Yeast Exonuclease 5 Is Essential for Mitochondrial Genome Maintenance

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Yeast exonuclease 5 is encoded by the YBR163w (DEMI) gene, and this gene has been renamed EXO5. It is distantly related to the Escherichia coli RecB exonuclease class. Exo5 is localized to the mitochondria, and EXO5 deletions or nuclease-defective EXO5 mutants invariably yield petites, amplifying either the ori3 or ori5 region of the mitochondrial genome. These petites remain unstable and undergo continuous rearrangement. The mitochondrial phenotypic of exo5Δ strains suggests an essential role for the enzyme in DNA replication and recombination. No nuclear phenotype associated with EXO5 deletions has been detected. Exo5 is a monomeric 5’ exonuclease that releases dinucleotides as products. It is specific for single-stranded DNA and does not hydrolyze RNA. However, Exo5 has the capacity to slide across 5’ double-stranded DNA or 5’ RNA sequences and resumes cutting two nucleotides downstream of the double-stranded-to-single-stranded junction or RNA-to-DNA junction, respectively.

Endonucleases and exonucleases are intimately involved in all aspects of DNA metabolism in the cell. In mitochondria, several constitutive nucleases have been identified that contribute to the proper maintenance of the mitochondrial genome through replication and recombination pathways. In addition, nucleases can localize to mitochondria in response to DNA stress in order to mediate appropriate DNA repair. Among the constitutive mitochondrial nucleases in Saccharomyces cerevisiae are the Nuc1 nuclease that contributes to DNA recombination efficiency and functions in apoptosis (4, 38) and the Cce1 endonuclease that resolves recombination intermediates (29). The Din7 endonuclease is a mitochondrially located 5’ flap endonuclease related to FEN1 (20). While deletion of the gene for either of these enzymes produced marginal mitochondrial phenotypes, more severe phenotypes were observed when combined deletions of these nuclease genes were studied or when they were combined with deletions of other genes involved in DNA recombination or repair, such as MHR1 or MSH1 (20, 22, 27). Recently, human Dna2 was shown to localize to both the nuclear and mitochondrial compartments and to participate in mitochondrial DNA replication and base excision repair (11, 39). Its function in yeast mitochondrial DNA maintenance has not been studied in detail. Finally, the 5’ flap endonuclease FEN1, which normally functions in primer RNA degradation during Okazaki fragment maturation in the nucleus, also localizes to the mitochondrion in response to DNA damage, participating in long-patch base excision repair (19, 23).

Since mitochondrial function is not essential to yeast survival, dysfunction caused by mutations of the mitochondrial genome can be readily detected as a loss of respiration function, which is scored as the inability to grow on nonfermentable carbon sources. A defect in the mitochondrial DNA polymerase γ MIP1 results in complete loss of the mitochondrial DNA, and the mutant fails to grow on glycerol-containing media lacking glucose (14). Such cells are designated ρ−. Genome maintenance defects can also result in the generation of petite mutants that still contain mitochondrial DNA. Generally, most of the mitochondrial genome has been deleted, and a small origin-containing region has been amplified (ρ−). S. cerevisiae contains eight such origin regions that are highly similar in sequence and are distributed over the 86-kb mitochondrial genome (8, 9, 15). Petites that have amplified the ori5 region have been studied more extensively (16, 22).

While the nucleases listed above participate in the proper maintenance of the mitochondrial genome through their replication and/or recombination functions, none appears to be essential for the integrity of the mitochondrial genome. One reasonable explanation for these observations is functional redundancy. Indeed, functional nuclease redundancy is quite common; it has been observed in the process of DNA degradation during mismatch repair in Escherichia coli, during Okazaki fragment maturation in yeast, and during the resection of double-stranded breaks in yeast (7, 25, 33). However, the possibility remains that an additional nuclease(s) is active in the mitochondrion. The present paper describes an essential mitochondrial exonuclease that is distantly related to the nuclease domain of RecB, a subunit of the bacterial RecBCD recombinase. This nuclease was discovered over 2 decades ago during a biochemical chromatographic survey of yeast exonucleases and was called exonuclease 5 (3). Initial studies with a partially purified enzyme preparation showed it to be a 5’ exonuclease specific for single-stranded DNA (ssDNA). Here we report the identification of the EXO5 gene and describe comprehensive biochemical and genetic studies that show a critical role for EXO5 in mitochondrial DNA maintenance, presumably through the processing of replication intermediates. Upon deletion of EXO5 or inactivation of its nuclease activity, only ρ− mutants could be recovered. EXO5 has previously been characterized as DEM1 (defects in morphology).
because the deletion mutant shows defects in growth and in mitochondrial morphology (10, 12). No nuclear defect associated with an EXO5 deletion has been detected.

