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Mms1 binds to G-rich regions in Saccharomyces cerevisiae and influences replication and genome stability

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ABSTRACT

The regulation of replication is essential to preserve genome integrity. Mms1 is part of the E3 ubiquitin ligase complex that is linked to replication fork progression. By identifying Mms1 binding sites genomewide in Saccharomyces cerevisiae we connected Mms1 function to genome integrity and replication fork progression at particular G-rich motifs. This motif can form G-quadruplex (G4) structures in vitro. G4 are stable DNA structures that are known to impede replication fork progression. In the absence of Mms1, genome stability is at risk at these G-rich/G4 regions as demonstrated by gross chromosomal rearrangement assays. Mms1 binds throughout the cell cycle to these G-rich/G4 regions and supports the binding of Pif1 DNA helicase. Based on these data we propose a mechanistic model in which Mms1 binds to specific G-rich/G4 motif located on the lagging strand template for DNA replication and supports Pif1 function, DNA replication and genome integrity.

INTRODUCTION

The preservation of genome stability is a major challenge for eukaryotic cells. The faithful duplication of DNA is necessary to maintain the integrity of the genome. During DNA replication, stalled replication forks are a major challenge to genome integrity, because they can cause DNA breakage. Mechanisms that repair or bypass stalled replication forks need to be activated in order to maintain genome stability (reviewed in (1)).

Recent studies indicate that Rtt101, Mms1 and Mms22 form the ubiquitin ligase Rtt101Mms1/Mms22 (2). All three components are required for replication fork progression in cells treated with the alkylating agent methyl methanesulfonate (MMS), which causes replicative stress (2,3). In line with this, it has been shown that Mms1 and Mms22 are required for homologous recombination (HR) at stalled forks, but not at HO endonuclease-induced double-strand-break (DSB) sites (4). Furthermore, it was shown that Mms1 binds near early origins of replication if replication forks are stalled by hydroxyurea (HU) and that this binding is dependent on the other two ligase components, Rtt101 and Mms22 (5).

In addition to the traditional regions (replication fork barriers) where DNA replication stalls (reviewed in (6,7)), DNA secondary structures such as G-quadruplex (G4) structures were shown to influence DNA replication fork progression (reviewed in (8,9)). G4 structures are guanine-rich, four-stranded structures that can form within nucleic acids if a defined nucleotide sequence, called G4 motif, is present (reviewed in (8,10)). Although the existence of G4 structures has long been controversial, increasing amounts of in vitro and in vivo data, such as immunostaining (11–13) and the detection of these structures by small molecules (14–16), now support a model in which G4 structures form in vivo. G4s can positively and/or negatively affect biological processes, such as telomere maintenance, gene expression, epigenetic regulation and DNA replication initiation (reviewed in (8,10,17)). In vitro and in vivo experiments have demonstrated that G4 structures hamper DNA replication and are prone for mutations and deletions (reviewed in (18)). Experiments in Saccharomyces cerevisiae, Caenorhabditis elegans and human cells have shown that formation of these structures leads to genomic instability (19–25). If a G4 structure is formed, helicases or translesion synthesis proteins are required to unwind or bypass these structures, respectively, to permit replication fork progression (reviewed in (26)). Yeast and human DNA helicases such as Pif1 and FANCJ bind to G4 motifs in vivo, prevent replication fork stalling, and support genome stability (19–25). The current
model is that the replication machinery slows down when it encounters a G4 structure and that helicases (e.g. Pif1 in yeast) unwind them efficiently (21). In this case, the replication fork has to restart behind the obstacle leaving a gap behind (reviewed in (26)).

Currently, it is unclear how the Rtt101Mms1/Mms22 ligase recognizes stalled forks and if it acts at all stalled forks or only at specific loci. Here, we show that Mms1 binds to specific G-rich motifs located on the lagging strand template for DNA replication in the genome. Replication forks stall at these G-rich motifs in mms1 cells, and this G-rich motif has the potential to form G4 structures in vitro. Additional genetic assays show that Mms1 binding sites promote gross chromosomal rearrangements in the absence of Mms1. Chromatin immunoprecipitation (ChIP) experiments indicate that the observed replication and genome stability defects are due to lower Pif1 binding to these G-rich/G4 motifs in the absence of Mms1. Our data suggest a new mechanistic model of how Mms1 supports DNA replication at specific G-rich sequences that could form G4 structures.

