Effects of the microtubule nucleator Mto1 on chromosomal movement, DNA repair, and sister chromatid cohesion in fission yeast

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ABSTRACT Although the function of microtubules (MTs) in chromosomal segregation during mitosis is well characterized, much less is known about the role of MTs in chromosomal functions during interphase. In the fission yeast Schizosaccharomyces pombe, dynamic cytoplasmic MT bundles move chromosomes in an oscillatory manner during interphase via linkages through the nuclear envelope (NE) at the spindle pole body (SPB) and other sites. Mto1 is a cytoplasmic factor that mediates the nucleation and attachment of cytoplasmic MTs to the nucleus. Here, we test the function of these cytoplasmic MTs and Mto1 on DNA repair and recombination during interphase. We find that mto1Δ cells exhibit defects in DNA repair and homologous recombination (HR) and abnormal DNA repair factory dynamics. In these cells, sister chromatids are not properly paired, and binding of Rad21 cohesin subunit along chromosomal arms is reduced. Our findings suggest a model in which cytoplasmic MTs and Mto1 facilitate efficient DNA repair and HR by promoting dynamic chromosomal organization and cohesion in the nucleus.

INTRODUCTION

The dynamic organization of chromosomes within the eukaryotic nucleus is essential for the proper regulation of gene expression, ribosome synthesis, RNA processing and transport, and DNA replication and repair (Schneider and Grosschedl, 2007; Misteli and Heldestad, 2009). One of the better characterized functions of the NE is the nuclear localization and attachment of cytoplasmic MTs to the NE. Here, we test the function of these cytoplasmic MTs and Mto1 on DNA repair and recombination during interphase. We find that mto1Δ cells exhibit defects in DNA repair and homologous recombination (HR) and abnormal DNA repair factory dynamics. In these cells, sister chromatids are not properly paired, and binding of Rad21 cohesin subunit along chromosomal arms is reduced. Our findings suggest a model in which cytoplasmic MTs and Mto1 facilitate efficient DNA repair and HR by promoting dynamic chromosomal organization and cohesion in the nucleus.
FIGURE 1: Microtubule-dependent movement of spindle pole bodies (SPBs) and DNA loci during interphase in *S. pombe*. (A) Schematic representation of interphase microtubule cytoskeleton in fission yeast, and its connections to the nucleus. The red arrow represents the direction of the force generated by MT polymerization after hitting the
Cytoskeleton-based forces may also affect the chromosomes inside the nucleus. There are well-documented cases showing cytoskeletal contributions to meiotic recombination. Meiotic chromosomal movements are driven by actin-based mechanisms in budding yeast (Conrad et al., 2008; Koszul et al., 2008; Christophorou et al., 2015) and by MTs in the fission yeast (Ding et al., 2004), Caenorhabditis elegans (Sato et al., 2009), and Drosophila (Hampole et al., 2011). In fission yeast, for instance, MTs drive dynein-dependent large oscillatory “horsetail” movements of the nucleus that are required for efficient meiotic homologous recombination (HR) (Ding et al., 2004).

The role of interphase MTs in nonmeiotic cells in chromosomal behaviors is less clear. During interphase in fission yeast, MTs are organized into multiple cytoplasmic bundles attached at the spindle pole body (SPB) and other sites on the cytoplasmic face of the NE. No MTs are present inside the nucleus during interphase (Hoog and Antony, 2007). Cytoplasmic MTs exert pushing forces to produce oscillatory movements or rotations of the SPB and nucleus and dynamically position the nucleus at the cell middle (Tran et al., 2001; Daga et al., 2006). These MT forces are transmitted to the centromeres of all three chromosomes via SPB, which is located just outside the NE, through LINC complexes, Csi1, and other centromeric proteins (Hou et al., 2012; Fernandez-Alvarez et al., 2016). Although the connection between the SPB and centromere is bridged by kinetochore MTs during mitosis, this connection is MT independent during interphase. Centromeric loci and multiple noncentromeric loci have been demonstrated to exhibit MT-dependent movements, suggesting that these forces from cytoplasmic MTs mediate large-scale chromosomal movements inside the nucleus (Kim et al., 2013).

The possible role of MTs in chromosomal organization and functions are not well understood. MTs, LINC complexes, and other NE factors have recently been implicated in DNA repair (Swartz et al., 2014; Lottersberger et al., 2015; Lawrence et al., 2016). In Schizosaccharomyces pombe, LINC complexes composed of Sad1/Unc84 and Klaricht/Anc1/SYNE1 homology protein Kms1 are recruited to sites of DNA damage (Swartz et al., 2014); however, whether and how MTs themselves contribute to DNA repair has not been thoroughly explored.

