (decondensed) is plotted. (F) SMC3-MCD1 fusion partially restores cohesion to eco1 $\Delta$  cells. SMC3-MCD1 (VG3940-2D) and eco1 $\Delta$  SMC3-MCD1 (MSB249-3A) were synchronously arrested in mid-M-phase using nocodazole (Materials and Methods). Cells were scored for cohesion (one GFP focus) and loss of cohesion (two GFP foci; sister separation) at the CEN-distal LYS4 locus as described in (B). The percentage of cells lacking cohesion (sister separation) is shown. The strains in this panel and in Figure 4C were analyzed for cohesion loss in the same experiment. The data were separated for clarity of presentation, and the SMC3-MCD1 cohesin data are presented here and in Figure 4C.

previous studies (Guacci and Koshland 2012), most sister chromatids remained tethered in wild-type cells, whereas ~70% of  $eco1\Delta$  wpl1 $\Delta$  cells exhibited separated sister chromatids at both loci (Figure 3D and Figure S4B in File S1). The pds5-S81R eco1 $\Delta$  and pds5-P89L eco1 $\Delta$  cells exhibited high levels of separated sisters at both CEN-proximal and distal loci, similar to that seen in eco1 $\Delta$  wpl1 $\Delta$  cells (Figure 3D and Figure S4B in File S1). Additionally, the pds5-E181K eco1-203 double-mutant and eco1-203 single-mutant strains also had high levels of sister separation when monitored at the restrictive temperature, 34° (Figure S4C in File S1). Thus, the *pds5* N-terminal suppressor mutants behave like *wpl1* $\Delta$ , as they restore viability and condensation but not cohesion to *eco1* $\Delta$  or *eco1*-*ts* cells.

The defects in cohesin function observed in *wpl1* $\Delta$  cells, along with evidence that Wpl1p destabilizes the interface between Smc3p and Mcd1p, suggest that this activity is important for promoting one or more of cohesin's biological functions. If so, one or more of these functions would be compromised when Wpl1p's destabilization activity was blocked by covalently fusing Smc3p to Mcd1p. Previously, a construct in which Smc3p was fused to the N-terminus of

Mcd1p (*SMC3-MCD1*) was shown to support viability when providing the sole sources of both Smc3p and Mcd1p in the cell (Gruber *et al.* 2006). Additionally, this fusion protein was able to restore viability to an *eco1* $\Delta$  mutant (Chan *et al.* 2012). This suppression of *eco1* $\Delta$  inviability suggested that the *SMC3-MCD1* fusion may suppress Wpl1p's ability to inhibit condensation.

To assess this possibility directly, we examined whether an SMC3-MCD1 fusion would suppress the condensation defect of an  $eco1\Delta$  like a  $wpl1\Delta$  does. We generated a strain in which the SMC3-MCD1 fusion was the sole source of both SMC3 and MCD1. Consistent with previous reports, we were then able to delete *ECO1* ( $eco1\Delta$ ) in this background. We examined rDNA condensation in wild-type, SMC3-MCD1, and SMC3-MCD1  $eco1\Delta$  cells that were synchronously arrested in M-phase (Materials and Methods). As before, we used an ECO1-AID strain as a control for decondensation. As expected, most wild-type cells had condensed rDNA loops, with few displaying decondensed rDNA, whereas most eco1-AID cells had decondensed rDNA (Figure 3E). Only a small percentage (~25%) of SMC3-MCD1 eco1 $\Delta$  cells had decondensed *rDNA*, similar to what was observed in wild-type and  $eco1\Delta$  wpl1 $\Delta$  cells, indicating that the SMC3-MCD1 fusion, like  $wpl1\Delta$ , is able to restore condensation to cells lacking Eco1p (Figure 3E).