MATERIALS AND METHODS

Strains, constructs and media

All yeast strains are listed in Supplementary data 1. All experimental strains are derivatives of the RAD5+ version of W303 (R. Rothstein) or YPH background (27). Deletions eliminated entire ORFs and were created according to (27). Epitope tagging to generate Mms1-Myc13, Pif1-Myc13 and DNA Pol2-Myc13 was carried out as described (28). Myc-tagged proteins were expressed from endogenous loci and promoters. The pif1-m2 mutant was created as described (29).

In vitro folding of G4

The folding was performed according to (30). To confirm G4 formation the samples were controlled by circular dichroism (CD) measurements.

Circular dichroism

15–20 μg DNA was subjected to CD measurements at 25°C using a Jasco J-810 spectropolarimeter (Jasco). The parameters were: continuous scanning mode (200–350 nm), accumulation 10, scanning speed 100 nm/min, response 0.25 s, band width 2 nm, and a data pitch of 0.2 nm.

Western analysis

Proteins for western analysis were isolated according to a protocol by Foiani (31). Western analysis was performed according to standard protocols. The primary antibodies for c-Myc (Clontech) and Hsp60 (Abcam) were used according to the manufacturer’s protocol. Hsp60 served as a reference protein. As secondary antibody, we used a HRP-coupled antibody (Santa Cruz Biotechnology). Proteins were detected by chemiluminescence. Quantification was performed using Image Lab (BioRad).

Gross chromosomal rearrangement assay

The gross chromosomal rearrangement (GCR) assay was performed as published (32) with minor modifications. Briefly, seven colonies per strain were grown for 48 h. Cells were plated on two different plates: YEPD as a reference plate and on a FOA/CAN selective plate. After incubation colonies were counted on both plates and the GCR rate was determined via fluctuation analysis using FALCOR and the MSS maximum likelihood method (33).

Chromatin immunoprecipitation (ChIP)

ChIP of asynchronous and synchronous samples was essentially performed as described (21). For ChIP-seq the average length was 200 bp using a M220Focused-ultrasonicator (Covaris) and for conventional ChIP the DNA was sheared to an average length of 250 bp using a Branson sonifier W250-D (50% amplitude, 50% duty cycle, 5 × 5 pulses) (Supplementary data 2A–C). The applied parameters for Covaris were 75 W, 25 duty and 200 cycles/burst for 20 min. C-Myc antibody was obtained from Clontech. Primers used for qPCR (Cycler and SYBR Green, Biorad) are listed in Supplementary data 3. For genome-wide sequencing, DNA was treated according to manufacturer’s instructions (NEBNext ChIP-seq Library Prep Master Mix Set for Illumina, NEB) and submitted to deep sequencing (Illumina Nextseq500 sequencer). Obtained sequence reads were aligned to the yeast reference genome (sacCer3) with BOWTIE (34). After alignment, the number of reads was normalized to the sample with the lowest number of reads. Binding regions were identified by using the program ‘Model-based Analysis for ChIP-Seq (MACS 2.0)’ with default settings, -no model option, and -extsize 180, which correlates to the minimal fragment size (35). See Supplementary data 4 for all Mms1 peaks. The ChIP input sample was used as a control. MEME-based motif elicitation was used to identify a consensus motif (36) within the FASTA file from the binding regions identified by MACS 2.0. G4 motifs were identified using a script previously published (37). Overlap of binding sites and qPCR regions with G4 motifs and genes was determined using bedtools’ window command (38). A window size of 400 bp was used when evaluating overlap of binding regions and 500 bp when evaluating qPCR regions. To determine the overlap with genomic features we took the annotations from saccharomyces_cerevisiae_R64-2-1.20150113.gff. To examine if the qPCR regions contain G4 motifs on the leading and/or lagging strand, we identified the closest ARS to the G4 motifs using bedtools and determined strand specificity.

Arrest of yeast cells

Cells were arrested in G1 phase according to (39) and in S and G2 phase according to (40). FACS analysis to confirm cell cycle arrest was performed as described (39) using a FACSCanto II (BD).