Here, we test the role of cytoplasmic MTs on DNA repair in S. pombe. We use the mto1Δ mutant as a tool to specifically disrupt the association of cytoplasmic MTs with the NE. Mto1 is a MT nucleator factor that forms a complex with Mto2 and the γ-tubulin ring complex to promote MT nucleation at cytoplasmic sites (Sawin et al., 2004; Venkatram et al., 2004; Zimmerman and Chang, 2005; Samejima et al., 2008; Bao et al., 2018). mto1Δ cells exhibit a uniquely strong and specific cytoplasmic MT nucleation defect; they either lack cytoplasmic MTs completely or form a small number of MT bundles that are not physically connected to the nucleus. In these cells, consistent with the lack of MT attachment, the nucleus is abnormally shaped and/or mispositioned, and the SPB oscillations are not observed (Sawin et al., 2004; Venkatram et al., 2004; Zimmerman and Chang, 2005; Daga and Nurse, 2008). In contrast, during mitosis, MTs in mto1Δ mutants are nucleated normally inside the nucleus for spindle assembly. Consistent with a cytoplasmic function, Mto1 localizes to cytoplasmic MTOCs but has not been detected in the nucleus (Sawin et al., 2004; Zimmerman and Chang, 2005). Here, we find that the inhibition of cytoplasmic MTs in the mto1 mutant or by drug treatment leads to significant defects in DNA repair and HR. In investigating the cause of this phenotype, we unexpectedly find that these cells have defects in sister chromatid pairing and loading or maintenance of the cohesin Rad21. Thus, these findings provide new insights into the role of MTs and this MT nucleation factor in chromosomal organization and maintenance.

**RESULTS**

**Interphase MTs are required for SPB and chromosomal movements**

We tested the effect of cytoplasmic MTs and Mto1 on the movement of the SPB and chromosomes. We imaged live fission yeast cells in which the SPB was marked with Sid2-tdTomato and two different chromosomal loci were marked with LacO arrays that were bound by green fluorescent protein (GFP)-LacI at lys1 and his2 loci, which are 30 kb away from centromere 1 and 0.6 Mb away from centromere 2, respectively (Figure 1A) (Molnar et al., 2003). In wild-type cells, the SPB and the lys1 locus moved together in oscillatory movements with approximately the same mean velocity (Figure 1, B–D). The his2 locus also displayed oscillatory movements similar to the SPB (Figure 1E). This chromosomal locus usually moved in the same direction as the SPB, but with reduced mean velocity relative to the SPB (Figure 1, E–G). These movements are dependent on MTs, as they were abolished after treatment with the MT-depolymerizing drug methyl benzimidazol-2-yl-carbamate (MBC) (Figure 1, B–G). Similarly, in the mto1Δ mutant, the oscillatory movements of the SPB and both chromosomal loci were absent (Figure 1, B–G). Thus, MTs and Mto1 are needed for large movements of chromosomes observed during interphase.

**mto1Δ and NE protein mutants are sensitive to DNA damage**

We next tested whether MT-dependent movements contribute to DNA repair. As MBC inhibits mitotic cell cycle progression, we initially focused on characterizing the mto1Δ mutant, which has a more specific defect in interphase MTs (Sawin et al., 2004; Venkatram et al., 2004; Zimmerman and Chang, 2005). We used a standard spot growth assay to measure sensitivity to DNA-damaging agents. We found that mto1Δ cells are sensitive to methyl-methane sulfonate (MMS), camptothecin (CPT), and γ-irradiation (Figure 2, A and B) (see also Swartz et al., 2014), which produce DNA damage that requires HR for its repair. They were not, however, sensitive to 254-nm UV irradiation and only mildly sensitive to hydroxyurea (Figure 2A), both of which cause different types of DNA damage.

cell tips. Black arrows indicate the position of the lys1 and his2 loci. ChrI/ChrII, chromosome I/II. The SPB is depicted in orange. Centromeres are depicted in yellow. (B) Kymographs showing movements of the SPB (marked with Sid2-tomTom) and chromosome at lys1 locus in wild-type (wt) cells, wild-type cells treated with 10μg/ml MBC, and mto1Δ cells. Three representative cells are shown in each case. Kymographs were prepared from maximal projections of three z-sections with a step size of 0.4 μm. Time between frames is 2 s with total time of 300 s. Scale bar: 5 μm. (C, D) Graphs showing mean instantaneous velocities of the SPB and the lys1 locus in the indicated strains and conditions (n = 50).

(E) Kymographs showing the SPB (marked with Sid2-tomTom) and chromosome at his2 locus. Three representative cells are shown in each case. Kymographs were prepared from maximal projections of three z-sections with a step size of 0.4 μm. Time between frames is 2 s. Total time is 300 s. Scale bar: 5 μm. (F, G) Graphs showing mean instantaneous velocities of the SPB and his2 locus (n = 50). **** denotes p < 0.0001 and ** denotes p < 0.001 from a Student’s t test.
This profile of DNA damage sensitivity suggests a defect of mto1Δ in HR-based DNA repair, or in the DNA damage checkpoint signaling.

We next tested what functions of Mto1 are needed for this phenotype by using different mto1 alleles and also mutants affected in MT dynamics. The mto1-9A2 mutant is defective in the interaction with the γ-tubulin complex and in MT nucleation and, as a consequence, shows reduced SPB movements (Samejima et al., 2008). We found that this mutant was sensitive to MMS (Figure 2A). In contrast, the mto1-1051 mutant, which is defective for MT attachment on the NE at non-SPB sites, but still exhibits MTs associated with the SPB and SPB movements (Samejima et al., 2008), was not sensitive to MMS (Figure 2A). Other MT regulatory mutants, such as mal3Δ (EB1) and tip1Δ (CLIP170), which have effects on MT dynamics but retain some SPB movement, were not MMS sensitive (Figure 2A). Note, the mto2 mutant, which has a weak MT nucleation effect and still exhibits SPB movements (Samejima et al., 2005; Venkatram et al., 2005), was not MMS sensitive (Figure 2A). Thus, these results suggest that efficient DNA damage response is mediated by Mto1 functions in MT nucleation and SPB movements.

