

DNA polymerases δ and λ cooperate in repairing double-strand breaks by microhomology-mediated end-joining in *Saccharomyces cerevisiae*

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Maintenance of genome stability is carried out by a suite of DNA repair pathways that ensure the repair of damaged DNA and faithful replication of the genome. Of particular importance are the repair pathways, which respond to DNA double-strand breaks (DSBs), and how the efficiency of repair is influenced by sequence homology. In this study, we developed a genetic assay in diploid Saccharomyces cerevisiae cells to analyze DSBs requiring microhomologies for repair, known as microhomology-mediated end-joining (MMEJ). MMEJ repair efficiency increased concomitant with microhomology length and decreased upon introduction of mismatches. The central proteins in homologous recombination (HR), Rad52 and Rad51, suppressed MMEJ in this system, suggesting a competition between HR and MMEJ for the repair of a DSB. Importantly, we found that DNA polymerase delta (Pol δ) is critical for MMEJ, independent of microhomology length and base-pairing continuity. MMEJ recombinants showed evidence that Pol δ proofreading function is active during MMEJ-mediated DSB repair. Furthermore, mutations in Pol δ and DNA polymerase 4 (Pol λ), the DNA polymerase previously implicated in MMEJ, cause a synergistic decrease in MMEJ repair. Pol λ showed faster kinetics associating with MMEJ substrates following DSB induction than Pol δ . The association of Pol δ depended on RAD1, which encodes the flap endonuclease needed to cleave MMEJ intermediates before DNA synthesis. Moreover, Pol δ recruitment was diminished in cells lacking Pol λ . These data suggest cooperative involvement of both polymerases in MMEJ.

DNA repair | genome stability | translocation

NA double-strand breaks (DSBs) are toxic lesions that can be repaired by two major pathways in eukaryotes: nonhomologous end-joining (NHEJ) and homologous recombination (HR) (1). Although HR repairs DSBs in a template-dependent, high-fidelity manner, NHEJ functions to ligate DSB ends together using no or very short (1-4 bp) homology. Recently, a new pathway was identified in eukaryotes, which uses microhomologies (MHs) to repair a DSB and does not require the central proteins used in HR (Rad51, Rad52) or NHEJ (Ku70–Ku80) (2–5). In mammalian cells, this pathway of repair is known as alternative end-joining (Alt-EJ) and is often but not always associated with MHs, whereas in budding yeast, the commensurate pathway, MH-mediated end-joining (MMEJ), will typically use 5–25 bp of MH (6, 7). These pathways are associated with genomic rearrangements, and cancer genomes show evidence of MH-mediated rearrangements (8–12). In addition, eukaryotic genomes contain many dispersed repetitive elements that can lead to genome rearrangements when recombination occurs between them (13-16). Therefore, controlling DSB repair in the human genome, which features a variety of repeats, is especially important given the fact that recombination between repetitive elements has been implicated in genomic instability associated with disease (17-20).

The original characterization of Alt-EJ in mammalian cells suggested it did not represent a significant DNA repair pathway and only operated in the absence of functional HR and NHEJ pathways. More recent analyses demonstrate a physiological role of Alt-EJ during DNA repair in the presence of active HR and

NHEJ pathways (2, 12, 21, 22). Furthermore, examination of I-SceI-induced translocation junctions in mammalian cells revealed the frequent presence of MHs (23, 24). NHEJ-deficient and p53-null mice develop pro-B-cell lymphomas, and non-reciprocal translocations characterized by small MHs are found at their break point junctions (25–28). Similarly, in human cancers, many translocation break point junctions contain MHs, suggesting a role for Alt-EJ in cancer development (29–31) and resistance to chemotherapy and genetic disease (32–36). Hence, the presence of many short repetitive sequences in the human genome is likely to increase rearrangements mediated by MHs following the creation of a DSB.

MMEJ is a distinct DSB repair pathway that operates in the presence of functional NHEJ and HR pathways (10, 37). The genetic requirements of MMEJ are being studied in the model eukaryote *Saccharomyces cerevisiae* and involve components traditionally considered specific to the NHEJ (Pol λ) and HR (Rad1–Rad10, Rad59, and Mre11–Rad50–Xrs2) pathways (4, 5, 10, 38). Although being clearly independent of the central NHEJ factor Ku70–Ku80 heterodimer (10, 37), the involvement of the key HR factor Rad52 in MMEJ remains uncertain. It has been reported that Rad52 is required for MMEJ repair (4, 10, 38), whereas in another assay system Rad52 suppresses MMEJ repair (37). More recently, it has been proposed that the replication

Significance

The human genome contains many segments of short repetitive DNA, known as microhomologies, which are potential sites for the types of rearrangements found in many different types of cancer. Therefore, understanding how microhomologies within DNA can initiate chromosome rearrangements and the genes involved in promoting this process is critical in understanding cancer development. We examined the repair efficiency of DNA containing microhomologies and found a positive correlation between microhomology length and quality with repair efficiency. Furthermore, our work directly implicates the major DNA replication and repair polymerase, DNA polymerase δ , in repair of DNA damage using microhomologies, which act in conjunction with DNA polymerase λ . Our data suggest a two-polymerase model for microhomology-mediated end-joining.

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protein A (RPA) regulates pathway choice between HR and MMEJ (37). In addition, several models have been proposed that identify specific pathways that may use MHs for the repair of DNA damage (39-41). Despite current advancements in our understanding of MMEJ, the precise involvement of DNA polymerases in supporting the repair of DSBs using MHs remains poorly understood. DNA polymerase λ (also called Pol4 in budding yeast) and its human homolog Pol λ are considered to be the primary candidates for the DNA polymerases working in NHEJ and MMEJ (4, 5, 42–46). Both genetic and biochemical evidence shows that Pol δ is recruited during HR to extend Rad51-dependent recombination intermediates (47-50). Recent analysis using pol32 mutants (5, 10) implicated the Pol32 subunit of Pol δ in MMEJ. Pol32 and Pol31 were also identified as subunits of the DNA polymerase zeta complex (Pol ζ) (51, 52), but previous analysis showed no effect of rev3 mutants in MMEJ (10). REV3 encodes the catalytic subunit of Pol ζ . However, an involvement of Pol δ had not been demonstrated directly before, and it is possible that Pol32 could act in conjunction with yet another DNA polymerase.

Here, we report the development of a series of interchromosomal MMEJ assays in diploid *S. cerevisiae* to assess the mechanisms underlying the repair of DSBs using varying MHs. We focus on diploid cells, as they represent the natural state of budding yeast, which is a diplontic organism (53). The yeast mating-type switching system represents a mechanism to return haploid yeast as efficiently as possible to diploidy (54). Using a combination of genetic, molecular, and in vivo chromatin immunoprecipitation (ChIP) experiments, we provide compelling evidence for a direct involvement of Pol δ in coordinating with Pol λ in MMEJ in budding yeast.