Endogenous mutation of G4 using Cre-Lox

G4 Chr VI (253592–255049) was mutated using Cre-Lox recombination. The mutated G4 motif was synthesized.
In order to get more insight into the binding properties of Mms1, and especially the potential contribution of G4 motifs, we performed ChIP-qPCR experiments using the same endogenous Myc-tagged Mms1 strain as for ChIP-seq. For the qPCR, we chose 12 different regions in the yeast genome. Out of these 12 sites, three were Mms1 binding regions and seven exhibited the presence of a G4 tract2. In addition, we selected two negative control regions based on the ChIP-seq data. For this and all subsequent ChIP experiments, IP values of each experiment were normalized to its input values. We considered it a positive binding if Mms1 levels were at least three times higher than those of the untagged control. In concordance with our earlier results, Mms1 bound three regions selected by ChIP-seq (Chr VIIBM, XBR, XI BM) with two regions harboring the G4 tract2 motif (Figure 1A). Strikingly, six out of seven regions with a G4 tract2 (Chr VI, IX, XIa, X Ib, XIII, XV) were also identified as Mms1 BR. Mms1 did not significantly bind to regions on Chr I, Chr XIV and an additional region from Chr XIII. Due to the lack of significant Mms1 binding, we treated these three regions as negative controls (NC) in the remaining analyses. To elucidate why Mms1 binds to specific regions, we further analyzed the ChIP-seq data. This analysis revealed that all regions which are bound by Mms1, using ChIP and qPCR, harbor one or multiple G4 tract2 or even G4 tract3 motifs (see Supplementary data 6 for details). However, the NC regions Chr INC and Chr XIII NC also harbor a G4 tract2 and a G4 tract3 motif, respectively. Considering the distance to the next ARS and the strand location of G4 tract2 motifs, we identified that all G-rich/G4 motifs bound by Mms1 are located on the DNA strand replicated by the lagging strand machinery and have a mean loop length smaller than eight nucleotides. The NC regions have either no such G-rich/G4 motifs or those are not located on the lagging strand template (Supplementary data 6). In summary, ChIP-seq and ChIP-qPCR revealed that Mms1 only binds to G-rich regions, if they harbor at least one G4 tract2 located on the lagging strand template for DNA replication. For simplicity, all regions that show significant binding of Mms1 either by ChIP-seq and/or ChIP-qPCR will be named in the text and figures as a BR.

To determine whether our identified G-rich binding regions can form G4 structures in vitro, we performed CD measurements on three of the 71 binding regions of Mms1. By these measurements, we confirmed that the regions from Chr VII BR, XIaBR, and X V BR (all harboring a G4 tract2 with a mean loop length < 8 nt, lagging strand template) folded into G4 structures in vitro (Figure 1C), whereas the regions from Chr I NC and XIII NC (mean loop length > 8 nt, lagging strand template) did not (Figure 1D, Supplementary data 7). These results show that many regions bound by Mms1, and also those harboring G4 tract2, have the ability to form G4 structures.

Mms1 binds in all cell cycle phases and independently of Rtt101 and Msms2

In order to determine the spatiotemporal function of Mms1, we determined in which cell cycle phase Mms1 binds to its target regions. First, we measured Mms1 levels at different cell cycle phases. Yeast cells expressing Myc-tagged
Mms1 were arrested in G1, S, and G2 phase using different reagents (α-factor, HU, and nocodazole, respectively). FACS analysis was performed to confirm cell cycle arrest (Figure 2A). Western blot analysis was performed to monitor Mms1 protein levels (Figure 2B, Supplementary data 8A). Although Mms1 protein levels peak in G1 phase (>5-fold more Mms1) Mms1 can be detected in all cell cycle phases (Figure 2B).

To address the question when Mms1 binds to sites containing G4 tract2 motifs, we performed ChIP and qPCR of Myc-tagged Mms1 cells arrested in G1, S, and G2 phase. The arrests were confirmed by FACS analysis (Supplementary data 8B). qPCR at Mms1 target regions Chr VI_{BR}, IX_{BR}, X_{BR}, XI_{BR}, XII_{BR}, XIII_{BR} and XV_{BR} revealed that Mms1 binds equally well throughout the cell cycle to all tested BR (Figure 2C).

Previous work has shown that Mms1 interacts with Rtt101 and either Crt10 or Mms22 (2). So far the only process Crt10 was shown to be involved in is ribonucleotide reductase (RNR) gene expression (47), while Mms22 and Mms1 together with Rtt101 (Rtt101 Mms1/Mms22) promote replication fork progression and HR at stalled replication forks (2–5). Due to the distinct functions of these two complexes, we hypothesized that the Rtt101^{Mms1/Mms22} complex could be recruited to our identified DNA target regions. In previous work, it was postulated that, in the Rtt101^{Mms1/Mms22} complex, Mms22 is the DNA interacting protein (2). Therefore, we first asked if Mms1 binding is dependent on Rtt101 or Mms22. We analyzed Mms1 binding in the absence of Mms22, Rtt101, and both proteins (MMS22/RTT101 double deletion) by ChIP-qPCR experiments. ChIP and qPCR analysis revealed that binding of Mms1 did not decrease in the rtt101 cells, but interestingly binding was significantly enriched in mms22 deficient cells (Figure 3A). This enhanced binding was observed at all tested regions including negative control regions. At Chr VI_{BR}, VII_{BR}, IX_{BR}, XI_{BR}, XIII_{BR}, I_{NC}, XIII_{NC} and XIV_{NC} we observed at least a 2-fold enriched binding of Mms1. In the mms22 rtt101 double mutant, Mms1 binding was similar to wild type or rtt101 cells (Supplementary data 8C).