To evaluate the effect of chromosomal movements on the sensitivity to DNA damage, we tested csi1Δ and INM protein Lem2 or yellow fluorescent protein (YFP) in live cells (Lisby et al., 2003; Meister et al., 2005). Wild-type cells typically form a small number of Rad52-YFP foci after S-phase during postreplication DNA repair. The disassembly of the repair factory is thought to occur upon repair (Meister et al., 2003). In asynchronous cultures, 15% of wild-type cells contained a single Rad52-YFP focus (Figure 3, A and B) (Meister et al., 2003, 2005). In mto1Δ cells, we found a twofold increase in the fraction of cells containing one Rad52 focus, and 10% of mto1Δ cells contained two foci (Figure 3, A and B). Time-lapse microscopy showed that, in wild-type cells, the lifetime of Rad52 focus is on average 60 min (Meister et al., 2003). In asynchronous cultures, 15% of wild-type cells contained a single Rad52-YFP focus (Figure 3, A and B) (Meister et al., 2003, 2005). In mto1Δ cells, we found a twofold increase in the fraction of cells containing one Rad52 focus, and 10% of mto1Δ cells contained two foci (Figure 3, A and B). Time-lapse microscopy showed that, in wild-type cells, the lifetime of Rad52-YFP foci was shorter than in Mto1Δ cells (Figure 3, C and D).

Increased sensitivity to DNA damage could be due to defects in DNA repair or, alternatively, in DNA checkpoint signaling. To test this, we conducted epistasis tests of mto1Δ with chk1Δ, cdc15Δ, and rad3Δ mutants that are defective in the DNA damage checkpoint, S-phase checkpoint, and general response to all types of DNA damage, respectively (Harrison and Haber, 2006). mto1Δ further increased MMS sensitivity of these checkpoint mutants (Figure 2C), suggesting that Mto1 does not merely act through these DNA checkpoint pathways. Cds1 regulates the response to S-phase DNA damage (Lindsay et al., 1998; Rhind and Russell, 2000), and accordingly, cdc15Δ cells were not sensitive to MMS (Figure 2C). Double cdc15Δ mto1Δ mutants showed sensitivity to MMS that was similar to that of mto1Δ. Together, these results suggest that mto1 mutants have a defect in HR-based DNA repair that is exacerbated in the absence of checkpoint signaling (rad3Δ, chk1Δ).

Mto1 and microtubules affect DNA repair factories

To examine DNA repair in vivo, we imaged DNA repair factories marked with Rad52–yellow fluorescent protein (YFP) in live cells (Lisby et al., 2003; Meister et al., 2005). Wild-type cells typically form a small number of Rad52-YFP foci after S-phase during postreplication DNA repair. The disassembly of the repair factory is thought to occur upon repair (Meister et al., 2003). In asynchronous cultures, 15% of wild-type cells contained a single Rad52-YFP focus (Figure 3, A and B) (Meister et al., 2003, 2005). In mto1Δ cells, we found a twofold increase in the fraction of cells containing one Rad52 focus, and 10% of mto1Δ cells contained two foci (Figure 3, A and B). Time-lapse microscopy showed that, in wild-type cells, the lifetime of Rad52 focus is on average 60 min (Meister et al., 2003). In asynchronous cultures, 15% of wild-type cells contained a single Rad52-YFP focus (Figure 3, A and B) (Meister et al., 2003, 2005). In mto1Δ cells, we found a twofold increase in the fraction of cells containing one Rad52 focus, and 10% of mto1Δ cells contained two foci (Figure 3, A and B). Time-lapse microscopy showed that, in wild-type cells, the lifetime of Rad52-YFP foci was shorter than in Mto1Δ cells (Figure 3, C and D).

In contrast, the repair factories in mto1Δ cells exhibited a twofold increase in Rad52-YFP intensity and an increase in average lifetime distribution (Figure 3, C–E). Mto1 therefore affects the dynamics of DNA repair factories.

To test whether MTs also affect Rad52 repair factory behavior, we determined the effects of depolymerizing cytoplasmic MTs in wild-type cells during a defined cell cycle period. For that, we imaged asynchronously growing cells in time lapse, introduced MBC (time 0), and then specifically tracked the individual cells in which MBC addition at t0 coincided with late anaphase B (just before S-phase). As cells went through S and G2 phases in MBC, we observed the
FIGURE 3: Mto1 and microtubules affect the dynamics of DNA repair factories. (A) Maximal-projection images of exponentially growing wild-type (wt) and mto1Δ cells expressing Rad52-YFP as marker of DNA repair factories (indicated by arrowheads). Scale bar: 5 μm. (B) Graphs showing the quantitation of factory number in wild-type and mto1Δ mutant (n = 120 cells). Error bars show SD of three independent experiments. (C) Time profiles of Rad52 factory intensity in wild-type cells, wild-type cells treated in late anaphase with 10 μg/ml MBC, and mto1Δ cells (three representative examples of each condition are shown). (D) Graphs showing total Rad52-YFP foci lifetime in the indicated strains and conditions. Error bars represent SD of three independent experiments. (E) Graph showing peak fluorescence intensity of Rad52-YFP foci in the indicated strains and conditions. There were 18 cells analyzed in D and E. Error bars represent SD. (F–G) Response to laser-induced DNA damage of wild-type and mto1Δ cells. Rad52-YFP was followed by time-lapse microscopy at 1-min time intervals. A single 10-ms pulse with a 355-nm laser was targeted at a fixed area of 500 nm of the nuclei of wild-type and mto1Δ cells at time 5 min. (F) Rad52-YFP repair factories after laser damage in a representative wild-type and mto1Δ cell. Time is indicated in minutes. N, nucleus. (G) Average intensity of Rad52-YFP repair factories is shown as a function of time (n = 15 cells for each strain). Error bars are SD.
Mto1 affects sister chromatid cohesion