We next assessed whether the *SMC3-MCD1* fusion restores cohesion to *eco1* $\Delta$  by comparing the *SMC3-MCD1* and *SMC3-MCD1 eco1* $\Delta$  strains. Strains were synchronously arrested in mid-M-phase (*Materials and Methods*) and cohesion was assessed at the *CEN*-distal *LYS4* locus. *SMC3-MCD1* exhibited a mild cohesion defect of ~20%. However, *SMC3-MCD1 eco1* $\Delta$ cells had an increased cohesion defect of ~40% (Figure 3F). This cohesion defect at the *CEN*-distal *LYS4* locus was less severe than the 70% seen in *eco1* $\Delta$  *wpl1* $\Delta$  cells and the 70– 80% seen in *eco1-AID* cells (Figure 3D; Guacci and Koshland 2012; Çamdere *et al.* 2015; Guacci *et al.* 2015). Therefore, the Smc3p-Mcd1p fusion protein can partially restore cohesion to *eco1* $\Delta$  cells, unlike *wpl1* $\Delta$  or the pds5 N-terminal mutants, which are unable to restore any cohesion.

# The Pds5p N-terminus and the Smc3p/Mcd1p interface promote cohesion establishment

A third phenotype characteristic of  $wpl1\Delta$  cells is their partial defect in cohesion establishment (Guacci and Koshland 2012). Budding yeast Pds5p-defective cells were already known to have a severe cohesion maintenance defect (Hartman *et al.* 2000; Panizza *et al.* 2000; Stead *et al.* 2003; Eng *et al.* 2014). The difference in the timing and severity of the cohesion defects suggested that Wpl1p and Pds5p might promote cohesion by distinct mechanisms. Alternatively, Pds5p might promote cohesion by two mechanisms, one dependent on Wpl1p and the other independent of Wpl1p. The phenotypic similarity between  $wpl1\Delta$  and the N-terminal alleles of pds5 described above suggested that the N-terminus of Pds5p might be involved in a Wpl1p-dependent pathway. We tested this possibility by monitoring the ability of pds5-S81R, pds5-P89L,

and *pds5-E181K* to mediate cohesion, either in the presence or in the absence of *WPL1*.

We first assessed cohesion in the *pds5* N-terminal mutants in an otherwise wild-type background (*WPL1*) in cells synchronously arrested in mid-M-phase. When cohesion was monitored at both *CEN*-proximal and *CEN*-distal loci, all three *pds5* N-terminal mutants exhibited cohesion defects of ~20 and ~30%, respectively, similar to that of *wpl1* $\Delta$  (Figure 4A and Figure S5A in File S1). The *pds5-P89L* mutant was previously shown to have a similar small cohesion defect at the *URA3* locus (Sutani *et al.* 2009). Additionally, the *pds5* N-terminal mutants lost cohesion with similar kinetics as *wpl1* $\Delta$  cells progressed from S- to M-phase (Figure S5B in File S1). These similarities suggest that Wpl1p and the Pds5p N-terminal domain act in a common pathway to promote efficient cohesion establishment.

To more directly assess whether Wpl1p and the Pds5p N-terminus function in a common pathway to promote cohesion, we examined cohesion in  $wpl1\Delta pds5$  N-terminal double-mutants. If Wpl1p and the Pds5p N-terminus function in a common pathway, we would expect the double-mutant to have cohesion defects similar to each single-mutant alone. In contrast, if Wpl1p and the Pds5p N-terminus function in distinct pathways, we would expect the double-mutants to have additive increases in cohesion defects approaching 60–70%. The cohesion defects of all three  $pds5 wpl1\Delta$  double-mutants (pds5-S81R  $wpl1\Delta$ , pds5-P89L  $wpl1\Delta$ , and pds5-*E181K* wpl1 $\Delta$ ) were ~30–40% when measured at *LYS4*, and  $\sim$ 20–30% when measured at TRP1. These cohesion defects were the same or only slightly higher than *pds5* single-mutants (*pds5-S81R*, *pds5-P89L*, and *pds5-E181K*) alone or *wpl1* $\Delta$  alone (Figure 4A and Figure S5A in File S1). The similar partial cohesion defects between pds5 single-mutants and pds5  $wpl1\Delta$  double-mutants suggest that the Pds5p N-terminus and Wpl1p promote cohesion through a common pathway. These results suggest that Wpl1p interacts functionally with Pds5p, both to inhibit condensation and to efficiently promote cohesion.