To evaluate whether the observed difference in binding was due to changes in the total protein level of Mms1, we performed western blot analysis of Myc-tagged Mms1 and quantified the amount of Mms1 in wild type, rtt101, and mms22 cells using Hsp60 as a reference protein. The ab-
Figure 2. Mms1 protein levels are highest in G1 phase and Mms1 binds throughout cell cycle. (A) FACS analysis of cells arrested in G1, S or G2 phase. Cells were arrested in G1 by treatment with α-factor, in S phase by HU and in G2 by nocodazole. (B) Western blot analysis of Myc-tagged Mms1 protein levels in G1, S and G2 phase. Level of Mms1 was quantified using Hsp60 as a reference protein. Shown are mean Myc-tagged Mms1 levels normalized to Hsp60 ± SD. N = 3 biological replicates. See Supplementary data 8A for the gel. (C) ChIP and qPCR analysis of Mms1-Myc to seven BR in G1-, S- and G2-phase. Plotted are IP/input values as means ± SD. N ≥ 3 biological replicates. In most cases Mms1 binds similar in G1, S and G2 phase. Statistical significance compared to cells arrested in G1 phase was determined by Student’s t-test. * P < 0.05.

Figure 3. Mms1 binds independently of Rtt101 and Mms22, supports Pif1 binding at G4 motifs and by this promotes DNA replication. (A) ChIP and qPCR analysis of Mms1-Myc at seven BR and two NC. Binding of Mms1 was monitored in wild type (light), rtt101 (grey) and mms22 (dark) cells. Statistical significance compared to Myc-tagged Mms1 wild type cells. For details on regions see Supplementary data 3. (B) Replication fork progression was analyzed by detected DNA Pol2 binding levels at Mms1 binding sites. ChIP and qPCR of DNA Pol2 binding in wild type and mms1 cells was performed at five BR and two NC. Statistical significance compared to Myc-tagged DNA Pol2 wild type cells. (C) Binding of Pif1 DNA helicase was analyzed at four Mms1 binding regions (BR) in wild type and mms1 cells. As control for Pif1 binding we used two known Pif1 binding sites, the replication fork barrier at the rDNA (rDNA) (54) and telomere VI-R (tel) (53) as well as one Pif1 independent site (tRNA). As done previously (53) IP/input values are compared to IP/input values of ARO1 where no Pif1 binds. Here, fold enrichment over ARO1 was plotted as mean value ± SD. For all ChIP, N ≥ 3 biological replicates. Statistical significance was determined by Student’s t-test. * P < 0.05, ** P < 0.01, *** P < 0.001.

The presence of Mms22 did not influence the abundance of Mms1, but Mms1 levels significantly increased in rtt101 cells (~4-fold). (Supplementary data 8D,E). These data (ChIP and Western analysis) suggested that Rtt101 impacts Mms1 protein levels, but that it does not impact Mms1 binding to G-rich target regions. In Summary, these results suggest that Mms1 does not require Rtt101 or Mms22 for binding to G-rich/G4 sites located at the lagging strand template of DNA replication. Furthermore, it raises the possibility that Mms1 binds easier/better to G-rich/G4 regions without Mms22. If Mms1 acts as part of the E3 ligase, Mms22 is the preferred DNA binding protein as previously suggested (2).
Mms1 promotes progression of the stalled DNA replication fork