Results

Development of an Interchromosomal MMEJ Assay with Varying MH **Length and Quality in Diploid S. cerevisiae.** To determine the role of DNA polymerases in MMEJ repair, we designed a genetic assay in diploid S. cerevisiae, which measures interchromosomal MMEJ frequency by using varying lengths of complete or incomplete MH shared between two truncated *his3* alleles, *his3-\Delta 3'* and *his3-\Delta 5'* (Fig. 1A). The his $3-\Delta 3'$ allele, located at the HIS3 locus on chromosome XV, and the $his3-\Delta5'$ allele, at the *LEU2* locus on chromosome III, share increasing amounts of complete (16 bp, 20 bp, or 25 bp) or incomplete (14-2-2 bp, 14-2-4 bp, or 14-2-9 bp) MHs (Fig. 1B). The chromosome XV homolog shares no homology to either his3 allele containing his $3\Delta 200$, which lacks the entire HIS3 ORF and ~ 200 bp of flanking sequence. Adjacent to each truncated his3 allele is an HO endonuclease recognition sequence allowing for the creation of DSBs upon expression of the galactose-inducible HO endonuclease at the trp1 locus (55). In addition to the truncated his3 alleles, the mating-type loci (MATa, $MAT\alpha$) on chromosome III (Fig. 1A) are targeted by HO endonuclease, resulting in DSBs, which are repaired by HR or NHEJ (54). To simulate an acute exposure to a DNA damaging agent, the expression of HO endonuclease, and thereby DSB formation, was limited to 4 h. The utilization of MHs for repair is measured as the frequency of histidine prototrophs, scored on medium lacking histidine, compared with cell growth on the nutrient-rich medium, YPD (Table 1). Following DSB formation, cell viability was determined for wild-type and several key genotypes using the 14-2-4 bp substrate to assess whether viability differences confound determination of MMEJ frequencies. Viability varied maximally only about twofold compared with wild type (Table S1) and does not affect the conclusions about repair frequencies.

The MMEJ frequency of diploid wild-type cells containing both complete and incomplete MHs of varying lengths showed a concomitant increase in MMEJ frequency with increasing MH length (Table 1 and Fig. 24). Further analysis showed that changing only a few base pairs of shared homology can significantly affect the MMEJ frequency, suggesting that MMEJ repair is sensitive to alterations within the sequences of the substrates. This is supported by an observed decrease in the repair frequency

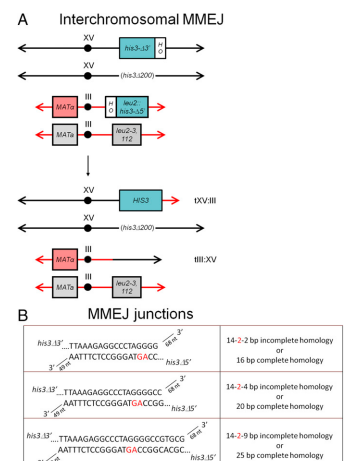


Fig. 1. Diploid MMEJ assays are defined by the length of MH and sequence continuity. (A) Recombination between *his3Δ3'* and *his3Δ5'* substrates on different chromosomes (interchromosomal MMEJ) generates *HIS3* recombinants measured by the number of histidine prototroph colonies on media lacking histidine. The homologous chromosome contains the *his3Δ200* allele, which is a complete deletion of the *HIS3* ORF. The 117 bp HO cut site (HO) is cut by a galactose-inducible HO endonuclease located at the *TRP1* locus (55). (*B*) The MMEJ junctions vary in homology length between the recombining *his3Δ3'* and *his3Δ5'* substrates from 16 to 25 bp and are fully complementary or contain a 2 bp mismatch (indicated in red) flanked by varying lengths of complementary sequences. The MMEJ junctions having incomplete homology are shown.

of the *his3-Δ3'* substrates containing a 2 bp mismatch (14-2-2 bp, 14-2-4 bp, and 14-2-9 bp) compared with the *his3-Δ3'* substrates having complete homology (16 bp, 20 bp, and 25 bp) in the shared MH with *his3-Δ5'* (Figs. 1B and 24). These results are consistent with previous studies, which showed an increase in the length of MH leads to an increase in MMEJ frequency (4, 10, 37), whereas the presence of mismatches within the recombining MH decreases MMEJ repair frequency (4, 10). Furthermore, the proximity of the mismatch to the 3'-hydroxy end (14-2-2 bp, 14-2-4 bp, and 14-2-9 bp) results in a differential decrease in MMEJ repair of 153-fold, 21-fold, and 10-fold, respectively, compared with the complete homology substrates (Fig. 24). These results demonstrate MMEJ occurs in diploid *S. cerevisiae* and is sensitive to the presence and location of sequence mismatches, in addition to the length of MH used for repair.

Analysis of wild-type unselected colonies grown on YPD and selected His⁺ MMEJ recombinants following expression of the HO endonuclease revealed an intact MATa, $MAT\alpha$, and leu2 locus for almost all of the colonies examined (Table S2). However, the presence of $his3-\Delta3'$ and $his3-\Delta5'$ was observed in only 8/16 and 7/16 unselected colonies tested, respectively. These results suggest that the repair of MATa and $MAT\alpha$ occurs independent of

Table 1. Frequencies of interchomosomal MMEJ in wild-type and mutant diploid and haploid strains