Because DNA repair based upon HR during G2 requires proper pairing of sister chromatids (Birkenbihl and Subramani, 1992; Hartsuiker et al., 2001; Jessberger, 2002), we next examined the effects of Mto1 and microtubules on sister chromatid cohesion. First, we imaged the dynamic behavior of GFP-labeled LacO arrays on chromosomal loci during the G2 phase in live cells. Time-lapse confocal imaging using GFP-labeled arrays at the his2 locus revealed that sister chromatids during interphase exhibited occasional “breathing” events in which the labeled loci moved apart (>200 nm) to appear transiently as two dots and then came back together in a period of 2 s (Figure 5, A and B). This behavior was seen infrequently in wild-type cells, but this transient separation of the sister loci occurred more frequently in mto1 mutant and cells treated with 200 nm) (Supplemental Figure S2A). Specifically we could not detect Mto1 at DNA repair factories upon MMS-induced DNA damage (Supplemental Figure S2B).

Together, these experiments show that Mto1 and MTs affect the behavior of repair factories for postreplication repair and the repair of exogenous DNA damage, and they may do so indirectly from the cytoplasm.

Homologous recombination is reduced in mto1Δ cells

As HR between sister chromatids is the primary pathway for DNA repair in G2 (Harrison and Haber, 2006), we assayed for possible defects in HR in the mto1Δ mutant. Intrachromosomal recombination (i.e., between sister chromatids) was assayed using a genetically based approach developed by Ahn et al. (2005). This assay uses strains containing the polar replication fork barrier (RFB) RTS1 oriented either in the direction of the replication (toward the centromere) or in the opposite direction, between a direct repeat of ade6 heteroalleles located at the ade6 locus on chromosome 3 (Chr III). This allowed us to measure HR induced by RFB, and much lower rates of spontaneous HR at this locus (Figure 4A). Recombination rates were decreased by 10-fold in mto1Δ strains in both recombination substrates (Figure 4B). The magnitude of the recombination defect is comparable to that seen in mutants in key recombination proteins such as Rhp51 (Ahn et al., 2005). We also found a similar decrease of recombination efficiency in both replication fork substrates when mto1 was deleted in the swi1Δ background (Ahn et al., 2005), which abolished the difference between the two substrates (Figure 4C). In contrast, mto1 had little effect on interchromosomal HR between the double strand break (DSB)-induced recombination of Chr III and an artificial homologous Chr III fragment (Prudden et al., 2003) (Supplemental Figure S3, A and B). These findings indicate that Mto1 is required for efficient sister chromatid-based HR.

formation and resolution of the postreplicative repair factories marked by Rad52 foci. The intensities and lifetimes of the Rad52 foci were increased in MBC-treated cells relative to untreated cells, similar to what was seen in mto1Δ cells (Figure 3, C–E). Thus, inhibition of MTs after mitosis leads to abnormal DNA damage response in interphase.

To measure the dynamics of the DNA damage response in a quantitative manner, we induced DNA damage at a defined time point by applying a focused 410-nm laser beam on the nucleus of living cells. In both wild-type and mto1Δ cells, Rad52-YFP started to accumulate in a single dot at 5 min after irradiation, which steadily increased in intensity over the 30-min time course (Figure 3, F and G). Measurements of fluorescence intensity showed that Rad52-YFP intensity and the rate of accumulation were increased in mto1Δ compared with wild-type cells throughout the 30-min time course.

We examined whether Mto1 localizes to DNA repair factories and found that Mto1-mCherry was not detectable within the nucleus, as previously shown (Sawin et al., 2004; Venkatram et al., 2005) (Supplemental Figure S2A). Specifically we could not detect Mto1 at DNA repair factories upon MMS-induced DNA damage (Supplemental Figure S2B).