Previous work has shown that DNA replication is slowed in *mms1* cells after MMS treatment (2). Due to this strong connection between Mms1 function and DNA replication we further analyzed the contribution of Mms1 to replication fork progression. We aimed to investigate if replication fork progression is affected at the G-rich motifs identified as Mms1 binding sites in *mms1* cells. DNA Polymerase 2 (DNA Pol2) is the catalytic subunit of DNA polymerase ε (48,49). It is assumed that regions with high DNA Pol2 levels are sites where DNA replication is slowed or stalled (21,50). Therefore, we performed ChIP-qPCR on asynchronous wild type and *mms1* cultures that express endogenous Myc-tagged DNA Pol2. Binding of DNA Pol2 was monitored at five Mms1 binding regions (Chr V_browser BR, X_browser BR, X_browser BR, X_browser I, and X_browser BR) as well as two non-Mms1 binding regions (Chr XIII_browser NC, XV_browser NC) (Figure 3B). Binding of DNA Pol2 was significantly (P-value < 0.05, 1.8- to 2.5-fold) enriched at Chr V_browser BR, X_browser BR, X_browser BR and X_browser BR in *mms1* cells. DNA Pol2 protein levels did not change in *mms1* cells (Supplementary data 9A and B). Furthermore, we tested if DNA Pol2 binding was also elevated in *rtt101*, *mms22* and *rtt101 mms22* cells at these G-rich sites. ChIP experiments showed no elevated DNA Pol2 binding in these single or double mutants (Supplementary data 9C). In summary, elevated DNA Pol2 level are observed in the absence of Mms1 at G-rich motifs indicating that replication fork movement is slowed at these regions if Mms1 is absent. At the control regions (Chr XIII_browser NC and XIV_browser NC) DNA Pol2 occupancy was independent of Mms1, correlating with no Mms1 binding at these sites (see Figure 1B). Furthermore, these results further strengthen the point that Mms1 acts independently of Rtt101 and Mms22 at such sites.

Mms1 does not recruit Mre11 to its binding sites

Previous studies revealed that Mms1 is required for HR at stalled replication forks (4). Recently, it has been shown that HR proteins, such as Rad51 and BRCA1, regulate HR at G4 structures during DNA replication (51). Additionally, work in yeast showed that breakage near a stalled fork induces recombination (19,21). In the next set of experiments, we wanted to determine if Mms1 binding at its target regions also causes recruitment of HR factors due to stalled forks. We examined if Mms1 recruits Mre11, a component of the MRX complex involved in HR (reviewed in (52)). We performed ChIP of asynchronous yeast cells expressing endogenous Myc-tagged Mre11 in wild type and *mms1* cells. Occupancy of Mre11 at Mms1 binding sites (from Figure 1B) was determined by qPCR, and the IP/input values were calculated (Supplementary data 10). We found that the association of Mre11 to all tested Mms1 binding regions was not dependent on Mms1, suggesting that Mms1 does not recruit HR factors to its target sites.

Mms1 and the Pif1 helicase act together at G-rich/G4 motifs

The Pif1 DNA helicase binds at the end of S phase to G4 motifs and supports DNA replication and promotes genome stability (19,21). The association of DNA Pol2 to G4 motifs is greatly increased in *pif1-m2* mutants (21). In the next set of experiments, we tested if Mms1 and Pif1 act together at G4 structures. We re-analyzed genome-wide Pif1 binding sites (21) and checked for overlap with Mms1 peaks (Supplementary data 4). We identified 38 Mms1 binding regions that overlap Pif1 binding sites (Supplementary data 11). This is significantly more overlap than expected if these sites were randomly distributed across the genome (*P* = 0.001).

Replication fork progression is impeded at G4 sites in the absence of Pif1. Therefore, we tested whether Mms1 associates more strongly to G4 sites in *pif1-m2* cells. In *pif1-m2* mutants, expression of the nuclear isoform of Pif1 is disrupted, but the mitochondrial isoform of Pif1 is still expressed (29). We found that Mms1 binding did not change in *pif1-m2* compared to wild type cells at all tested regions (Supplementary data 12A).

Previous work revealed that Pif1 binds at the end of S phase to G4 motifs (21) and that Mms1 binds throughout the cell cycle to specific G4_tract2 motifs (Figure 2). We speculate that Mms1 binding supports Pif1 function at those sites. In this model, we expect reduced Pif1 binding to target regions in the absence of Mms1. In three independent ChIP experiments, we observed that Pif1 binding was more than two-fold reduced in the absence of Mms1 at G4_tract2 motifs (these sites were previously also identified as Pif1 binding sites (21)) (Figure 3C). In contrast, Pif1 binding to three different control regions, telomere VI-R (Tel-VI-R (53)), the replication fork barrier at the rDNA repeat (rDNA (54)), and a tRNA gene (no Pif1 binding site (21)), was not altered. Pif1 levels in the cell did not change upon *MMS1* deletion (Supplementary data 12B and C).