Genotype*	14- 2 -2 bp	14- 2 -4 bp	14- 2 -9 bp	16 bp	20 bp	25 bp
Wild type	1.57×10^{-8} (1)	4.51×10^{-7} (1)	2.41×10^{-6} (1)	1.2×10^{-6} (1)	9.4×10^{-6} (1)	2.7×10^{-5} (1)
	[1.1–1.9]	[3.5–5.0]	[1.7-4.9]	[0.53-2.0]	[5.4–11.2]	[0.9–5.5]
ku70∆/ku70∆	ND^{t}	$3.51 \times 10^{-7} \ (\downarrow 1.2)$	ND	ND	$5.1 \times 10^{-6} \ (\downarrow 1.8)$	ND
		[2.9-4.5]			[1.4–11.2]	
rad52∆/rad52∆	$1.8 \times 10^{-6} \ (\uparrow 115)$	$2.3 \times 10^{-5} (\uparrow 51)$	$1.5 \times 10^{-4} \ (\uparrow 62)$	$1.3 \times 10^{-5} \ (\uparrow 10.8)$	$7.3 \times 10^{-5} \ (\uparrow 7.8)$	$6.4 \times 10^{-5} \ (\uparrow 24)$
	[1.2–2.2]	[1.3-4.2]	[1.1–1.9]	[0.64–1.6]	[5.5–8.9]	[4.2–8.8]
rad51∆/rad51∆	$1.1 \times 10^{-6} \ (\uparrow 71)$	$9.0 \times 10^{-6} \ (\uparrow 20)$	ND	ND	$8.0 \times 10^{-5} \ (\uparrow 7.8)$	ND
	[0.8–1.4]	[7.5–10.5]			[4.9–9.1]	
$rad1\Delta / rad1\Delta$	ND	$1.1 \times 10^{-6} \ (+2.4)$	ND	ND	ND	ND
rad52∆lrad52∆		[0.37–2.5]				
pol 4Δ /pol 4Δ	ND	$2.15 \times 10^{-7} (\downarrow 2.1)$	ND	ND	ND	ND
po. :=:po: :=		[1.8–2.5]				
dnl4∆/dnl4∆	$1.0 \times 10^{-8} \ (\downarrow 1.6)$	$3.9 \times 10^{-7} (\downarrow 1.2)$	ND	ND	$5.8 \times 10^{-6} \ (\downarrow 1.6)$	ND
a	[5.3–1.2]	[3.4–4.5]	2	2	[4.7–7.5]	
pol4∆/pol4∆	ND	$6.0 \times 10^{-6} (+13.3)$	ND	ND	ND	ND
rad52Δ/rad52Δ	110	[4.4–7.3]	110	ND.	110	110
pol4∆/pol4∆	ND	$1.7 \times 10^{-10} (\downarrow 2441)$	ND	ND	ND	ND
pol32Δ/pol32Δ	ND	[0.8–2.1]	ND	ND	ND	ND
pol4Δ/pol4Δ	ND	$1.3 \times 10^{-10} \ (\downarrow 3192)$	ND	ND	ND	ND
pol3-ct/pol3-ct	ND	[1.0–3.3]	ND	ND	ND	ND
pol32Δ/pol32Δ	$3.8 \times 10^{-9} \; (\downarrow 4.2)$	$5.2 \times 10^{-8} (\downarrow 8.3)$	$5.1 \times 10^{-8} \ (\downarrow 50)$	ND	$5.5 \times 10^{-8} \ (\downarrow 167)$	ND
μοισεΔιμοισεΔ	[2.1–6.3]	[2.1–6.9]	[3.3–10.5]	ND	[1.4–9.6]	ND
pol3-ct/pol3-ct	$1.0 \times 10^{-9} \ (\downarrow 15.6)$	$2.6 \times 10^{-9} \ (\downarrow 167)$	$3.0 \times 10^{-7} (\downarrow 8.3)$	ND	$[1.4-3.0]$ $1.2 \times 10^{-6} (\downarrow 7.7)$	ND
	[0.6–2.3]	[1.9–3.6]	[1.7–4.4]	ND	[0.8–2.1]	ND
rev3Δ/rev3Δ	$[0.0-2.3]$ $1.2 \times 10^{-8} (\downarrow 1.2)$	$[1.9-3.0]$ $2.4 \times 10^{-7} (\downarrow 1.9)$	$[1.7-4.4]$ $1.2 \times 10^{-6} (\downarrow 2)$	ND	$[0.8-2.1]$ $8.6 \times 10^{-6} (\downarrow 1.1)$	ND
rev3\(\Delta\)rev3\(\Delta\)	[1.0–3.0]	2.4 × 10 (\$1.9) [1.4–3.4]	[0.6–1.6]	ND	[4.7–11.9]	ND
144/ 144	[1.0–3.0] ND	$[1.4-3.4]$ $1.4 \times 10^{-7} (\downarrow 3.2)$	[0.6–1.6] 1.5 × 10 ^{–6} (↓1.6)	ND	[4.7–11.9] ND	ND
pol4∆/pol4∆ rev3∆/rev3∆	ND	[0.9–2.0]	[0.7–2.0]	ND	ND	ND
	$3.7 \times 10^{-9} \; (\downarrow 4.2)$	$[0.9-2.0]$ $1.55 \times 10^{-7} (\downarrow 2.9)$	$[0.7-2.0]$ $1.7 \times 10^{-6} (\downarrow 1.4)$	ND	ND	ND
pol3-01/pol3-01	[2.5–6.0]			ND	ND	ND
		[1.2–2.7] 1.2 × 10 ^{−7} (↓3.8)	[0.8–2.5]	ND	$3.2 \times 10^{-6} \; (\downarrow 2.9)$	ND
$msh2\Delta/msh2\Delta$	ND		ND	ND		ND
	ND	[0.8–1.8]	ND	ND	[1.4–4.9]	ND
rad59∆/rad59∆	ND	$1.4 \times 10^{-7} \ (\downarrow 3.2)$	ND	ND	$4.2 \times 10^{-6} \ (\downarrow 2.2)$	ND
150.4 150.	ND	[0.5–2.3]	ND	ND	[1.3–6.5]	NB
rad59∆/rad59∆	ND	$7.4 \times 10^{-6} \ (+16.4)$	ND	ND	ND	ND
rad52∆/rad52∆		[5.0–8.4]				
Wild type [‡] Mata-inc	ND	1.93×10^{-7} (1)	ND	ND	ND	ND
(Haploid)		[1.3–3.2]				
Wild type [‡] <i>Mat</i> α	ND	$1.57 \times 10^{-7} \ (\downarrow 1.2)$	ND	ND	ND	ND
(Haploid)		[0.9–2.8]				
rad52∆ [‡] Mata-inc	ND	2.3×10^{-6} (11.9)	ND	ND	ND	ND
(Haploid)		[1.2–4.0]				
rad52Δ [‡] Matα	ND	1.51×10^{-6} (7.8)	ND	ND	ND	ND
(Haploid)		[1.1–2.2]				

The median translocation frequency was determined for each strain from at least 15 independent cultures. Fold increase (1), or decrease (1), rounded to integers, from the median frequency obtained with the wild-type strain, is indicated in parentheses. The 95% confidence intervals were determined using Table S5 and are indicated in brackets.

MMEJ. The loss of $his3-\Delta3'$ and $his3-\Delta5'$ is consistent with repair by HR using the homolog as a template. Furthermore, an intact leu2 locus demonstrates at least one chromosome III homolog is recovered following DSB induction. Identification of the reciprocal translocation (see Fig. 1A) occurred only in a small subset of His⁺ recombinants (1/14), which is similar to previous reports (55, 56) (Table S2).

MMEJ Is Independent of Ku70 and Ligase 4 but Suppressed by Rad52.

To ascertain whether our assay fulfills the criteria that define MMEJ, we examined the effect of $ku70\Delta$, $lig4\Delta$, and $rad52\Delta$ single mutants on the frequency of MMEJ repair. Because MMEJ in

budding yeast was initially defined as being independent of Ku70, Ligase 4, and Rad52 (5, 10), we expected the loss of either Ku70 Ligase 4 or Rad52 function to have little effect on the overall frequency of MMEJ repair. However, recent findings have shown MMEJ requires or is inhibited by Rad52 depending on the length of MH (4, 10, 37, 38). Using the intermediate MMEJ substrates (14-2-4 bp and 20 bp), we observed no significant difference in MMEJ repair frequency between $ku70\Delta/ku70\Delta$ or $lig4\Delta/lig4\Delta$ mutants and wild-type cells (Fig. 2B and Table 1). In contrast, the $rad52\Delta/rad52\Delta$ mutants showed a dramatic increase in MMEJ repair frequency for both substrates tested (Fig. 2B and Table 1), suggesting Rad52 can inhibit DSB repair using MHs. We confirmed

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^{*}All strains possess the $his3-\Delta5'$ substrate at the LEU2 locus on one copy of chromosome III and the $his3-\Delta3'$ substrate at the HIS3 locus (Fig. 1). The $his3-\Delta5'$ and $his3-\Delta3'$ substrates share varying MHs of HIS3 coding sequence indicated above each column. Complete genotypes are given in Table S3.

†Frequency not determined (ND).

[‡]Haploid strains.

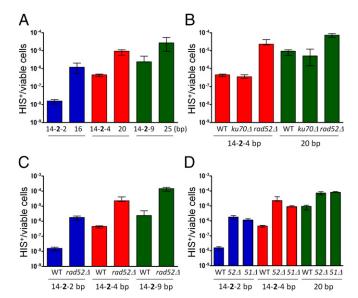


Fig. 2. MMEJ is influenced by Rad52, Rad51, the length and continuity of MH, but is independent of Ku70. (A-D) The median frequency, $\pm 95\%$ confidence interval, of interchromosomal MMEJ between the his3Δ3' and his3 $\Delta 5'$ substrates was calculated from a minimum of 20 independent cultures. Homology shared between $his3\Delta3'$ and $his3\Delta5'$ varies in length and the presence (14-2-2, 14-2-4, or 14-2-9 bp) or absence (16, 20, or 25 bp) of a 2 bp mismatch. All strains are diploid and homozygous for the indicated mutation. (A) Analysis of MMEJ frequencies in wild-type (WT) strains containing MH of 14-2-2 (WDHY2948), 16 (WDHY3427), 14-2-4 (WDHY2949), 20 (WDHY3188), 14-2-9 (WDHY3001), and 25 bp (WDHY3426). (B) MMEJ frequencies of ku70∆ (WDHY3281 and WDHY4391) or rad52∆ (WDHY3475 and WDHY3598) single mutants in the MMEJ assay with (14-2-4 bp) or without (20 bp) a 2 bp mismatch within the MH. (C and D) Influence of rad524 (WDHY3644, WDHY3475, and WDHY3597) single mutants (C) and rad514 (WDHY4352, WDHY4423, and WDHY4502) single mutants (D) in the MMEJ assay containing varying lengths of homology. The rad524 single mutant MMEJ frequencies using 14-2-2, 14-2-4, or 20 bp of MH are represented in panels C and D, B-D, and B and D, respectively.