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FIGURE 5: Defects in sister chromatid cohesion in mto1Δ and microtubule depolymerized cells. (A) Kymographs of representative examples of increased separation between sister chromatids as shown by GFP at his2 locus in the indicated strains and conditions (blue arrowheads). Images are maximal projections of six z-sections with a step size of 0.4 μm acquired every 2 s. Scale bar: 2.5 μm. (B) Plots showing dynamics of loci in time-lapse images in the indicated strains and conditions. Distances between his2 loci on sister chromatids are plotted over 300 s. Each color represents an individual cell (n = 10 cells). (C) Percentage of cells of the indicated strains and conditions showing separated GFP dots on his2 loci at a single time point (n = 50 cells for each condition). psc3-1T cells were imaged at 25°C or after 2.5 h at 36°C. Error bars represent the SD from three independent experiments. (D) Graph showing the average maximum GFP-his2 loci separation observed over 300 s in cells shown in B. Error bars represent the SD. (E) Graph showing frequency of separation events per second in cells showed in B. Error bars represent the SD. **, p < 0.01; ***, p < 0.001; ****, p < 0.0001. wt, wild type.
MBC (Figure 5, A–C, and Supplemental Movies S1 and S2). In wild-type cells, loci showed on average 4.8 ± 2.3 separation events over a period of 5 min (Figure 5, A, B, and E, and Supplemental Figure S5). In cells treated with MBC and in mto1Δ cells, we observed an average of 18.4 ± 12.8 events and 27.2 ± 15.7 events, respectively, during the same time period (Figure 5, A, B, and E, and Supplemental Figure S5). The maximum distance reached between the loci was also increased in wild-type cells treated with MBC and in mto1Δ cells (0.61 ± 0.11 and 0.59 ± 0.05, respectively) compared with wild-type untreated cells (0.45 ± 0.09) (Figure 5, A, B, and D, and Supplemental Figure S5). The time that the loci spent apart was also increased. In wild-type cells, sister loci spent on average 2.2 ± 0.25 s apart, whereas in cells treated with MBC and mto1Δ cells, the loci spent on average 3.1 ± 1.1 and 4.6 ± 1.9 s apart, respectively, although we occasionally observed breathing events that persisted for >25 s (Figure 5, A and B, and Supplemental Figure S5).

We tested whether this breathing behavior is due to decreased cohesion between sister chromatids. To test this, we compared this behavior with that of a cohesin subunit mutant psc3-1T (Nonaka et al., 2002). Cohesin mutants have been shown to exhibit a higher percentage of cells with sister chromatid or centromere separation in fixed samples (Tomonaga et al., 2000; Bernard et al., 2001; Tanaka et al., 2001; Nonaka et al., 2002), but the dynamics of chromosomal movements using live-cell imaging has not been previously shown. In time-lapse imaging of the his2Δ loci, we found increased sister chromatid breathing in the temperature-sensitive psc3-1T mutant at nonpermissive temperature (Figure 5, A–E). These behaviors were similar but slightly more severe than those seen in the mto1Δ mutant. Thus, the mto1Δ mutant and cells treated with MBC exhibit defects similar to those of the psc3 cohesin mutant.

We also assayed breathing behavior of chromosomal arrays at the lys1 site, close to the centromere (Supplemental Figures S4, A–E, and S6). Here, we also detected similar breathing events. However, in contrast to the his2Δ site, the frequency of these events at lys1 was significantly elevated in the psc3-1T mutant, but not in mto1Δ and MBC-treated cells. This suggests that cohesion at the centromere is defective in the psc3-1T mutant but not in the mto1Δ mutant.

This defect in sister chromatid cohesion raises the possibility that Mto1 and microtubules affect the distribution or function of the cohesin complex. We used cohesin subunit Rad21 as a marker for the cohesin complex (Tomonaga et al., 2000; Tanaka et al., 2001; Nonaka et al., 2002), but the dynamics of chromosomal movements using live-cell imaging has not been previously shown. In time-lapse imaging of the his2Δ loci, we found increased sister chromatid breathing in the temperature-sensitive psc3-1T mutant at nonpermissive temperature (Figure 5, A–E). These behaviors were similar but slightly more severe than those seen in the mto1Δ mutant. Thus, the mto1Δ mutant and cells treated with MBC exhibit defects similar to those of the psc3 cohesin mutant.

We next tested for Rad21 binding at chromosomal sites by chromatin immunoprecipitation (ChIP). We sampled binding at five loci that have been previously shown to have different levels of cohesin loaded in normal conditions (Figure 6C) (Schmidt et al., 2009): The centromeric dh repeat locus represent cohesin-rich sites. Additional cohesin-associated regions include positions Chr II 2.13 Mb (site d), which is 10 kb apart from the his2 locus, Chr II 1.25 Mb (site a), and Chr II 1.26 Mb (site b). Chr II 0.178 Mb (site c) is a cohesin-poor position. At four of the five chromosomal sites, we found a significant decrease (>50%) in cohesin binding in mto1Δ compared with wild type (Figure 6C). However, at centromeric dh repeats, we found no detectable difference in Rad21 binding between wild-type and mto1Δ cells. Similar results were seen in three independent experiments. Thus, mto1Δ cells exhibit a defect in Rad21 cohesin binding at chromosomal loci on chromosomal arms but retain normal Rad21 binding at centromeres. These results are consistent with live-cell imaging results showing increased sister chromatid breathing at the his2Δ locus (near site d), but not the lys1 locus (near the centromere) (Figure 5 and Supplemental Figures S4–S6).

In summary, these results show that mto1Δ cells have a defect in sister chromatid cohesion that is likely due to abnormal cohesin distribution along chromosomal arms. It is well established that cohesion defects result in defects in DNA repair (Birkenbihl and Subramani, 1992; Sjogren and Nasmyth, 2001; Wu and Yu, 2012), and thus, this cohesin defect provides an explanation for the DNA repair and HR defects in mto1Δ cells.