The cohesion defect that we observed in SMC3-MCD1 fusion was similar to that of  $wpl1\Delta$  cells (compare Figure 3F and Figure 4A). Similar results for both the SMC3-MCD1 fusion and the  $wpl1\Delta$  strains were previously reported at the more CEN-proximal URA3 locus (Gruber et al. 2006; Rowland *et al.* 2009). Given that Wpl1p is thought to destabilize the Smc3p/Mcd1p interface, these similarities suggested that the cohesion defect could be due to an inability of Wpl1p to modulate the interface in the fusion. Therefore, we reexamined cohesion in the  $wpl1\Delta$  mutant and the SMC3-*MCD1* fusion in the same experiment. *SMC3-MCD1* and *wpl1* $\Delta$ did indeed have similar moderate defects in cohesion of  $\sim$ 20-30% at the CEN-distal LYS4 locus when cells were arrested in mid-M-phase (Figure 4B). If the moderate cohesion defect in SMC3-MCD1 cells was due to an inability of Wpl1p to modulate this interface, then deleting WPL1 in the SMC3-MCD1 strain should have no further impact on the cohesion defect. Indeed, the SMC3-MCD1 wpl1 $\Delta$  strain had a similar partial



Figure 4 pds5 N-terminal mutants and SMC3-MCD1 fusion are defective for Wpl1p-mediated cohesion. (A) pds5 N-terminal mutants have a modest cohesion defect similar to  $wpl1\Delta$  alone or to pds5wpl1a double-mutants. Strains PDS5 (VG3349-1B), wpl1 $\Delta$  (VG3360-3D), pds5-S81R (MSB183-1A), pds5-P89L (MSB184-3A), pds5-E181K (MSB101-3C), pds5-S81R wpl1 $\Delta$  (MSB133-3C), pds5-P89L wpl1 $\Delta$ (MSB134-1L), and pds5-E181K wpl1 $\Delta$  (MSB223-1A) were synchronously arrested in mid-M-phase and scored for cohesion at the CEN-distal LYS4 locus, as described in Figure 3D. The percentage of cells lacking cohesion (sister separation) is shown. (B) SMC3-MCD1 has a similar modest cohesion defect to wpl1D. PDS5 (VG3349-1B), wp/1A (VG3360-3D), and SMC3-MCD1 (VG3940-2D) were synchronously arrested in mid-Mphase and scored for cohesion at CEN-distal LYS4, as described in Figure 3D. The percentage of cells lacking cohesion (sister separation) is shown. (C) Deletion of WPL1 has little effect on the cohesion of SMC3-MCD1 cells. SMC3-MCD1 (VG3940-2D) and SMC3-MCD1 wpl1 $\Delta$  (VG3957-1C) cells were synchronously arrested in mid-M-phase and scored for cohesion at the CEN-distal LYS4 locus, as in Figure 3D. The percentage of cells lacking cohesion (sister separation) is shown.

cohesion defect as the *SMC3-MCD1* strain (Figure 4C). Together, these results suggest that Wpl1p promotes efficient cohesion through the destabilization of the Smc3p/Mcd1p interface.

# Regulation of the Smc3p/Mcd1 interface, but not the Pds5p N-terminus, is important for resistance to DNA-damaging agents

The final characteristic  $wpl1\Delta$  phenotype is their sensitivity to the DNA-damaging agents CPT and MMS. As our results suggested that the N-terminus of Pds5p and Wpl1p function together in cohesion and condensation, we wondered whether they also function together to promote DNA-damage repair. To test this possibility, we examined effects on the growth of the *pds5* N-terminal mutants alone or in the *wpl1* $\Delta$  background by plating the single- and double-mutants on media containing either CPT or MMS. Surprisingly, cells containing pds5-S81R, pds5-P89L, and pds5-E181K alone grew similarly to wild-type cells on 20 µg/ml CPT and 0.015% MMS, and significantly better than  $wpl1\Delta$  cells. The  $wpl1\Delta$  pds5 double-mutants and  $wpl1\Delta$  cells were equally sensitive to both CPT and MMS (Figure 5A). These results suggest that Wpl1p's role in DNA-damage repair is independent of its functional interaction with the Pds5p N-terminus.