this inhibitory effect of Rad52 using other MMEJ substrate variants, which are independent of MH length, sequence continuity, and cell viability (Fig. 2C, Table 1, and Table S1). In addition, unselected colonies and His⁺ recombinants from rad52∆ mutants retained at least one chromosome III homolog, but only 12/18 YPD colonies retained both $MAT\alpha$ and $his3\Delta5'$ located on either side of the chromosome III centromere (Table S2; see Fig. 1A). This suggests that two simultaneous breaks on chromosome III in rad52Δ mutants lead to increased chromosome loss. Rad52 has several functions in DSB repair, which include annealing of single-stranded homologous sequences or the replacement of RPA-coated single-strand DNA with Rad51 to form a nucleoprotein filament required for HR. To determine if the inhibitory effect observed in rad52\Delta mutants is due to Rad51 filament formation, we examined $rad51\Delta/rad51\Delta$ mutants. MMEJ frequencies were very similar in the $rad51\Delta$ and $rad52\Delta$ mutants (Fig. 2D and Table 1), suggesting HR directly competes with MMEJ in DSB repair through the formation of Rad51 nucleoprotein filaments mediated by Rad52.

To test if HR competes with MMEJ through the formation of Rad51 nucleoprotein filaments and the subsequent repair using the homologous chromosome, we analyzed MMEJ repair in wild-type and *rad52Δ* mutants in haploid strains possessing the 14-2-4 bp substrate. Under these conditions, haploid wild-type cells showed a small, but significant, decrease in MMEJ repair frequency compared with diploid wild-type cells. This effect is independent of DSB formation at the *MAT* locus and is not accompanied by a significant drop in cell viability (Table 1 and

Table S1). However, haploid $rad52\Delta$ mutants display a smaller increase in MMEJ repair frequency than diploid $rad52\Delta$ mutants compared with haploid or diploid wild-type cells, respectively. Again, this effect is not due to decreased cell viability or DSB formation at the MAT locus (Table 1 and Table S1). These results suggest Rad52 functions to inhibit MMEJ by promoting Rad51 filament formation and thereby HR but does not require the presence of a homologous template.

DNA Polymerase δ Promotes MMEJ Repair. The major emphasis of previous studies analyzing the genetic requirements of MMEJ has focused on components predicted to function in the early steps of resection, annealing, and end-processing of the nonhomologous 3' flap (4, 5). However, only a few of these studies have examined the role of DNA polymerases in the final steps of MMEJ repair. Although a role for Pol λ in MMEJ has been previously reported (4, 5), an involvement of other DNA polymerases is unclear in S. cerevisiae. A recent report demonstrated a role of the nonessential DNA polymerase subunit Pol32 in MMEJ (10). Pol32 associates with Pol31 and the catalytic subunit Pol3 to form the hetero-trimeric DNA polymerase delta (Pol δ), which functions in lagging strand synthesis during DNA replication and in the extension of HR intermediates (47, 48, 50, 57–59). In addition, DNA polymerase ζ (Pol ζ) is composed of the Rev3-Rev7 heterodimer and functions in DNA translesion synthesis as a four-subunit Pol ζ complex with Pol32 and Pol31 (51, 52). Therefore, the involvement of Pol32 in MMEJ repair suggests that either Pol δ or Pol ζ , or both, is potentially involved in the repair of DSBs using MHs during the extension step of MMEJ. To examine the role of Pol32 further, we tested $pol32\Delta$ / pol32∆ mutants in our diploid MMEJ assay and found a significant decrease in the frequency of MMEJ repair compared with wild type regardless of MH length or the presence of a mismatch (Fig. 3A and Table 1). Depending on the specific substrate, 76% to \geq 99% of the recombinants depended on Pol32 (Table 1). Because this decrease in $pol32\Delta$ mutants could be due to its interaction with Pol ζ , we decided to test the effect of the rev3 Δ mutation on MMEJ repair to determine if a similar decrease was also observed. A modest decrease of twofold was observed in $rev3\Delta/rev3\Delta$ mutants using the 14-2-4 bp and 14-2-9 bp MH substrates for MMEJ repair, whereas the 14-2-2 bp and 20 bp MH substrates showed no defect in MMEJ repair (Table 1). Instead, the $pol32\Delta$ mutation exerted its largest effect on the 20 bp MH substrate (Table 1), demonstrating that Pol32 and Rev3 show different substrate specificity. Taken together, these results demonstrate a robust and consistent role of Pol32 in promoting MMEJ repair, which is mainly independent of Rev3 function.

The above results suggest the defect observed in $pol32\Delta$ / pol32Δ cells is through its interaction with Pol3 and Pol31 in forming Pol δ . The involvement of Pol δ in MMEJ is more difficult to test directly, as both POL31 and POL3 are essential genes. However, there are several alleles of POL3, which alter the protein function but allow for cell growth. One such allele is pol3-ct, which is missing the last four C-terminal amino acids and was originally identified as a mutant defective in meiotic gene conversion (60). Subsequent analysis revealed that in addition to its meiotic phenotypes, pol3-ct was also defective in mitotic gene conversion and break-induced replication (BIR) due to a decrease in the interaction between Pol3 and Pol31 resulting in an unstable Pol δ complex (48, 61, 62). To determine if the role of Pol32 in MMEJ repair was due to a defect in the Pol δ complex, we tested the frequency of MMEJ in pol3-ct/pol3-ct mutants. A significant decrease in the frequency of MMEJ was observed in pol3-ct/pol3-ct, which was similar to the $pol32\Delta/pol32\Delta$ mutants (Fig. 3B and Table 1). The cell viability of pol3-ct and pol32 Δ mutants is similar to wild type and each other (Table S1). These results show the involvement of Pol32 in MMEJ is due to the action of Pol δ in promoting the repair of DSBs using MHs and is

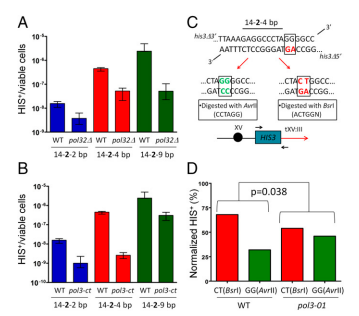


Fig. 3. DNA polymerase δ promotes MMEJ independent of MH length and can correct mismatched nucleotides near the 3'-hydroxyl end. (A and B) The median frequency, ±95% confidence interval, of interchromosomal MMEJ between the $his3\Delta3'$ and $his3\Delta5'$ substrates was calculated from a minimum of 20 independent cultures. Homology shared between his3 $\Delta 3'$ and his3 $\Delta 5'$ varies in length (14-2-2, 14-2-4, or 14-2-9 bp) and contains a 2 bp mismatch. The contribution of Pol32 (WDHY4390, WDHY3126, and WDHY3165) (A) and the DNA polymerase δ allele, *pol3-ct* (WDHY3872, WDHY3809, WDHY3810, and WDHY3807) (B), in promoting MMEJ using varying lengths of MH compared with wild-type (WT) (WDHY2948, WDHY2949, and WDHY3001). (C and D) Sequence analysis of histidine prototrophic colonies following MMEJ repair in WT (WDHY2949) and cells harboring the exonucleasedefective DNA polymerase δ allele, pol3-01 (WDHY3809 and WDHY3807). Correction of the 2 bp GG mismatch on the top strand would result in a CT conversion, thereby making the sequence sensitive to Bsrl digestion, Retention of the 2 bp GG mismatch on the top strand would allow for digestion with AvrII. Initially 20 colonies were chosen for genomic DNA extraction, PCR amplification of the His+ recombinant (using primers indicated as arrows in C), digestion with both AvrII and BsrI, and then verification by sequencing. Subsequent His+ recombinants were collected as above but were digested with both AvrII and BsrI to determine sequence and not sent for sequencing. A total of 121 WT and 100 pol3-01 His+ recombinants were analyzed, and the two sets of data were compared using a contingency table and Fisher's exact test.

the first direct evidence, to our knowledge, demonstrating the involvement of Pol δ in MMEJ repair. Although the overall quantitative effect of the $pol32\Delta$ and pol3-ct mutation was similar, the substrate specificity pattern is somewhat different between both. Although $pol32\Delta$ mutants had the largest effect on the 20 bp MH substrate, the pol3-ct mutation affected the 14-2-4 bp substrate most. This suggests greater complexity and potential for the involvement of multiple DNA polymerases in a single MMEJ event.