Mms1 binds to G4 structures

Our data led us to the hypothesis that Mms1 binds to G4 structures rather than to G-rich motifs. To determine whether Mms1 binds G-rich regions or G4 structures itself, we performed an affinity purification approach using biotinylated G4 structures or biotinylated control DNA sequences as bait. We used four G4 structures (Chr IG4_tract2, Chr IXG4_tract3, Chr XIIIG4_tract2, Oxy2G4_tract4) as well as three controls (G-rich, non-G-rich, and a mutated G4 motif (95% identical to the G4 motif), none of which can form a G4 structure). The biotinylated oligodeoxynucleotides containing G4 motifs were folded into G4 structures. Control oligodeoxynucleotides, which cannot form a G4 structure, were treated in parallel. Formation of G4 structures as well as non-folding of the control sequences was confirmed by CD. Folded G4s and controls were incubated with total yeast protein lysate (from *pif1-m2* cells: to reduce G4 unwinding potential in the lysate) in which Mms1 was endogenously tagged with a Myc tag. After wash steps and incubation with streptavidin-coupled beads, we isolated bound proteins. Western blot analysis revealed that Mms1 bound to all four G4 oligodeoxynucleotides but not to the three control oligodeoxynucleotides. This experiment showed that Mms1 did not bind to G-rich sequences in general or non-G-rich linear DNA (Supplementary data...
13 lane 1–3), but that it specifically binds to G4 structures (Supplementary data 13, lanes 4–7).

To further support the finding that Mms1 acts on G4 structures rather than just G-rich sequences, we mutated the G4 motif from Chr VI (IV\textsubscript{G4}) (GGGGCACACGTGCGGGAGTTCAAGGGGGCAGAATAAGTGGGCTAGCGGGG) by Cre-Lox recombination to abolish its G4-forming potential (Chr VI\textsubscript{G4mut}) (GGGGCACACGTGCGGGAGTTCAAGGGGGCAGAATAAGTGGGCTAGCGGGG). We endogenously tagged Mms1 in this strain. ChIP and qPCR analyses revealed that Mms1 binding is lost once the G4 motif is mutated (Figure 4A). The binding of Mms1 to non-mutated G4 motif (X\textsubscript{G4}), in the same strain background was not altered (Figure 4A). This indicates that the G-rich nature of the motif is not responsible for Mms1 binding, but the G4 structure itself. To reveal if DNA replication pausing also depends on G4 motif at these sites, we endogenously tagged DNA Pol2 in the strain harboring the mutated G4 on Chr VI (VIG\textsubscript{G4mut}). ChIP and qPCR analyses were performed in wild type and mms1 cells. As observed in Figure 3B, DNA Pol2 was enriched in mms1 at VIG\textsubscript{G4} and X\textsubscript{G4}. Interestingly, DNA Pol2 binding was no longer enriched at VIG\textsubscript{G4mut} in mms1 cells upon mutation of the G4 motif (Figure 4B, black). Binding of DNA Pol2 in MMS1 cells did not change upon G4 mutation. These results further strengthen the hypothesis that Mms1 binds to G4 structures, including sequences with the G\textsubscript{4tract} motif as well as the conventional G\textsubscript{4tract}.