We also tested whether the ability to modulate the Smc3p/ Mcd1p interface was required for resistance to DNA-damaging agents. We compared the sensitivities of wild-type, *wpl1* $\Delta$ , and *SMC3-MCD1* fusion strains to 20 µg/ml CPT and to 0.01% MMS. The *SMC3-MCD1* fusion strain and the *wpl1* $\Delta$ strain showed similar growth inhibition to both drugs (Figure 5B). Together, our analyses of cells containing the Smc3-Mcd1p fusion protein suggest that Wpl1p destabilization of the Smc3p/Mcd1p interface is a common underlying mechanism necessary to promote the cohesion and repair of S-phase-induced DNA damage, as well as to inhibit condensation.

# Physical interaction between Wpl1p and Pds5p is not sufficient for regulation of cohesin function

After determining that Wpl1p and the N-terminus of Pds5p share a common function in promoting cohesion and the inhibition of condensation, we sought to determine if these functions derive from formation of the Wpl1p-Pds5p complex. If so, *pds5* N-terminal mutants might abrogate formation of the Wpl1p-Pds5p complex. Support for this idea came from a recent crystal structure of human Pds5B (Ouyang *et al.* 2016). This crystal structure also contained a short peptide from the N-terminus of Wapl, the human ortholog of Wpl1p, which bound to the N-terminus of Pds5B. As this Pds5B region is highly conserved with yeast Pds5p, we were able to map the analogous residues of the Pds5p N-terminal mutations onto the crystal structure of Pds5B (Figure 6A and Figure S6 in File S1). These residues were located either within, or in very close proximity to, the Wapl-binding site (Figure 6A).

Given this structural information, we asked whether the Pds5p N-terminal mutations disrupted the physical interaction between Pds5p and Wpl1p. We C-terminally tagged Wpl1p with the Flag epitope (Wpl1p-3FLAG) and then performed anti-FLAG immunoprecipitation from extracts of asynchronously growing cells. We compared Wpl1p co-immunoprecipitation with wild-type Pds5p and each of the N-terminal Pds5p mutants. The anti-FLAG immunoprecipitation robustly co-immunoprecipitated wild-type Pds5p when Wpl1p-3FLAG was present, but not when Wpl1p was untagged, confirming



**Figure 5** The *SMC3-MCD1* fusion, but not *pds5* N-terminal mutants, is defective for WpI1p-mediated DNA repair. (A) Assessment of sensitivity of *pds5* N-terminal mutants to camptothecin (CPT) and MMS. Cultures of cells in the *WPL1* background [*PDS5* (VG3349-1B) *pds5-S81R* (MSB183-1A), *pds5-P89L* (MSB184-3A), and *pds5-E181K* (MSB101-3C)] and the *wpI1*Δ background [*wpI1*Δ (VG3360-3B), *pds5-S81R wpI1*Δ (MSB204-1B), *pds5-P89L wpI1*Δ (MSB205-4C), and *pds5-E181K wpI1*Δ (MSB223-1A)] were serially diluted 10-fold and plated on YPD media either containing no drug, 20 µg/ml CPT, or 0.015% MMS, and incubated at 23° and assessed at 3 days postplating. (B) Assessment of sensitivity of *SMC3-MCD1* fusion to CPT and MMS. Wild-type (*WT*) (VG3349-1B), *wpI1*Δ (VG3360-3B), and *SMC3-MCD1* (VG3940-2D) were serially diluted 10-fold and plated on YPD media alone, or containing either 20 µg/ml CPT or 0.01% MMS, then incubated at 23° and assessed 3 days postplating.

that Pds5p co-immunoprecipitation is due to a specific interaction with Wpl1p (Figure 6B). The Wpl1p-3FLAG immunoprecipitates contained very little Pds5p-P89L and clearly reduced amounts of Pds5p-E181K (Figure 6B). Thus, both mutations disrupt Pds5p binding to Wpl1p. In contrast, Pds5p-S81R retained binding to Wpl1p-3FLAG at a level similar to wild-type Pds5p (Figure 6B). These differences in Wpl1p binding between the three N-terminal Pds5p mutants are surprising, as all three mutants similarly disrupt promotion of cohesion and restore viability to *eco1* mutants through restoration of condensation. These results suggest that binding to the Pds5p N-terminus is required for Wpl1p's function as both an inhibitor of condensation and efficient promoter of cohesion. However, this interaction is not sufficient for these functions, as Pds5p-S81R binds Wpl1p-3FLAG at close to wild-type levels.