To further investigate the role of Pol δ in MMEJ repair, we analyzed the junction sequence of His⁺ recombinants from cells, which used 14-2-4 bp of shared MH between the *his3-\Delta3'* and *his3-\Delta5'* alleles for repair (Fig. 3C). The exonuclease activity of Pol δ has been shown to be active during DSB repair (40, 63). We chose the 14-2-4 bp substrate to be within the range of the Pol δ proofreading window (64). The 2 bp mismatch, which exists between the *his3-\Delta3'* and *his3-\Delta5'* alleles, can be corrected by mismatch repair, resulting in two distinct classes of His⁺ recombinants in equal proportions that can be distinguished by restriction digestion using either *Avr*II or *Bsr*I (Fig. 3C). Alternatively, the two mismatched guanine nucleotides on *his3-\Delta3'*

located close to the 3'-hydroxyl end could be corrected by the proofreading exonuclease activity of Pol δ , which would bias the outcome toward His+ recombinants digested by the BsrI restriction enzyme. In fact, 82 out of 121 (68%) of all His⁺ recombinants recovered from wild-type cells were digested by BsrI, demonstrating a significant bias toward one class of MMEJ recombinants, which is potentially mediated by the exonuclease function of Pol δ (Fig. 3D). This is in contrast to His⁺ recombinants from cells using the 14-2-2 bp or 14-2-9 bp substrates for repair, which showed 20 out of 20 and 12 out of 32 were digested by BsrI, respectively. To test if the exonuclease function of Pol δ biased the His+ recombinants toward the BsrI class, we examined 100 independent His⁺ recombinants from cells containing the 14-2-4 bp substrate in the exonuclease-defective pol3-01 strain (65). The AvrII and BsrI classes of His⁺ recombinants recovered from pol3-01/pol3-01 were in near equal proportion and significantly different from wild type (P value = 0.0385; Fig. 3D). All His⁺ recombinant junctions were susceptible to either AvrII or BsrI, showing that all products underwent faithful repair. These results provide further independent evidence for a role of Pol δ in the extension of intermediates during MMEJ repair.

DNA Polymerase δ Associates with DSBs During MMEJ Repair. Because the results of our genetic experiments demonstrate that Pol δ is involved in MMEJ repair, we decided to investigate the physical association of Pol3 with the $his3-\Delta3'$ MMEJ substrate before and after DSB formation using ChIP (Fig. 4A). These experiments were performed in $rad52\Delta/rad52\Delta$ strains that contained the 14-2-4 bp of shared MH between the $his3-\Delta3'$ and his3-Δ5' alleles, allowing for an increased probability of crosslinking Pol3 to the MMEJ substrate due to the higher frequency of repair. Following the addition of galactose to the cell cultures, a progressive increase in DSB formation was observed, which peaked at 90% after 2 and 4 h of HO endonuclease induction (Fig. 4C), consistent with previous analyses (66). The kinetics of Pol3 recruitment at his3-Δ3' follows DSB formation, with an apparent 2 h delay, showing a 3.5-fold and sixfold enrichment of $his3-\Delta3'$ Pol3 ChIP over background at the 2 and 4 h time points, respectively (Fig. 4C). To simulate the genetic MMEJ assay, cells were collected, washed, and placed in YPD after 4 h to repress HO endonuclease expression and continue growth. Following this shift to YPD, we saw a decrease in the presence of the DSB from 6 to 12 h, which is a measure of cells rejoining the broken chromosome ends by NHEJ and subsequent growth of those cells within the culture over the indicated time course (Fig. 4C). In contrast, the 11-fold and fourfold enrichment of his $3-\Delta 3'$ at 6 and 8 h, respectively, demonstrates that Pol3 is functioning in MMEJ after DSB formation (Fig. 4C). By 12 h, there is no detectable enrichment of $his3-\Delta3'$, indicating the repair process is completed by this time (Fig. 4C). Taken together, these results provide direct evidence for the involvement of Pol3, and thereby Pol δ , in the repair of DSBs using MHs.

The involvement of Pol δ in extending MMEJ repair intermediates suggests its role is dependent on the activities of proteins that function to process the DSBs before extension. Resection of the DSBs and annealing using MHs are required to form an intermediate containing 3' heterologous flaps derived from HO endonuclease cutting of the 117 bp HO endonuclease recognition sequence (Fig. 1B). The removal of these 3' heterologous flaps requires the action of the Rad1-Rad10 endonuclease (67) and should be necessary for Pol δ extension. Therefore, rad1 mutants should diminish the association of Pol δ with the MMEJ recombination intermediate and reduce the overall repair efficiency. To test this, we repeated the above ChIP experiments with $rad1\Delta/rad1\Delta$ $rad52\Delta/rad52\Delta$ double mutants and examined the association of Pol3 with $his3-\Delta3'$ over the indicated time course (Fig. 4C). The absence of Rad1 results in a significant decrease in the association of Pol3 with his3-Δ3' from 2 to 8 h compared with cells containing Rad1 (Fig. 4C), which correlates

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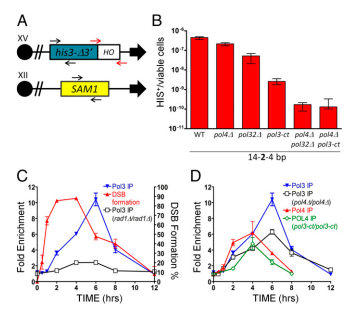


Fig. 4. DNA polymerases δ and λ associate with a DSB containing MHs and cooperate in the subsequent repair of a DSB using MMEJ. (A) Diagram of the chromosomal loci and the relative location of primers used in the quantification of DSB formation (red arrows) or ChIP analysis (black arrows) of DNA polymerase δ and λ . (B) The median frequency, $\pm 95\%$ confidence interval, of $pol4\Delta$, $pol32\Delta$, pol3-ct, $pol4\Delta$ $pol32\Delta$, and $pol4\Delta$ pol3-ct single and double mutants using 14-2-4 bp of MH between the his3 $\Delta 3'$ and his3 $\Delta 5'$ substrates was calculated from a minimum of 20 independent cultures. The pol32∆ and pol3-ct single mutants are also represented in Fig. 3 A and B, respectively. (C) DSB formation was analyzed using qPCR by comparing the formation of a PCR product using primers flanking the HO endonuclease recognition cut site (red arrows in A) relative to the PCR product of SAM1 (black arrows in A), before and after HO endonuclease induction over the indicated time course (red triangle). DNA following ChIP, using antibodies to Pol3, was analyzed using qPCR by comparing the PCR product from his3\(\Delta\)3' to SAM1 (black arrows in A) both before and after DSB induction over the indicated time course (Pol3 IP, rad52Δ/rad52Δ, blue triangle; rad1Δ/rad1Δ rad52Δ/ $rad52\Delta$, black square). The mean \pm SEM of samples taken at the indicated time points from two independent cultures (WDHY3475, WDHY4487), grown concurrently, is represented. The Pol3 IPs in rad52Δ/rad52Δ (blue triangle) strains are represented in C and D. (D) ChIP analysis as in C using antibodies to Pol3 or Pol λ as indicated (Pol3 IP, $rad52\Delta/rad52\Delta$, blue triangle; Pol3 IP, $pol4\Delta/pol4\Delta$ $rad52\Delta/rad52\Delta$, black square: Pol λ IP, $rad52\Delta/rad52\Delta$, red triangle: Pol λ IP. pol3-ct/pol3-ct rad52 Δ /rad52 Δ , green diamond). The mean \pm SEM of samples taken at the indicated time points from two independent cultures (WDHY3475 and WDHY3793), grown concurrently, is represented.