MATERIALS AND METHODS

Plasmids and oligonucleotides. Plasmid pBL253 contains the YBR163w (EXO5) open reading frame (ORF) together with 249 nucleotides (nt) of 5' untranslated sequence and 300 nt of 3' untranslated sequence cloned into centromere vector yCP50 (ARS1 CEN4 URA3). Plasmid pBL256 contains the same EXO5 region cloned into centromere vector yCpa2 (ARS1 CEN4 TRP1). Plasmids pBL256-270 and -320, with active-site mutations exo5-D320A and exo5-D320H, respectively, were made by site-directed mutagenesis, and the correct sequences were confirmed by sequencing of the entire gene. Plasmid pBL254 (bluescript 2 um ori HIS3 M13 ori GAL1-10 GST-EXO5) contains the Schistosoma japonicum glutathione S-transferase (GST) gene fused to the N terminus of the EXO5 gene in vector pRS424-GALGST (5). The GST tag is separated from the EXO5 gene by a recognition sequence for the human rhinoviral 3C protease (LEVLFQGP). After cleavage by the protease, the N-terminal sequence of the EXO5 polypeptide is extended with the GEF sequence. Plasmids pBL254-270 and -320 have active-site mutations as in pBL256. Plasmids and sequences are available upon request.

Oligonucleotides were purchased from IDT (Coraville, IA) and purified by high-performance liquid chromatography or urea-polyacrylamide gel electrophoresis (PAGE) (v31, GCCCATCAACGTTCCAGACC; c41, GGTCTGGAAACGTTGATGG; c44, GGTCTGGAACGTTGATGG; c43, GGTCTGGAAACGTTGATGG; c44, GGTCTGGAAACGTTGATGG; c43, GGTCTGGAAACGTTGATGG). PAGE (PAG) of v31, GCCCATCAACGTTCCAGACC; c41, GGTCTGGAAACGTTGATGG; c44, GGTCTGGAACGTTGATGG; c43, GGTCTGGAAACGTTGATGG. The 5' 32P-labeled oligonucleotide was further purified on a 20-cm column, and washed at 1 ml/min with 100 ml of buffer A 300. Bound chaperones, as gene replacements with KanMX4 in BY4741 (MATa his3-D1 leu2-D1 met15-D0 ura3-D0), strain PY207 (MATa his3-D1 leu2-D1 met15-D0 ura3-D0), strain PY207 (MATa his3-D1 leu2-D1 met15-D0 ura3-D0) strand transferase or the human rhinoviral 3C protease and, after dilution with buffer A to equal buffer A 300, loaded onto a 1 ml heparin-Sepharose column. Protein was eluted with a linear gradient of buffer A100 to buffer A300. Pure Exo5 eluted at 250 to 300 mM NaCl. Mutant forms of Exo5 were purified similarly using the GST-Sepharose step.

Enrichment of Exo5 from yeast extracts. Yeast single-ORF deletion derivatives of BY4741 were grown in 1 liter of YPD (2% peptone, 1% yeast extract, 2% glucose) to mid-log phase. The cells were harvested, and extracts were prepared by homogenization with glass beads, polyvinyl precipitation, and ammonium sulfate precipitation as described previously (5). Protein (1 mg) in buffer A was gently agitated with 2 ml of S-Sepharose plus 200 mM NaCl in a total volume of 1.5 ml. Under these buffer conditions, Exo5 does not bind to either matrix. Beads were spun down, and the supernatant was used for nucleases assays with 5'-32P-labeled dT12 substrate.