The three *pds5* mutants have similar functional defects *in vivo*, despite their differences in the ability to bind Wpl1p. These differences suggest that the molecular function of Wpl1p must be attenuated through a mechanism other than Pds5p binding. Thus, we wondered whether the interaction between Wpl1p and cohesin might be compromised in these mutants. We probed the Wpl1p-3FLAG immunoprecipitates for the cohesin subunit, Mcd1p. In the wild-type Pds5p strain, Mcd1p exhibited robust co-immunoprecipitation with Wpl1p (Figure 6B). In contrast, in all three Pds5p N-terminal mutant strains, there was reduced

Mcd1p co-immunoprecipitation with Wpl1p (Figure 6B). We conclude from this result that formation of a functional Wpl1p-Pds5p complex is important for efficient recruitment of Wpl1p to cohesin. Additionally, Wpl1p was still able to interact with Mcd1p in the *pds5-P89L* mutant cells despite the Wpl1p interaction with Pds5p being abolished. This result indicates that Wpl1p can bind cohesin independently of Pds5p, which corroborates previous studies in yeast and other organisms that show that Wpl1p can interact directly with the cohesin subunit Scc3p/SA/STAG (Rowland *et al.* 2009; Shintomi and Hirano 2009).

In contrast to our conclusion that the Pds5p N-terminus functions with Wpl1p to inhibit condensation and promote cohesion, our studies suggest that Wpl1p promotes DNA repair independently of the Pds5p N-terminus. Consistent with this idea, pds5-*P89L* abrogates Wpl1p's interaction with Pds5p, but Wpl1p retains the ability to bind Mcd1p (Figure 6B). However, it is possible that DNA damage causes a modification to Pds5p or Wpl1p that promotes the formation of the Wpl1p-Pds5p complex. If so, the interaction between Wpl1p and Pds5p-P89L might be restored upon the induction of DNA damage. We assessed this possibility by treating asynchronously growing PDS5 WPL1-3FLAG and pds5-P89L WPL1-3FLAG cells with either DMSO or 20 µg/ml CPT for 3 hr. We immunoprecipitated Wpl1p in extracts from these cells and assayed for Pds5p binding. Wild-type Pds5p and Wpl1p co-immunoprecipitated at similar levels with or without CPT, whereas Pds5p-P89L remained unable to co-immunoprecipitate with Wpl1p under either condition (Figure 6C). These findings further corroborate our conclusion that Wpl1p promotes DNAdamage repair independently of its interaction with Pds5p.

#### Discussion

Previous studies in budding yeast have demonstrated roles for Wpl1p in promoting efficient sister chromatid cohesion and in inhibiting condensation (Guacci and Koshland 2012; Lopez-Serra *et al.* 2013). Here, we provide evidence for a biological function of Wpl1p in the timely repair of DNA damage in S-phase, beyond its roles in cohesion and condensation. We report that cells blocked for Wpl1p function grow slowly when they experience DNA damage induced during S-phase by CPT and MMS. This slow growth results from a delay in the onset of chromosome segregation, likely reflecting activation of the DNA damage checkpoint in G2/M because of slow repair of the damage. Consistent with this view, Wpl1p was shown to promote the timely repair of DSBs and homologous recombination in budding yeast meiotic chromosomes (Challa *et al.* 2016).