with a 21-fold decrease in the frequency of His⁺ recombinants compared with $rad52\Delta/rad52\Delta$ (Table 1). These results demonstrate a strong genetic dependence of MMEJ on the heterologous flap cleavage activity of Rad1–Rad10 as a prerequisite for extension by Pol δ .

DNA Polymerases δ and λ Cooperate to Repair DSBs Using MMEJ. Studies on MMEJ using shorter MHs of 8–12 bp have implicated the involvement of Pol λ (4, 5), but longer MHs were not examined. Therefore, we tested $pol4\Delta/pol4\Delta$ mutants in the MMEJ repair assay using 14-2-4 bp of MH and observed a twofold decrease in the frequency of MMEJ while maintaining cell viability similar to wild type (Fig. 4B, Table 1, and Table S1). The involvement of Pol4 in NHEJ prompted the examination of $pol4\Delta$ mutants for intact MATa, $MAT\alpha$, leu2-3,112, $his3\Delta3'$, and $his3\Delta5'$ loci in unselected and His+ recombinant colonies. A similar proportion of intact MATa, $MAT\alpha$, and leu2 loci in unselected and His+ recombinant colonies was observed in wild-type and $pol4\Delta$ mutants (Table S2). However, a noticeable decrease of intact $his3\Delta3'$ and $his3\Delta5'$ loci in $pol4\Delta$ mutants suggests a defect in NHEJ, resulting in a

higher use of HR in repair using the homologous chromosome (Table S2).

Next, the interaction between Pol λ and Pol δ in MMEJ was determined through epistasis analysis of $pol4\Delta/pol4\Delta$ $pol32\Delta/pol32\Delta$ and $pol4\Delta/pol4\Delta$ pol3-ct/pol3-ct double mutants using 14-2-4 bp of MH for repair. A similar synergistic decrease in MMEJ frequency of over 2,500-fold was observed in both double mutants and is not due to a decrease in cell viability (Fig. 4B, Table 1, and Table S1). This suggests that Pol λ and Pol δ cooperate in MMEJ. To exclude the possibility of Pol32 exerting its effect with Pol4 as part of the Pol ζ complex, the MMEJ frequency of $pol4\Delta/pol4\Delta$ $rev3\Delta/rev3\Delta$ double mutants was tested and showed no significant decrease from single $pol4\Delta$ mutants (Table 1).

To provide additional evidence that Pol λ is functioning in MMEJ, we tested its physical association with the *his3-\Delta3'* MMEJ substrate using ChIP, as performed above with Pol3. We observed a fivefold and sixfold enrichment of *his3-\Delta3'* over background at the 2 and 4 h time points, respectively (Fig. 4D). The above enrichment is specific, as the *pol4\Delta/pol4\Delta rad52\Delta/rad52\Delta* double mutants show no significant enrichment of *his3-\Delta3'* above background for the 2 and 4 h time points. However, unlike Pol3, enrichment of Pol λ peaked at 6 h and decreased to 3.5-fold at 6 h and was indistinguishable from background at 8 h (Fig. 4D), suggesting a possible earlier role for Pol λ than Pol δ in MMEJ. Finally, recruitment of Pol3 and Pol4 to the DSB was interdependent (Fig. 4D), consistent with a two-polymerase model for MMEJ-mediated DSB repair.

Discussion

Here, we developed an interchromosomal MMEJ repair assay in diploid S. cerevisiae to assess the repair of simultaneous DSBs using MHs of varying lengths and sequence continuity. The study of MMEJ in diploids represents a departure from previously published reports (5, 10, 37) in haploid budding yeast and was chosen because it is the natural state of S. cerevisiae (53). Although NHEJ is suppressed in diploids (68–70), we find the broken loci at MATa, $MAT\alpha$, his $3\Delta 3'$, and his $3\Delta 3'$ are often repaired by HR or NHEJ potentially due to acute HO endonuclease induction, which results in similar cell viability in many mutants tested compared with wild type. Similar to the results of Villarreal et al. with haploid yeast cells and shorter MHs of 6-18 bp, we find a correlation between increasing MH length and MMEJ repair in wild-type cells with MHs of 16–25 bp (10). Using the his 3 MMEJ substrates in haploid yeast yields a significantly lower repair frequency, possibly due to the requirement of the reciprocal translocation in MMEJ His⁺ recombinants for viability. In addition, the presence of a di-nucleotide mismatch significantly decreased the frequency of MMEJ. The most dramatic decrease of nearly 100-fold was observed when using the 14-2-2 bp MH substrate for repair, indicating that the nearer the dinucleotide mismatch was to the 3'-hydroxyl end of the MH, the greater its repressive effect on MMEJ repair. This suggests that the mismatch had a destabilizing effect on the MH pairing intermediate and/or the subsequent extension of the 3'-hydroxyl end by a DNA polymerase. This effect is potentially due to the proximity of the mismatch to the 3'-hydroxyl end, which could destabilize the terminal base pairing, resulting in only 14 bp of MH used for repair. Hence, the length and degree of mismatching of the MH have a major impact on the efficiency of repair, which is possibly mediated by the thermostability of the MH pairing intermediate.

The role of Rad52 in MMEJ repair has been unclear, as initial studies suggested MMEJ was independent of Rad52 (3). In contrast to more current reports, which show the use of longer MHs (≥15 bp) for repair is Rad52-dependent, while Rad52 is inhibitory when short MHs are used for repair (≤14 bp) (10, 37). In our system, we observed an inhibitory effect of Rad52 on the frequency of MMEJ repair regardless of homology length, similar to other published observations (37). This is in contrast to Villarreal et al., who saw a dependence on Rad52 for MMEJ repair using MHs ≥15 bp but an inhibitory effect of Rad52 when MHs used to repair

the DSB were ≤ 13 bp (10). These disparate results could be the consequence of differences in the experimental design (acute vs. chronic, DSB formation), the presence of longer nonhomologous flaps, and the effect of ploidy on MMEJ. Rad52 functions to promote HR by loading Rad51 on RPA-coated ssDNA to form a Rad51 nucleoprotein filament that searches the genome for homologous sequences (71). In addition, Rad52 efficiently anneals protein-free and RPA-coated ssDNA (72, 73). One possible mechanism for an inhibitory effect of Rad52 on MMEJ could be through the loading of Rad51 and subsequent allelic recombination after extensive resection on either side of the DSB to reveal homologous sequences for recombination, thereby generating a histidine auxotroph. This is supported by a similar inhibitory effect of rad51Δ and rad52Δ mutants on MMEJ frequency, indicating there is little if any inhibitory role of Rad52 on MMEJ, which is independent of Rad51. In addition, Rad59 has been shown in diploids to function in single-strand annealing (SSA) independent of Rad52 (55), and we observe a differential effect of rad59Δ and rad52\Delta mutants on MMEJ frequency (Table 1), which supports a Rad51-dependent inhibition by Rad52. Taken together, a Rad51independent effect of Rad52 in annealing MHs during MMEJ is not supported by our data, but we cannot rule out a supportive effect of Rad52 on MMEJ using MHs shorter than 16 bp.