Exonuclease assays. The standard 20-pll assay mixture contained 20 mM HEPES-NaOH (pH 7.6), 20 plg/ml bovine serum albumin, 1 mM DTT, 5 mM Mg-acetate, 50 mM NaCl, 100 fmol of 32P-labeled oligonucleotide substrate, and enzyme. Incubations were at 30°C for the indicated times. Variations from the standard assay are indicated in the legends to the figures. Reactions were stopped with 10 mM (final concentration) EDTA plus 0.2% SDS and analyzed by thin-layer chromatography (TLC) on polyethyleneimine-cellulose in 0.7 M LiCl or stopped with 10 mM (final concentration) EDTA plus 40% formamide and analyzed by 20% PAGE–7 M urea electrophoresis. After the TLC plate or gel was dried, it was subjected to Phosphorimagery analysis.

PCR and Southern analysis. Total cellular chromosomal DNA preparations were made from 5-ml cultures grown overnight in selective medium (select). These were subjected to 20 cycles of PCR in a 50-pll reaction mixture with 100 ng of chromosomal DNA and 50 pmol each of primers ori-pnm (GGGGGTTCAATTATTTCCT and TAGGGGAGGGTGGGTG), ori-pnt (primers complementary to ori-pnm: GAAAAATAATTTGGGACCCCC and ACCACCCGCT GACGGC), and CEN4 (primers complementary to CEN4: CAGCCCCCGCATTTTCGAGCCAGCAC and GCACCCCGCAGAAAATACCAAACTCTATGAT, and CEN5 (GATGATCATAA GGAGAGACTGTCACGAGGAGGGAGG))). Analysis was on a 2.5% agarose gel.

To label the ori sequences, the isolated ori3 or ori5 PCR fragment was reamplified by PCR with ori primers but with 10 ml [a-32P]dATP and the other three deoxynucleotide triphosphates at 200 pM. The labeled ori3 and ori5 fragments were mixed and used as a probe of EcoRV-cut total cellular DNA. EcoRV cuts either origin near the end of the fragment (see Fig. 6).

RESULTS

Identification of the S. cerevisiae EXO5 gene. Previously, we had partially purified Exo5 from yeast and established some of the properties of the enzyme (3). Specifically, we noted that the enzyme preferentially degraded ssDNA and produced dinucleotides as the main products of digestion. In order to further studies of this unique enzyme and gain an understanding of its physiological role in the cell, we first identified the gene for Exo5.

The enzyme was partially purified from yeast as described before, through four chromatographic steps (3). This preparation was further fractionated on a MonoQ column. Unfortunately, sodium dodecyl sulfate (SDS)-PAGE analysis still did
not allow positive identification of a protein band that comigrated with exonuclease activity. Therefore, the fraction with the highest exonuclease activity and a neighboring fraction with much reduced enzyme activity but with a similar protein banding pattern were fractionated by SDS-PAGE and each of the two lanes was cut into 15 slices and subjected to liquid chromatography-tandem mass spectrometry analysis. Six yeast proteins (Ade3, Tos9, Ngl1, Rfx1, Dem1, and Gon3) that were present in greater abundance in the active lane than in the lane without activity were identified. In order to determine which one corresponded to Exo5, we obtained deletion strains for each one of these genes (none of the genes identified was essential for growth). Extracts were prepared, partially enriched for Exo5, and assayed for activity. Only the extract of the dem1Δ mutant strain lacked a dinucleotide-producing activity (Fig. 1A). This analysis suggests that the DEM1 gene is required for Exo5 activity, and the simplest explanation is that it actually encodes Exo5.

An analysis of yeast protein databases revealed some information about YBR163w = EXO5 = DEM1. The deletion mutant is respiration deficient and shows morphological defects of the mitochondrion, hence the original gene designation DEM1 (defects in morphology) (12). Consistent with a mitochondrial function for EXO5/DEM1, the localization of the protein is mitochondrial (18). In support of this observed localization, the protein sequence contains a strong mitochondrial localization signal, using several prediction programs (MitoProt, Predotar), predicting cleavage at Ser26/Leu27 during mitochondrial import.