The defect in DNA repair in cells blocked for Wpl1p function cannot be explained by their partial cohesion defect. We showed that *pds5* N-terminal mutants have the same partial cohesion defect as *wpl1* $\Delta$  and *SMC3-MCD1* cells, yet these *pds5* mutants exhibit growth similar to wild-type cells when exposed to CPT and MMS. These results suggest that Wpl1p modulates a cohesin function in the repair of S-phase-induced DNA damage beyond simply its role in promoting sister chromatid cohesion. When DNA damage is induced in G2/M,



Figure 6 Pds5p N-terminus promotes Wpl1p binding to the cohesin complex. (A) Crystal structure of Pds5B bound to the YSR motif of Wapl from Ouyang et al. (2016). Gray: Pds5B; blue: Wapl peptide; red: eco1-ts suppressors from Rowland et al. (2009) and Sutani et al. (2009). Yeast residues were mapped to analogous residues on Pds5B through alignment. (B) Pds5p N-terminal mutants impair Wpl1p's binding to cohesin and but have different effects on Wpl1p's interaction with Pds5p. Wpl1p-3FLAG was immunoprecipitated from protein extracts in asynchronous cultures containing Wpl1p-3FLAG and either PDS5 (MSB192-2A), pds5-S81R (MSB193-1B), pds5-P89L (MSB194-1C), or pds5-E181K (MSB195-2D), as described in the Materials and Methods. No tag control contains wild-type (WT) untagged WPL1 and PDS5 alleles (VG3349-1B). For western blot analysis, Wpl1p was detected using mouse anti-FLAG, Pds5p was detected using rabbit anti-Pds5, and Mcd1p was detected using rabbit anti-Mcd1 antibodies (Materials and Methods). For anti-FLAG, a nonspecific species present in all cells is denoted by an asterisk. (C) Assessment of interaction between Wpl1p-3FLAG and Pds5p (MSB192-2A) or Pds5p-P89L (MSB194-1C) when treated with CPT. Asynchronous cultures were treated either with DMSO or 20 µg/ml CPT for 3 hr before being harvested. No tag control is the WPL1 PDS5 (VG3349-1B) strain, which was treated with DMSO. Immunoprecipitation (IP) and western blot analysis were performed as described in (B) and the Materials and Methods.

cohesin loading around the break site is stimulated (Ström *et al.* 2004; Unal *et al.* 2004). It may be that Wpl1p promotes cohesin binding at either sites of DNA damage or at replication forks to reinforce them upon damage, and thereby promotes DNA repair.

The results from our study suggest that Wpl1p regulates cohesin function in DNA repair, cohesion, and condensation through a common mechanism. We show that cells expressing the Smc3p-Mcd1p fusion protein, like *wpl1* $\Delta$ , have partial cohesion defects, are sensitive to S-phase DNA-damaging agents, and restore viability and condensation to cells lacking Eco1p. As Wpl1p destabilizes the interface between Smc3p and Mcd1p (Beckouët *et al.* 2016), the Smc3-Mcd1p fusion likely makes this interface refractory to Wpl1p function. The common phenotypes of *wpl1* $\Delta$  and the Smc3-Mcd1p fusion make it likely that Wpl1p-mediated regulation of the Smc3p/ Mcd1p interface is required for efficient cohesion, timely repair of DNA damage, and the inhibition of condensation.

Disruption of the Smc3p/Mcd1p interface by Wpl1p is thought to be one mechanism to remove cohesin from DNA (Chan *et al.* 2012). Our results suggest that Wpl1p-mediated removal of cohesin has both positive (efficient promotion of cohesion and DNA repair) and negative (inhibition of condensation) consequences. One model posits that cohesin and Pds5p regulate chromosome condensation by first binding DNA at sites along a chromatid to divide the chromosomes into domains, followed by axial shortening generated by looping out the intervening DNA (Guacci et al. 1997; Hartman et al. 2000). Interactions between cohesins may contribute to condensation by mediating looping (Guacci et al. 1997; Hartman et al. 2000). Wpl1p may inhibit condensation by preventing or inhibiting cohesin from tethering DNA in cis along a chromatid. This process could entail either destabilizing interactions between nonacetylated cohesins and/or destabilize the DNA binding of nonacetylated cohesins (Figure 7A). Indeed, studies showed that Wpl1p depletion increased the size of DNA loops in interphase human cells, induced precocious condensation in human and yeast cells, and induced hyper-condensation in yeast mitotic and meiotic cells (Lopez-Serra et al. 2013; Tedeschi et al. 2013; Challa et al. 2016; Haarhuis et al. 2017). Moreover, the ability of  $eco1\Delta$  wpl1 $\Delta$  cells to condense their rDNA demonstrates that nonacetylated cohesin can promote condensation (Guacci and Koshland 2012). It is curious, though, that destabilization of cohesin's interaction with DNA would promote DNA repair and cohesion. These positive aspects may reflect cohesin's burden to carry out diverse biological functions.