Mms1 prevents genome instability

Our findings revealed that Mms1 binds to G-rich/G4 regions (Figure 1A and B) and supports Pif1 function (Figure 3C) at such sites. Previously, it was shown that, in the absence of Pif1, DNA replication is not only impeded, but genome stability is also challenged (19–23). We tested if genome instability at G4 motifs is increased in mms1 cells, in which Pif1 binding is reduced, using a previously published GCR assay (20,32). In this assay, we monitor G-rich versus G4 induced genome instability quantitatively. It allows us to measure complex genome rearrangement by simultaneous selection against two counter-selectable markers (URA3, CAN1) (20,55). We created four experimental strains in which a G\textsubscript{4tract} motif from Chr I (G4-LEU2), a G-rich region from Chr I (GR-LEU2), or a non-G-rich region from Chr VII (NG-LEU2) was inserted into the yeast genome on the left arm of Chr V, replacing the non-essential PRB1 gene (Figure 4C, see Supplementary data 14A for detailed information on regions and sequences). Because the insertions are done using a LEU2 gene, which harbors many G\textsubscript{4tract} on the template for lagging strand replication, that are Mms1 binding regions based on our ChIP and ChIP-seq data, we also inserted only the LEU2 gene into this region. The two counter-selectable markers (URA3 and CAN1) are located downstream of the PRB1 locus (Figure 4C). If the inserted sequence induces genomic instability, the markers are lost, and cells can grow on selective media. By counting the colonies on selective media compared to those on rich media plates, the GCR rate can be determined via fluctuation analysis (33). The GCR rate of the wild type cells with-
out an insert was \(\sim 0.1 \times 10^{-9}\) events per generation, as published (56). The observed GCR rates, normalized to the rate of wild type without insert, are depicted in Figure 4D. None of the inserts induced a significant increase of the GCR rate in wild type cells. Interestingly, even without any insertion, the deletion of MMS1 caused an increased (3.3-fold) GCR rate over wild type without insert (Figure 4D). Normalizing these GCR rates by the rate obtained from mms1 without an insert, we measured 2.5- to 3.8-fold higher GCR rates using GR-LEU2, NG-LEU2, and the LEU2 gene as an insert. However, if a G4-LEU2 insert is present in mms1 cells, the GCR rate increased \(\sim 19\)-fold compared to wild type cells (Figure 4D), which is significantly \((P = 0.003)\) higher than in the presence of the other inserts. As mentioned above, the LEU2 gene contains a G4 tract2 on the lagging strand template for replication and therefore increased GCR rates were expected for all inserts. Strikingly, insertion of an additional G4 motif resulted in an even higher GCR rate in mms1 cells (see Supplementary data 14B for data normalized against LEU2). Similarly, increased G4-dependent genome instability (GCR rates in pif1-m2 are without insert 76-fold and with a G4 insert 200-fold) was also detected using the same GCR assay in pif1-m2 cells (see (20)). These data suggest that loss of Mms1 causes genome instability and that this effect is even more severe if G4 tract3 motifs are present.

DISCUSSION

In this study, we investigated the functional role of the ubiquitin ligase component Mms1 at G-rich/G4 regions during DNA replication. We demonstrated that Mms1 binds to G-rich regions, and more specifically to G4 tract2 motifs on the template for lagging strand replication (Figure 1A, B, Supplementary data 6). CD experiments showed that a subset of the identified G4 tract2 motifs form G4 structures in vitro (Figure 1C). Additionally, we observed that replication slows in mms1 cells at such sites, contributing to increased genome instability (Figures 3 and 4). Using ChIP experiments, we further revealed that Mms1 binding to G-rich/G4 sites supports Pif1 DNA helicase function at G4s (Figure 3C).

Based on our experiments and published data, we propose a new mechanistic model for G4 function during replication (Figure 5). Mms1 binds throughout the cell cycle to specific G-rich regions with a potential to form G4 structures. Interestingly, Mms1 does not bind to all G4 motifs it is specific for G4 motifs located on the lagging strand template. It is important to note that we cannot exclude the possibility that this binding is indirect. Regardless, once Mms1 binds to these G-rich/G4 regions it supports Pif1 binding at the end of S phase, allowing Pif1 to unwind structured DNA. Hence, replication fork progression and genome integrity are maintained. In the absence of Mms1, Pif1 binding is diminished, which results in replication fork pausing and genome instability. We present six results that support this model.

First, the consensus Mms1 binding motif is more G-rich than expected \((P < 0.001)\) and most, 61 of 71 binding sites, harbor a G4 tract2 motif located on the lagging strand template. Using CD analysis, we showed that a subset of Mms1 binding sites that harbor a G4 tract2 can fold into stable G4s in vitro (Figure 1C). Second, Mms1 binding to G4 tract2, as well as G4 tract3 motifs with a potential to form G4s was shown by ChiP and qPCR (Figure 1B). The low abundance of Mms1 (57) and the unpredictable nature of G4 structure formation in vivo (8) supports the notion that Mms1 likely binds to more G4 sites in vivo than identified by ChiP-seq. This hypothesis is supported by ChiP-qPCR experiments that showed robust Mms1 binding to G4 tract2 and G4 tract3 motifs that were not discovered by ChiP-seq (Figure 1B, Chr V I B R, IX B R, XI I B R, XI I I B R, XII I B R). Third, we showed that Mms1 binding sites overlap significantly with Pif1 binding regions \((P = 0.001,\) Supplementary data 11). Furthermore, in the absence of Mms1, less Pif1 binds to G-rich/G4 motifs (Figure 3C). This result indicates that either Mms1 binding supports Pif1 function at G4 structures or that Mms1 itself stabilizes G4 structures that are in need of unwinding by Pif1 helicase. From the current state of knowledge, we exclude the second scenario because without Mms1 more genome instability is observed (GCR assay, Figure 4D). Interestingly, Mms1 has a preference for G-rich/G4 tract2 on the lagging strand template whereas Pif1 binds to G4 tract3 motifs with no preference for leading and the lagging strand templates (21). This suggests that G4 structure regulation is not as straightforward as initially thought and that other factors or proteins will be found in the future that support Pif1 function at other G4 structures.