Overproduction and purification of Exo5 and Exo5 mutants. Exo5/Dem1 was overproduced in yeast from a multicopy plasmid with the EXO5/DEM1 gene placed under the control of the galactose-inducible GAL1-10 promoter. A cleavable GST tag was added to aid in purification (see Materials and Methods). Following glutathione affinity column chromatography, the GST tag was proteolytically cleaved and Exo5/Dem1 was further purified by heparin-agarose chromatography to more than 98% purity (Fig. 1B). Upon SDS-PAGE analysis, the protein migrated as a 64-kDa protein, in agreement with the size calculated from the ORF sequence (67 kDa). A gel filtration analysis showed that Exo5 migrated as a 58-kDa protein, indicating that Exo5 is a compact monomeric enzyme (Fig. 1C).

An analysis of the gel filtration fractions, using a 5′-labeled oligonucleotide as the substrate, showed potent exonuclease activity that comigrated with the protein peak (data not shown). In agreement with previous studies of the partially purified enzyme, dinucleotides were the main products of digestion (Fig. 2C and 3). Since these studies strongly suggest that the DEM1 gene actually encodes exonuclease 5, we will from now on refer to this gene as EXO5.

We subjected EXO5 to a PSI-BLAST analysis, followed by a protein threading analysis (http://toolkit.tuebingen.mpg.de/hhpred) (32). The PSI-BLAST analysis identified EXO5 as the member of a poorly conserved protein family. Interestingly, the threading analysis that was carried out with the consensus sequence obtained from the PSI-BLAST analysis yielded bacterial RecB-type nuclease as top-scoring structural homologs of EXO5 (Fig. 2A). Among the putative homologs in other model eukaryotic organisms are ORFC185.02 in Schizosaccharomyces pombe and C1orf176 in humans, both uncharacterized ORFs. While the conservation with E. coli RecB was restricted to just three small motifs in the C-terminal domain of RecB (Fig. 2A), the members of the eukaryotic Exo5 family showed conservation of a large number of small motifs over a core domain of ~250 amino acids (alignment not shown). Remarkably, a set of conserved cysteine residues in the C-terminal domain of the Exo5 members was conserved with the AddB nuclease domain of the Bacillus subtilis AddAB recombinase (36). In AddB, these cysteines coordinate an iron-sulfur cluster, suggesting that the Exo5 class may also possess this structural domain.

The first and second motifs conserved with the E. coli RecB nuclease are in two β strands that contain the two aspartates that chelate the divalent metal ion in the active site of the nuclease (31). The third conserved motif is in an α helix that overlies the active-site aspartates (Fig. 2B). Although the function of this α helix is not known, the proximity of the invariant glutamine and tyrosine to the active site is remarkable. Previous studies of RecB had shown that mutation of D1080, one of
FIG. 2. Mutational analysis of EXO5. (A) Threading analysis of Exo5. Motifs conserved with E. coli RecB nuclease are shown. Sc, S. cerevisiae; Sp, S. pombe; Hs, Homo sapiens. (B) Active site of the RecB nuclease domain from the crystal structure of E. coli RecBCD (31). The active-site divalent metal ion (black) is coordinated by Asp1067 and Asp1080 (dark gray), which are part of the two gray beta strands, 1057 to 1072 and 1078 to 1082, respectively. The active site is closed off by the aspartates that ligate the divalent metal ion, abolishes the exonuclease activity of RecB (37). Therefore, we mutated either putative active-site residue Asp270 or Asp320 to alanine, overproduced and purified the mutant proteins analogously to the wild type (Fig. 1B), and determined the resulting exonuclease activity. No exonuclease activity was detectable in Exo5-D270A, while that of Exo5-D320A was less than $10^{-4}$ times that of the wild type (Fig. 2C). This analysis not only confirms that EXO5 actually encodes the exonuclease but also lends strength to the threading analysis indicating that Exo5 belongs to the RecB family of nucleases.