Cohesin is loaded onto DNA at cohesion-associated regions prior to S-phase to establish cohesion, but cohesin is also loaded *de novo* at sites of damage and at stalled replication forks (Ström *et al.* 2004; Unal *et al.* 2004; Tittel-Elmer *et al.* 



Figure 7 Model for Wpl1p promotion and inhibition of cohesin function through recycling. (A) Model for Wpl1p inhibition of condensation. Cohesin mediates condensation through chromosome looping. Nonacetylated cohesin (red) promotes condensation, while acetylated cohesin (blue) promotes cohesion. Wpl1p antagonizes condensation by countering interactions of nonacetylated cohesin and/or by removing nonacetylated cohesin from DNA. (B) Wpl1p promotes cohesion and DNA-damage repair. Top: when Wpl1p is present, nonacetylated cohesin (red) is loaded onto DNA by Scc2/4p, and is removed from DNA by Wpl1p, maintaining a soluble pool of cohesin. Cohesin loading, followed by Eco1p acetylation, promotes cohesion, which is refractory from Wpl1p. Upon DNA damage, nonacetylated cohesin is removed from other sites in the genome and/or is recruited from the soluble pool and loaded around the damage site. Bottom: in the absence of Wpl1p (wpl1 $\Delta$ ), cohesin is loaded onto DNA by Scc2/ 4p. The smaller amount of cohesin on DNA and in the soluble pool makes cohesion establishment less efficient or less robust. Upon DNA damage, cohesin removal from DNA and/or mobilization from the smaller soluble pool is limited. Thus, cohesin loading around damage sites is less efficient.

2012). These spatially and temporally distinct functions may require cohesin's mobilization, either from a DNA-bound or a nucleoplasm pool of cohesin (Figure 7B, top). In the absence of Wpl1p function, cohesin is stabilized on DNA, so is less efficiently mobilized and perhaps trapped in nonproductive sites. Additionally,  $wpl1\Delta$  cells exhibit a twofold decrease in cohesin (Mcd1p) bound on DNA and in cells (Figure S7A in File S1; Rowland et al. 2009; Sutani et al. 2009; Guacci et al. 2015). In wild-type cells, most yeast cohesin is bound to chromosomes but a small soluble pool remains (Tóth et al. 1999; Tong and Skibbens 2014). *wpl1* $\Delta$  cells show decreases in both the chromosomal and soluble pools (Figure S7B in File S1). This decrease in cohesin levels in  $wpl1\Delta$  cells may limit the pool of dynamic cohesin, regardless of whether it derives from a chromosomal or nucleoplasmic source. Thus, cohesion promotion is less efficient both during a normal cell cycle and in response to DNA damage (Figure 7B, bottom). We cannot rule out that the premature/hyper-condensation or aberrant chromosome structures in wpl1-depleted cells could contribute to DNA-damage sensitivity by hindering repair.

The necessity of maintaining a dynamic pool of cohesin is supported by a number of observations. First, only  $\sim$ 20–30% of cohesin is acetylated during S-phase to establish sister chromatid cohesion (Zhang *et al.* 2008). As acetylated cohe-

sin is thought to be refractory to Wpl1p activity (Rolef Ben-Shahar et al. 2008; Unal et al. 2008), this small percentage of acetylated cohesin may ensure that most cohesin remains responsive to Wpl1p. Second, we previously showed that a three- to fourfold reduction in the total cellular pool of cohesin leads to more severe defects in condensation and DNA repair than in cohesion (Heidinger-Pauli et al. 2010). These phenotypes may arise from a larger proportion of the remaining cohesin being locked onto the DNA in the cohesive (acetylated) state that is refractory to Wpl1p, and thus not available for recycling to promote condensation and DNA repair. In this light, the primary biological function of Wpl1p may not be to inhibit cohesin's function by removing it from the DNA. Rather, it would be to generate a dynamic cohesin pool for redistribution to different chromosomal sites to perform cohesin's distinct biological functions.