**DISCUSSION**

In this study, we show that the MT nucleation factor Mto1 is needed for efficient DNA repair, HR, and sister chromatid cohesion at chromosomal arms in interphase fission yeast cells. We find that mto1Δ mutants have defects in cohesin binding and sister chromatid pairing (Figures 5 and 6), which explain at least in part the defects in DNA repair and HR (Figures 3 and 4). We also show that transient inhibition of cytoplasmic MTs by treating interphase cells with MBC produces similar phenotypes in DNA repair factories and sister chromatid pairing (Figure 3 and 5). As Mto1 and MTs are outside the nucleus during interphase, these data lead to an intriguing model that the attachment of cytoplasmic MTs to the nucleus is needed for chromosomal functions such as cohesion and DNA repair. The results presented here provide the initial demonstration of a role for MTs and Mto1 in sister chromatin pairing and cohesin loading or distribution onto chromosomal arms.

How might cytoplasmic MTs and Mto1 affect chromosomal processes? The cytoplasmic MT bundles are physically attached to chromosomes through the NE, the SPB, and other sites containing SUN-KASH complexes (Hagan and Yanagida, 1995; Ding et al., 2004; Chikashige et al., 2006; Hou et al., 2012; Fernandez-Alvarez et al., 2016; Bao et al., 2018). Pushing forces of these MT bundles at the cell tips produce oscillatory movements of the SPB and the chromosomes inside the nucleus (Daga et al., 2006; Daga and Nurse, 2008; Swartz et al., 2014; Schreiner et al., 2015). We show that these chromosomal movements, which affect centromeric chromatin and chromosomal arms, are dependent on MTs and Mto1 (Figure 1). One attractive model is that these MT-dependent forces might facilitate cohesin loading or distribution early in the cell cycle (G1/S) and consequently affect the efficiency of DNA repair (discussed below). An alternate model is that MTs and Mto1 are needed for proper organization or functioning of NE complexes with roles in general chromosomal organization (Swartz et al., 2014). In a third model, Mto1 and perhaps postmitotic MTs function directly at the chromosome inside the nucleus. In mammalian cells, for instance, β-tubulin has been found in complex with Rad51 in the nucleus in response to genotoxic treatments (Lescà et al., 2005; Oakley et al., 2015). However, we did not detect Mto1 inside the nucleus and it does not appear to colocalize with DNA repair factories (Supplemental Figure S2), although our experiments could not rule out, for instance, whether Mto1 associates with the nuclear face of the SPB.

Chromosomal movements driven by the cytoskeleton are likely to play diverse functions in nuclear processes, including dynamic chromosomal organization, homology searching and pairing, detangling of chromosomes, and stimulation of mechanosensitive processes. In meiosis, cytoskeleton-based forces from actin or MTs facilitate recombination and synaptonemal complex formation in fission yeast (Ding et al., 2004), budding yeast (Conrad et al., 2008; Koszul et al., 2008), and C. elegans (Sato et al., 2009). There are a
number of differences, however, between mitotic and meiotic processes. The magnitude of nuclear oscillation during interphase is much smaller compared with meiotic horsetail movement, during which the nucleus rapidly moves between the two ends of the cell (Chikashige et al., 1994; Sjogren and Nasmyth, 2001; Chacón et al., 2016). In addition, meiotic movements are driven by force application to telomeres, while interphase movements are primarily transmitted to centromeres and have less effect on distal chromosomal areas (Chikashige et al., 1994; Swartz et al., 2014). Moreover, the precise nature and molecular control of HR events are quite different between HR in homologous chromosomes in meiosis and in the mitotic cell cycle, wherein the main source of an HR donor sequence for DNA repair is the sister chromatid linked by cohesin complexes to the damaged DNA sequence. Chromosomal movements may also contribute to DNA repair and recombination processes. Microtubules may directly or indirectly contribute to the increased mobility of chromosomal loci observed in response to DNA damage in budding and fission yeast (Dion et al., 2012; Mine-Hattab and Rothstein, 2012; Swartz et al., 2014; Lawrimore et al., 2017). In budding yeast, microtubules are needed for the DNA damage induction of telomere mobilization that contributes to increased mobility of the genome (Lawrimore et al., 2017). While our work was in revision, a report appeared showing that, in budding yeast, abnormally strong MT forces may cause increased DNA repair defects through effects of compressive forces on the nucleus as it migrates through the small bud neck (Estrem and Moore, 2019); as fission yeast nuclei do not encounter a similar bud neck constriction, it is unlikely that they experience these types of compression forces.