Fourth, in the absence of Mms1, DNA replication slows at G4/G-rich motifs (Figure 3B) and genome instability increases (Figure 4D). This is in line with the fact that without Mms1 less Pif1 binds to G4 structures (Figure 3C), consequently G4 structures are not unwound, thus DNA repli-
cation is impeded and genome instability is observed using a yeast genetic assay (GCR). Interestingly, if a G4tract3 motif was inserted in addition to the G4tract2, the GCR rate was further increased (Figure 4D). Previously, it was shown that in the absence of Pif1, GCR rate increases 2.5-fold upon addition of G4 motifs (increase from 76-fold in without insert to 200-fold with G4 insert) (20). The higher GCR rates in pif-m2 compared to the GCR rate in mms1 can be explained by the observation that without Mms1 Pif1 binding is reduced at G4 structures but not abolished and some Pif1 still binds to G4s. Additionally, without Pif1 the GCR rate (without insert) is already over 35-fold higher than in mms1 cells. This might be due to the fact that Pif1 is a multifunctional helicase involved in multiple steps important for genome stability (58). Nevertheless, the GCR results agree with the finding that more stable G4 structures, harboring a G4tract3, are more challenging for genome stability (Piazza et al., 2015) than G4tract2.

Fifth, we show that at G4 motifs located on the lagging strand template Mms1 is the core component which recognizes the G4 motif and supports Pif1 function. This function is independent of the E3 ligase complex (Figures 1 and 3A and Supplementary data 8). ChIP experiments showed that Mms1 binding is dependent on neither Mms22 nor Rtt101, which suggests that Mms1 is either binding the region itself or via a so far unknown interaction partner. However, we did not identify other Mms1 interaction partners by co-IP followed by mass spectrometry. In two independent approaches, we could only identify Rtt101 as a significant and relevant binding partner (Supplementary data 15). We think it is unlikely that Mms22 or Rtt101 support Mms1 binding at such sites, because deletion of one or both proteins did not result in reduced ChIP signals of Mms1-Myc (Figure 3A, Supplementary data 8) nor in replication fork pausing (Supplementary data 9). This conclusion is in contrast to previous publications where it was suggested that Mms1 binds via Mms22 to DNA (2,5,59). This difference could be explained by different experimental set ups and the fact that we look at a very specific target (G4 motifs on the lagging strand) of Mms1.

Sixth, our results show for the first time that Mms1 binds to G4 structures rather than to G-rich DNA regions (Supplementary data 13). Mms1 binding is even lost upon disruption of the G4 forming potential (Figure 4A). Consequently, DNA replication is independent of Mms1, and thus Pif1, if the G4 motif is mutated (Figure 4B). These results and the observed finding that Mms1 binds throughout the cell cycle to these G4 motifs indicate that G4 structures form already before the onset of S phase. This argument is supported by the fact that G4 structures can form in duplex DNA in vivo (14), but also raises the question how and why G4 structures form prior to S phase.

Previous work in yeast using a human minisatellite sequence had shown that only G4s with very short loops challenge genome stability. In this manuscript, we show that G4s with either longer loops or shorter G-tracts located on the laggings strand template are bound by Mms1 and that Mms1 assist DNA replication and genome stability by supporting Pif1 function here. This is in agreement with other publications showing that in the absence of helicases (e.g. Pif1, FANCJ, Dog-1) genome stability is at risk at G4 motifs (20,21,23–25,60,61). Our data strengthen the hypothesis that G4 regulation is done by specialized helicases and that a specific helicase regulates G4 unwinding at a specific loci or during a specific mechanism (e.g. Pif1 during DNA replication). In summary, the data presented here highlights the still mysterious nature of G4 structure, function and regulation in vivo, and also calls for further future experiments. Our study provides new insights into G4 structure regulation during replication. In yeast it was shown that Pif1 DNA helicase is responsible for genome stability at G4 structures. We provide data that, depending on the location of the G4 and maybe other unknown factors, G4 structures are unwound/unfolded with the support of other proteins. We revealed, that Mms1 supports Pif1 binding to G4 structures located on the lagging strand template for replication, leading to the question, which other proteins support Pif1 function at other G4 loci.

ACCESSION NUMBER
The accession numbers for the ChIPseq data reported in this paper are NCBI GEO: GSE98524.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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