The idea that Wpl1p mobilizes cohesin to perform different biological functions can explain two seeming paradoxes from our studies. First, we found that the three *pds5* N-terminal mutants dramatically differ in their ability to bind Wpl1p, yet mimic *wpl1* $\Delta$  in their failure to efficiently promote cohesion and inhibit condensation. Second, we found that all three *pds5* mutants differ from *wpl1* $\Delta$  in their resistance to DNA-damaging agents. As *wpl1* $\Delta$  cells and *pds5-P89L* cells exhibit the same almost twofold decrease in cohesin bound to chromosomes (Rowland

et al. 2009; Sutani et al. 2009; Guacci and Koshland 2012), the difference in sensitivity cannot be due to different levels of cohesin binding. However, these paradoxes may be explained by the similar reduction in the amount of Wpl1p bound to cohesin that we observed in the three *pds5* N-terminal mutants. The regulation of both cohesion and condensation entails the modulation of cohesin at many sites genome-wide. pds5 N-terminal mutants may reduce the amount of Wpl1p bound to cohesin below a threshold required to mobilize the global pool of cohesin, thereby impairing the proper regulation of cohesion and condensation. In contrast, repair of DNA damage is likely to involve a small number of genomic sites, so may require a smaller pool of cohesin. The reduced levels of Wpl1p bound to cohesin in the pds5 N-terminal mutants may be sufficient to mobilize enough cohesin to promote DNA repair. As there is no Wpl1p present in  $wpl1\Delta$  cells, all three cohesin biological functions would be defective.

In addition to the insights into the relationship between Wpl1p and cohesin, our work furthers our understanding of the relationship between Wpl1p and Pds5p. Two pds5 N-terminal alleles either entirely (pds5-P89L) or partially (pds5-E181K) disrupt the interaction with Wpl1p. However, Wpl1p still binds cohesin in these cells, albeit at reduced levels. These data suggest that one function of the Wpl1p-Pds5p complex is to help recruit Wpl1p to cohesin. However, Wpl1p can also bind cohesin independently of its ability to complex with Pds5p. This independence has previously been demonstrated in vitro (Rowland et al. 2009; Shintomi and Hirano 2009). The pds5-S81R mutation preserves the interaction between Pds5p and Wpl1p, yet its effects on cohesin function are the same as *pds5-P89L*, which abolishes this interaction. This result indicates that the formation of a Wpl1p-Pds5p complex is not sufficient for Wpl1p's function. The human Wapl N-terminus contains a conserved motif, [K/R][S/T]YSR, which has been shown to interact with the Pds5B N-terminus through crystal structure analysis (Figure 6A; Ouyang et al. 2016). This YSR motif is critical for the interaction between Wapl and Pds5B, and for Wapl function in vertebrate cells (Ouyang et al. 2016). The N-terminus of budding yeast Wpl1p contains a partial consensus of this motif, suggesting that Wpl1p may bind the yeast Pds5p N-terminus in a similar manner (Ouyang et al. 2016). As other cohesin regulators have also been shown to contain YSR-like motifs (Ouyang et al. 2016; Goto et al. 2017; Zhou et al. 2017), mutating the N-terminus of Pds5p may also compromise interaction with these factors, contributing to defects in cohesin function and regulation. We find that binding of Wpl1p to Pds5p requires interaction with the N-terminal Pds5p residues, but that this interaction is not sufficient for Wpl1p function. It may be that this complex must be activated, possibly through a conformational change in either one or both proteins for proper function, to enable interaction with other proteins.

Our studies provide new insights into how the Wpl1p-Pds5p complex regulates cohesin's functions *in vivo*. Our proposed role for Wpl1p in recycling cohesin, in part through its association with Pds5p, assigns a biological function for the previous finding that the Wpl1p-Pds5p complex promotes

both cohesin loading and unloading *in vitro* (Murayama and Uhlmann 2015). Finally, exploring the roles of Wpl1p and cohesin in DNA-damage repair will open up an exciting new direction in the cohesin field.

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