Our findings elucidate a new function for the cytoskeleton in DNA repair and recombination: Mto1 and MTs are needed for sister chromatid cohesion. In the absence of Mto1 or the cohesin subunit Psc3, sister loci undergo transient, frequent breathing events in which chromosomal loci move apart and then come back together (Figure 5). Cohesins are loaded at centromeres and other sites by...
| Strain origin           | Strain no. | Genotype                                                                 |
|------------------------|------------|---------------------------------------------------------------------------|
| This study             | 2452       | h90 sid2-tdTomato::NatR lys1::LacOp his7::LacI-GFP-NLS                    |
| This study             | 2822       | h90 mto1Δ::kanMX6 sid2-tdTomato::natMX6 lys1::LacOp his7::LacI-GFP-NLS    |
| Chang lab collection   | 1845       | h- mto1Δ::kanMX6 ura4-D18 leu1-32                                        |
| Daga lab collection    | 2020       | h+ rad3Δ::ura4+ ura4-D18 leu1-32                                         |
| This study             | 2022       | h- rad3Δ::ura4+ mto1Δ::kanMX6 ura4-D18 leu1-32                            |
| Daga lab collection    | 1994       | h- chk1Δ::ura4+ ura4-D18 leu1-32                                         |
| This study             | 1845       | h- mto1Δ::kanMX6 chk1Δ::ura4+ ura4-D18 leu1-32                            |
| Daga lab collection    | 2034       | h cds1Δ::ura4+ ura4-D18 leu1-32                                          |
| Sawin lab              | KS2010     | h+ mto1-9A2 ade6-M216 leu1-32                                            |
| Sawin lab              | KS1957     | h- mto1Δ::ura4+ ade6-M210 leu1-32                                        |
| Daga lab collection    | 1805       | h+ tip1Δ::kanMX leu1-32 ade6-                                          |
| Jia lab                | 2686       | h+ csi1Δ::natMX6 ade6-leu1-32ura4-D18                                   |
| Hiraoka lab            | 2691       | h+ his2 ade6-M216 leu1 ura4-D18 lys1 ima1Δ::hphNT1                        |
| Hiraoka lab            | 2693       | h+ his2 ade6-M216 leu1 ura4-D18 lys1 ish-GFP::kanMX6 man1Δ::LEU2+         |
| Hiraoka lab            | 2694       | h+ his2 ade6-M216 leu1 ura4-D18 lys1 ish-GFP::kanMX6 lem2Δ::ura4+         |
| Hiraoka lab            | 2695       | h+ his2 ade6-M216 leu1 ura4-D18 lys1 ish-GFP::kanMX6 ima1Δ::hphNT1 lem2Δ::ura4 |
| Meister lab            | 1803       | h+ rad52-YFP::kanMX6 leu1-32ura4-D18                                    |
| This study             | 1785       | h- rad52-YFP::kanMX6 mto1Δ::kanMX6 ura4-D18 leu1-32                      |
| This study             | 2712       | h mto2Δ::kanMX6 rad52-YFP::kanMX6 ura4-D18 leu1-32                       |
| This study             | 2340       | h90 sid2-tdTomato::natMX6 his2::kanMX6-ura4+ LacOp his7::lacl-GFP leu1-32|
|                        |            | ade6- leu1-32                                                            |
| This study             | 2341       | h90 mto1Δ::kanMX6 sid2-tdTomato::natMX6his2::kanMX6-ura4+ LacOp his7::lacl-GFP leu1-32 ade6- leu1-32 |
| This study             | 7700       | h psc3-1T-kanMX6 his2::kanMX6-ura4+ LacOp his7::lac Sid2:tomato:kanMX leu1-32|
|                        |            | ade6- leu1-32                                                            |
| This study             | 7701       | h psc3-1T-kanMX6 lys1::kanMX6-ura4+ LacOp his7::lac Sid2:tomato:kanMX leu1-32|
|                        |            | ade6- leu1-32                                                            |
| TH805                  | 2203       | h ade6-M210 leu1-32 ura4-D18 Ch16-MG                                      |
| This study             | 2206       | h mto1Δ::natR ade6-M210 leu1-32 ura4-D18 Ch16-MG                         |
| TH844                  | 2250       | h ade6-M210 leu1-32 ura4-D18 Ch16-MG pREP81X-HO                          |
| This study             | 2253       | h+ mto1Δ::natR ade6-M210 leu1-32 ura4-D18 Ch16-MG pREP81X-HO             |
| TH877                  | 2272       | h rad3Δ::ura4+ ade6-M210 leu1-32 ura4-D18 Ch16-MG pREP81X-HO             |
| This study             | 2287       | h rad3Δ::ura4+ ade6-M210 leu1-32 ura4-D18 Ch16-MG                        |
| This study             | 2397       | h mto1Δ::natR rad3Δ::ura4+ ade6-M210 L- U- Ch16-MG pREP81X-HO             |
| This study             | 2399       | h mto1Δ::natR rad3Δ::ura4+ ade6-M210 L- U- Ch16-MG                       |
| MCW1262                | 2814       | h- ura4-D18 leu1-32 his3-D1 arg3-D4 ade6-M375 int::pUC8/his3+/RTS1 siteA orientation1/ade6-L469 |
| This study             | 3662       | h- mto1Δ ura4-D18 leu1-32 his3-D1 arg3-D4 ade6-M375 int::pUC8/his3+/RTS1 siteA orientation1/ade6-L469 |
| MCW1362                | 2816       | h- swi1Δ ura4-D18 leu1-32 his3-D1 arg3-D4 ade6-M375 int::pUC8/his3+/RTS1 siteA orientation1/ade6-L469 |
| This study             | 3666       | h- mto1Δ swi1Δ ura4-D18 leu1-32 his3-D1 arg3-D4 ade6-M375 int::pUC8/his3+/RTS1 siteA orientation1/ade6-L469 |
| MCW1433                | 2815       | h- ura4-D18 leu1-32 his3-D1 arg3-D4 ade6-M375 int::pUC8/his3+/RTS1 siteA orientation2/ade6-L469 |

TABLE 1: Strains used in this study.
cohesin-loading complexes before DNA replication and may spread laterally along chromosomes to sites of convergent transcription (Tanaka et al., 2001; Lengronne et al., 2004; Schmidt et al., 2009; Peters and Nishiyama, 2012). Cohesin is also recruited during postreplication to sites of double-stranded breaks (Strom et al., 2004). In the mto1 mutant, we found defects in sister chromatid cohesion and a significant reduction of Rad21 cohesin at several cohesin associated sites, even in the absence of DNA damage (Figure 6). This defect may be a consequence of defective cohesin loading, maintenance, or distribution. We speculate that cohesin loading or its dynamic distribution may be a mechanism sensitive process dependent on MT-based forces. Specifically, the oscillatory movements of the SPB linked to centromeres provide tension on the pericentric regions of the chromosomes that could be necessary for efficient cohesin spreading. Whether cohesin defects may account for other cytoskeletal effects on chromosomal processes remains to be tested. Future studies may focus on how mechanical forces regulate cohesin dynamics and function. In general, understanding how chromosomal processes sense and respond to mechanical forces promises to reveal new dimensions in chromosomal biology.