**Exo5 is a single-strand-specific 5′ exonuclease.** Previous enzymatic studies with a partially purified preparation of Exo5 showed that the enzyme is specific for ssDNA and releases primarily dinucleotides as products. (3). We carried out a more thorough investigation of the enzymatic properties of the pure enzyme. First, we tested whether Exo5 has endonuclease activity. A 34-mer oligonucleotide was circularized, and enzyme. First, we tested whether Exo5 has endonuclease activity. No exonuclease activity was detectable in Exo5-D270A, while that of Exo5-D320A was less than $10^{-4}$ times that of the wild type (Fig. 2C). This analysis not only confirms that Exo5 actually encodes the exonuclease but also lends strength to the threading analysis indicating that Exo5 belongs to the RecB family of nucleases.

Exo5 yielded mainly the dinucleotide as a product, while longer products, 4 to 7 nt in length, made up a total of $\sim 5\%$ of the digestion products (Fig. 4A, lanes 3 to 6). These longer products disappeared at longer incubation times. On the other hand, incubation of the 3′-32P-labeled oligonucleotide under the exact same conditions generated a ladder of intermediates of odd lengths, consistent with a model in which the enzyme sequentially releases dinucleotides from the 5′ terminus of the 21-mer oligonucleotide (Fig. 4B, lanes 4 to 7). The starting 21-mer oligonucleotide was contaminated with $\sim 10\%$ 22-mer due to the addition of two dAMP residues during labeling (Fig. 4B, lane 3). This contamination likely accounted for the presence of small amounts of even-sized oligonucleotides in the digest. These data strongly indicate that Exo5 is a 5′ exonuclease and releases dinucleotides as main products of catalysis.

Exo5 showed no detectable nuclease activity when the 5′-32P oligonucleotide was completely double stranded (Fig. 4A, lane 12). Activity at the 5′-32P-labeled end was not affected when just the 3′ end was made double stranded by hybridization of a 14-mer, leaving 6 nt single stranded at the 5′ end (Fig. 4A, lanes 7 to 10). However, no dinucleotide was produced when only the 5′ end was made double stranded (Fig. 4A, lane 11). Interestingly, a 16-mer product was made at an $\sim 50\%$-reduced rate compared to the single-stranded control. This 16-mer could not have resulted from endonuclease activity since Exo5 has none (Fig. 3). However, generation of the 16-mer product could have resulted from 3′ loading of Exo5 with low efficiency, followed by cutting 4 nt in from the terminus or, alternatively, Exo5 could have loaded at the double-stranded end at a reduced rate and, after sliding across the double-stranded DNA (dsDNA) region, cut 2 nt downstream from the dsDNA-ssDNA junction. Based upon studies with the...
The 3'-labeled substrate discussed below, we conclude that the latter model is most likely.

Next, we assayed enzymatic activity on 3'-end-labeled substrates. No nuclease activity was detected on the fully double-stranded substrate (Fig. 4B, lane 13). When only the 3' end was made double stranded, the primary product was 17 nt in length and was followed by a 15-mer at a reduced rate. This again indicates that Exo5 digests from the 5' end but cuts with limited efficiency at the dsDNA-ssDNA junction (Fig. 4B, lanes 8 to 11). When just the 5' end was made double stranded, di- and trinucleotides were produced, similar to the results observed with the single-stranded oligonucleotide, but they were produced at an ~50-fold reduced rate (Fig. 4B, lane 12). Together with the data from the 5'-labeled substrate, these data are most consistent with the ability of Exo5 to load at dsDNA ends with low efficiency and slide across dsDNA, followed by initiation of degradation at a position 2 nt down from the dsDNA-ssDNA junction.

To determine Exo5 activity at or close to junctions, we carried out a comprehensive analysis with different partially double-stranded substrates (summarized in Table 1). A partially dsDNA substrate with a 4-nt 5' overhang is cut as efficiently as ssDNA (compare entry 5 with entry 1); however, a 2-nt 5' overhang shows only 2.8% activity and a single-nucleotide overhang is inactive (entries 4 and 3). Exo5 also cuts pseudoknots with a similar substrate preference, showing full activity as long as 4 nt of the 5' strand are single stranded (entry 7). As expected from the substrate specificity displayed by the enzyme, model Holiday junctions were not cut by Exo5 (data not shown).