The key finding reported here is a direct role of DNA polymerase δ in MMEJ. Our understanding of the identity of the DNA polymerases involved in MMEJ is incomplete. In Drosophila melanogaster, DNA polymerase theta was shown to be critical for Alt-EJ (74). However, budding yeast does not contain a homolog for this DNA polymerase (75). In S. cerevisiae, MMEJ seems to be dependent on Pol32, and a role of Rev3 has been excluded (10). Our results demonstrate directly that Pol δ is required for DSB repair using MMEJ, which is largely independent of MH length and sequence continuity. First, consistent with previous results, we observed a significant decrease in the repair frequency of DSBs using varying lengths of MH with mismatches in $pol32\Delta$ mutants (Fig. 3A). Furthermore, a decrease of over 170-fold occurred in pol32\Delta mutants when the MH without mismatches (20 bp) was used for repair, suggesting Pol32 becomes critical for MMEJ when the MH has no mismatches. Second, we demonstrate a similar decrease in MMEJ repair in pol3-ct mutants affecting the catalytic subunit of Pol δ, suggesting the function Pol32 in the Pol δ holoenzyme, as opposed to alternative complexes containing Pol32, affects MMEJ (Fig. 3B). The difference in the specific effects on the different MMEJ substrates in pol32 and pol3-ct mutants raises the possibility of a more complex situation involving multiple polymerases in a single MMEJ event, consistent with the small but significant effect of rev3\Delta mutants on certain MMEJ substrates. Third, we made use of a proofreading-deficient pol3-01 mutant to show that Pol δ directly accesses the 3'-OH end during the DNA synthesis step in MMEJ and can act on proximal mismatches (Fig. 3 C and D). Fourth, we provide physical evidence by ChIP that Pol3 directly associates with DSBs showing kinetics that are consistent with a role in the DNA synthesis step of MMEJ. The maximum enrichment of his $3-\Delta 3'$ at 6 h occurs 2 h after repression of HO endonuclease expression and thus DSB formation. Furthermore, the continued association of Pol3 with his3- $\Delta 3'$ at 8 h suggests the repair process using MHs is slow and takes longer to complete than HR-mediated DSB repair (55, 76, 77). Finally, we demonstrate the association of Pol3 with the DSB is largely dependent on the activity of the Rad1-Rad10 heterodimer, as the enrichment of his3- $\Delta 3'$ is reduced up to fivefold in rad1 Δ /rad1 Δ mutants compared with cells with functional Rad1. This observation is further supported by the genetic dependence of MMEJ on Rad1 in His+ recombinant formation.

The involvement of Pol δ in MMEJ prompted us to examine other DNA polymerases involved in DSB repair using MHs. An early study in MMEJ by the Lee laboratory showed a modest role of Pol λ in MMEJ (5), which is supported by in vivo evidence showing Pol λ is proficient in gap filling of annealed MHs (38, 42).

Consistent with the above results, we observed a decrease in the MMEJ frequency of $pol4\Delta/pol4\Delta$ mutants and early association of Pol λ with MMEJ repair substrates. The synergistic decrease in $pol4\Delta/pol4\Delta$ $pol32\Delta/pol32\Delta$ and $pol4\Delta/pol4\Delta$ pol3-ct/pol3-ct double mutants suggests dynamic, mutually dependent roles for Pol λ and Pol δ in promoting MMEJ. These results are a departure from a previous report examining $pol4\Delta pol32\Delta$ double mutants (5), which show no difference between the single and double mutants. This difference is potentially due to several factors, which includes the acute expression of HO endonuclease, strain differences, use of diploid strains, longer MHs, and longer nonhomologous flaps in our study. One possibility is Pol λ and Pol δ function in separate pathways to promote MMEJ, with Pol δ-dependent repair contributing more to the overall number of MMEJ recombinants. Alternatively, Pol λ and Pol δ could function in the same pathway and cooperate in the repair process. We favor the latter explanation. Pol λ is proficient at extension in the absence of PCNA, lacks proofreading, and can extend even when the 3'-hydroxyl end contains a mismatch (42). These properties may endow Pol λ with greater efficiency than Pol δ, which requires PCNA and proofreads, at extending specific, possibly transient, heteroduplex molecules, which have low thermostability. The ChIP data show interdependence of the Pol λ and Pol δ association with the break site, not allowing a clear conclusion about a possible temporal sequence of action between both polymerases. The early Pol4 association with the DSB MMEJ substrate is reduced in pol3-ct mutants, and the partial dependence of the Pol3 association with the DSB substrate at 4 and 6 h suggests a more dynamic interaction during MMEJ, which is consistent with a two-polymerase model. This interaction is supported by previous results demonstrating the cooperation between Pol4 and Pol3 in gap filling during the repair of broken plasmids (78). A two-polymerase model has been proposed for lesion bypass of DNA damage in yeast and mammalian cells (79, 80) and similarly may allow for the most efficient polymerase to function during the initial and subsequent extension of MMEJ intermediates.

The importance of MMEJ has been highlighted by recent reports of MH use in various cellular repair processes and the presence of MH at chromosomal breakpoints (6, 81-83). However, whether alt-EJ represents a distinct repair pathway in mammalian cells and its role in cancer development is under debate (84). One potential problem is alt-EJ being defined as any DSB repair event independent of the Ku70-Ku80 heterodimer. These events are often associated with large deletions, insertions, and either no MHs or MHs ranging from 1 to 25 nt (85), but they may not all represent the same pathway for repair. It has been proposed that the repair junctions featuring limited or no MHs were repaired by a NHEJ pathway using alternative components to substitute for the canonical NHEJ components (84). Although the use of alternative NHEJ components could affect repair efficiency and the types of junctions formed, it does not explain the use of longer MHs, which are used at similar frequencies in both NHEJ-deficient and NHEJ-proficient cells (10, 37, 85). Examination of MH use in mammalian cells and yeast shows a similar range of 5-22 nt used for repair, but the frequency of repair appears to be higher in mammalian cells compared with yeast (4, 10, 21, 37, 86-88). We favor a model where longer MHs, such as those used in this study, are revealed following resection and form a stable base pairing relationship, which is further stabilized by DNA polymerase extension of the repair intermediate. Perhaps the rate-limiting step in the repair of DSBs using MHs is the formation and stability of the pairing intermediate and subsequent extension of the 3'-hydroxyl end by a DNA polymerase. Therefore, a DSB not repaired by NHEJ using limited or no MHs is subject to resection and repair by HR or MMEJ depending on the length of homology and the availability of the repair proteins.

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Materials and Methods

Strain and Plasmid Construction. All yeast strains used in this study were isogenic with W303-1A but were wild type at the RAD5 locus (Table S3). The rad52::TRP1, ku70::TRP1, rad1::LEU2, rad51::LEU2, and pol3-01 mutants were kindly provided by A. Bailis, Beckman Research Institute of the City of Hope, Duarte, CA (55, 89, 90). The pol32::KANMX and pol3-ct mutants were a generous gift from L. Symington, Columbia University, New York, (91) and L. Maloisel, Commissariat à l'énergie atomique et aux énergies alternatives, Fontenay-aux-Roses, France (48), respectively. The pol4::KANMX mutant was generated by PCR amplification of KANMX using pWDH517 as a template. The primers contain 50 nucleotides on their 5' end, which share homology to sequences upstream and downstream of the POL4 ORF. The deletion of the POL4 ORF was verified by PCR in the transformant. Because the pol3-01 is a mutator allele, all diploid strains with this mutation were constructed from haploid strains derived from freshly sporulated heterozygous diploid strains (see Table S3). The pol3-ct and pol3-01 genotypes were determined by PCR and DNA sequencing.