**MATERIALS AND METHODS**

**Strains and media**

Strains used in this study are listed in Table 1. Standard fission yeast techniques and growth media were used (Moreno et al., 1991). Cells were grown in YES media at 25 or 30°C, as indicated. Strains were prepared by crossing and (in the case of the HR assays) by genomic integration.

**Microscopy**

A spinning-disk confocal microscope (Olympus IX81; Roper Scientific) was used. Cells were imaged using 461- and 562-nm lasers. Unless otherwise stated, z-stacks of 10 images with a z-step of 0.4 μm were collected using the 100x objective (UPlanSAPO, NA 1.4), and maximum projections were used for signal quantification. MetaMorph and ImageJ were used for image acquisition and analysis.

**Quantification of Rad52 intensity and dynamics**

To follow Rad52-YFP factory dynamics, we acquired time-lapse images with a time interval of 2 min. We quantified peak nuclear signal of Rad52 over time from maximum-projection images, after subtracting the nuclear background signal. Alternatively, we also quantified background-corrected the total Rad52-YFP signal after thresholding the images to only include the Rad52-YFP factories.

**Microtubule depolymerization**

For depolymerization of MTs, cells were grown to mid-log phase and attached to lectin-coated glass-bottom plates, and freshly prepared 10 μg/ml MBC was added to the cells. Late anaphase cells were marked and either directly imaged for 5–6 h to follow Rad52-GFP dynamics or tracked for 40–110 min and then imaged for 5 min to visualize Rad52 dynamics (Figure 3) or his2 and lys1 LacI-GFP dot “breathing” (Figure 5).

**Generation of DNA damage by laser**

A single 10-ms pulse using an iLas system (Roper Scientific) with a 355-nm laser was targeted at a fixed area of 500 nm of the nuclei. The Rad52-YFP signal was subsequently followed by time-lapse imaging for an additional 25 min.

**Western blotting**

To detect Pk-tagged Rad21, we used anti-Pk monoclonal antibody (kindly provided by I. Hagan, Cancer Research UK, Manchester Institute). Anti-tubulin antibody TAT-1 (kindly provided by Keith Gull, University of Oxford) was used to detect α-tubulin.

**Chromatin immunoprecipitation and real-time PCR**

Strains expressing Rad21-Pk–tagged protein at endogenous levels were fixed with 1% formaldehyde for 20 min, and chromatin was prepared according to Shan et al. (2016). Anti-Pk monoclonal antibody was used for immunoprecipitation, and primers targeting various several genomic regions were used for subsequent real-time PCR amplification of Rad21-bound DNA. Primer sequences (Table 2) correspond to positions 2.13, 0.178, 1.26, and 1.25 Mb on Chr II according to Schmidt et al. (2009). Real-time PCR signals were normalized to wild-type signal.

**Recombination assays**

For HR detection, strains containing mto1 deletion were prepared by crossing (in the case of the Chr16-MG–based system) and by homologous integration (for the intrachromosomal HR substrate). At least five independent clones of each genotype were tested to...
determine recombination frequency. In the RFB system, strains were streaked to single colonies on YES medium, then replica plated on selective ade primers, and individual ade colonies were inoculated into liquid cultures and grown for 8 h at 30°C before plating. Cells were counted and plated in parallel on YES plates to determine total viable cell numbers and on plates lacking adenine to determine the number of adenine prototrophs generated by recombination. The numbers of cells plated were calculated to result in 100–200 growing colonies per plate. Colonies were counted after 3 d at 30°C, and recombination frequency was calculated as in Ahn et al. (2005). At least 300 adenine prototrophs from three biological repeats were counted for each condition.

For quantification of HR with the Chr16-MG substrate (Prudden et al., 2003), cells were grown on ade−, leu− plates containing thiamine and then transferred into liquid Edinburgh minimal media (EMM) lacking leucine for 16 h to induce HO endonuclease expression before plating. G418-resistant cells were replica plated on ade− plates to determine adenine prototrophy. Before calculating recombination frequency (Prudden et al., 2003), we subtracted background rates of Chr16 loss unrelated to HR, which was calculated by plating uninoculated cells on YES and then replica plating them on ade− plates.

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TABLE 2: Primer sequences used for real-time PCR of Rad21-Pk9 ChiP

| A forward | A reverse | C forward | C reverse | D forward | D reverse |
|-----------|-----------|-----------|-----------|-----------|-----------|
| AATTGCAATCCCTGAAGCTGGC | CTTCAAGCTAACTCCGCTATGC | AGAAGTTCAGCTCTCGGAAAA | GTGTAATTTCCGTGAATCGTCA | GGAACAGGTTCCGAGATTCTT | TGGACGCGCTACTCTCAATGG |

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