The his3-Δ3'-HOcs MMEJ substrate cassettes were generated by PCR from the fusion of two smaller PCR fragments into a single larger his3-Δ3'-HOcs fragment. Using pLAY500 (55) as a template, an 810 bp fragment was amplified using a primer (olWDH946; sequences for all oligonucleotides are listed in Table S4) complementary to nucleotides upstream of the HIS3 coding sequence and another corresponding to nucleotides 294-318 within the HIS3 coding sequence (ol1659, 14-2-2 bp; ol1657, 14-2-4 bp; ol1661, 14-2-9 bp; ol1128, 16 bp; ol1074, 20 bp; ol1129, 25 bp), A second 600 bp fragment was amplified from pLAY500 using a primer complementary to the 5' end of the 117 bp HOcs plus varying MHs complementary to nucleotides 294–318 of the HIS3 coding sequence (ol1660, 14-2-2 bp; ol1658, 14-2-4 bp; ol1662, 14-2-9 bp; ol1130, 16 bp; ol1075, 20 bp; ol1131, 25 bp) and another primer complementary to nucleotides downstream of the HIS3 coding sequence (ol949). The shared MH between the 810 bp and 600 bp PCR fragments was modified to create all of the various MHs used in this study (14-2-2 bp, 14-2-4 bp, 14-2-9 bp, 16 bp, and 25 bp). The 810 bp and 600 bp PCR fragments corresponding to each of the above MHs were paired and used as templates to amplify a larger fragment of his3- $\Delta 3'$ -HOcs, each with a defined MH length and sequence continuity. The his3-Δ3'-HOcs fragments were digested with BamHI and cloned into YIp365R, which possesses a URA3-selectable marker that had been digested with BamHI. Each YIp365R+ his3-\(\Delta 3'\)-HOcs was linearized by digestion with MscI to target its transplacement into the HIS3 locus following transformation and selection for uracil prototrophy. Uracil prototrophs were plated to 5-FOA media to select for plasmid loss, and the resulting 5-FOAr colonies assessed for integration by PCR and sequencing.

Construction of the other translocation assay components his3-Δ200, $leu2::HOcs-his3-\Delta5'_{(300)}$, and trp1::GAL1-HO-KAN-MX were published previously (55). All strains possess the his3- $\Delta 5'$ substrate at the LEU2 locus on one copy of chromosome III. The his3- $\Delta 3'$ substrates are located at the HIS3 locus on one copy of chromosome XV. The his3- $\Delta 5'$ and his3- $\Delta 3'$ substrates share varying amounts of MH (14-2-2 bp, 14-2-4 bp, 14-2-9 bp, 16 bp, 20 bp, and 25 bp) within the HIS3 coding sequence.

Interchromosomal MMEJ Assay and Analysis. We inoculated 2 mL cultures of YP-Raffinose medium containing 1% yeast extract, 2% (wt/vol) peptone, and 2% (wt/vol) raffinose with single colonies and incubated them for 24 h at 30 °C. The next day, galactose was added to the cultures to a final concentration of 2% (vol/vol) to induce expression of the HO endonuclease. After 4 h of induction, the cells were plated to medium lacking histidine, and the interchromosomal MMEJ frequency was determined by dividing the number of histidine prototrophic colonies by the total number of viable cells as determined by plating dilutions onto YPD. The median translocation frequencies represent at least 20 independent cultures, and the 95% confidence intervals were determined using Table S5. Selected YPD and His⁺ recombinant colonies from substrates using the 14-2-4 bp of MH for repair were subjected to PCR amplification of the MMEJ product (ol968 and ol1013), reciprocal translocation (ol936 and ol1549), leu2-3,112 (ol936 and ol1623), MATa (olAP53 and olAP54), MAT α (olAP52 and olAP54), $his3\Delta3'$ (ol1013 and ol1549), and $his3\Delta5'$ (ol953 and ol936) loci. Subsequent analysis of the MMEJ PCR product by sequencing and restriction digestion was used to determine which two nucleotides within the mismatched MH were retained. Analysis of DSB formation by HO endonulcease, following the addition of galactose, was done using quantitative PCR (qPCR) with primers flanking the 117 bp HO recognition sequence located adjacent to the his3△3′ allele (ol935 and ol960). The qPCR signal was normalized to the PCR amplification signal from SAM1, which is not located near any DSB and the percent of broken molecules at the 0 h time point.

Cell Viability. Cell viability was determined from YP-Raffinose cultures of the indicated genotypes with the 14-2-4 bp substrate after the addition of galactose. HO endonuclease was induced for 4 h at 30 °C, at which point the cell number was assessed by hemocytometer. Approximately 200-500 galactosetreated cells were plated to YPD, incubated at 30 °C for 4 d, and the number of colonies counted. Cell viability was determined by dividing the number of colonies by the number of cells plated and multiplying the quotient by 100. The mean plating efficiencies from at least 10 independent trials \pm SEM were reported. Differences between plating efficiencies for which the ±SEM did not overlap are considered significantly different.

ChIP Assay. Newly dissected mutant spores of rad52::TRP1, pol4::KANMX rad52:: TRP1 pol3-ct rad52::TRP1, and rad1::LEU2 rad52::TRP1 double mutants possessing the interchromosomal MMEJ substrates, which share 14-2-4 bp of MH, were grown in YPD overnight, diluted to an OD₆₀₀ of 0.2 in fresh YP-Raffinose, and allowed to grow to an OD_{600} of ~ 1.0 , at which time 45 mL was removed (0 h), pelleted, and immediately subject to the ChIP procedure described previously (56). Galactose was added to the remaining culture to a final concentration of 2% (vol/vol), which induces the expression of HO endonuclease and DSB formation at the $his3\Delta3'$ allele. At varying intervals following the addition of galactose, 45 mL were removed and immediately subject to the ChIP procedure.

Briefly, the pelleted cells were exposed to formaldehyde, washed, lysed in a buffer containing protease inhibitors, and the DNA sheared to $\sim\!500$ bp by sonication. The whole cell extract was subjected to immunoprecipitation by incubating with 5 µL of Pol3 antiserum, kindly provided by Burgers, at 4 °C overnight with shaking. The next day, protein A agarose beads were added to the samples and incubated at 4 °C for 3 h. The beads were then washed successively in SDS, high salt, and LiCl solutions, and finally TE buffer. DNA/protein/ antibody complexes were eluted from the beads with elution buffer (1% SDS/TE) and the DNA/protein cross-links reversed by incubation at 65 °C overnight. Proteins were then digested with proteinase K for 2 h, extracted with phenol: chloroform, and the DNA precipitated with ethanol. Selected DNA sequences were analyzed by qPCR in triplicate using a Roche LightCycler480 with the following primer sets: A 198 bp SAM1 sequence was amplified with the primers ol1653 and ol1654 and a 166 bp sequence specific to the 5' end of HIS3 amplified with the primers ol1651 and ol1652. Relative quantification of the PCR products was performed using the Roche LightCycler480 software. The HIS3 and SAM1 PCR signals obtained by ChIP were normalized to HIS3 and SAM1 PCR signals obtained from the above ChIP strain inputs that have not been exposed to the Pol3 antibody. The normalized HIS3 and SAM1 PCR signals were used to obtain the H/S ratios. The mean H/S ratios \pm 1 SEM reported were derived from two independent strains for each genotype and repeated a minimum of four times. The SAM1 PCR was chosen as the control locus for the HIS3 PCR because it is not located near an induced DSB sequence.

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