Exo5 slides across RNA regions to engage downstream DNA. Incubation of Exo5 with labeled RNA yielded little or no digestion products, indicating that the enzyme shows sugar specificity (data not shown). However, how does the enzyme react with chimeric RNA-DNA molecules such as those that might arise as a result of RNA-primed DNA synthesis? Exo5 was incubated with oligonucleotides with increasing-length sections of 5' RNA (Fig. 5A). Remarkably, while incubation with pT12 released the dinucleotide pTpT, incubation with pUT11, pU2T10, and pU3T9, yielded tri-, tetra-, and pentanucleotides, respectively, indicating that the enzyme cuts selectively 2 nt downstream of the RNA-DNA junction. Since we know from the analyses described above that Exo5 shows neither endonuclease activity nor 3' exonuclease activity, we conclude that Exo5 binds to 5' RNA termini and, after sliding across the RNA substrate, cuts the DNA 2 nt from the RNA-DNA junction. The rates of hydrolysis do not decrease significantly with the addition of increasing 5' RNA sections, indicating that loading of Exo5 at the 5' terminus and sliding across RNA do not constitute a rate-limiting step in the reaction. This type of analysis was repeated with a chimeric oligonucleotide containing 10 ribonucleotides, followed by 11 deoxyribonucleotides, and in addition a 5' biotin label that allows blocking of the 5' end by streptavidin binding (Fig. 5B). With the unblocked RNA10DNA11, oligonucleotide, the major cut site was at position 12, 2 nt downstream of the RNA-DNA junction, and the rate of cutting was reduced to about 50% of that of the comparable DNA oligonucleotide (compare lanes 8 and 9 with lanes 2 and 3). However, when the 5' end was blocked with a biotin-streptavidin moiety, only residual nuclease activity was observed, indicating that Exo5 has to load at the 5' RNA end and slide across the RNA prior to cutting downstream DNA (lanes 14 and 15). As demonstrated above, converting the oligonucleotides to a double-stranded state abrogated all nuclease activity (lanes 4 and 5, lanes 10 and 11, and lanes 16 and 17). This specificity suggests that if Exo5 were involved in the processing of RNA-primed DNA replication intermediates, it would require prior strand displacement synthesis by the DNA polymerase or the participation of a 3'-5' helicase, e.g., Hml, in order to generate the appropriate substrate for digestion (30).

Mitochondrial defects of EXO5 mutants. The large-scale gene deletion project already indicated that EXO5 (DEM1) deletions are respiration deficient (12). To confirm this mitochondrial phenotype, we made a his3Δ::HIS3 diploid strain heterozygous for EXO5/exo5Δ::HIS3. After sporulation, tet-
rads were dissected onto YPD plates. A 2:2 segregation was observed for histidine prototrophy, and all His\(^+\)/H11001 colonies failed to grow on YPG plates, i.e., containing 5% glycerol as a non-fermentable carbon source, while all His\(^+\)/H11002 colonies did grow on YPG (data not shown). The tetrad analysis was repeated, but this time spores were directly germinated onto YPG plates. Each of the 14 tetrads dissected yielded a maximum of two colonies that were all His\(^+\)/H11002. Therefore, even under selective conditions, cells lacking EXO5 are respiration deficient.

In order to determine whether the nuclease activity of Exo5 was required to maintain respiration proficiency, we first made strain PY209, which is exo5\(^+/H9004::HIS3\) but is wild type (\([\rho_0]/H11001\]) because of the presence of a complementing EXO5 plasmid with a URA3 selectable marker. We then transformed this strain with a TRP1 plasmid with wild-type EXO5, exo5\(^{-}D270A\), exo5\(^{-}D320A\), or the empty vector and asked whether the URA3 plasmid could be evicted by growth on 5-fluoroorotic acid (FOA) media, either on plates containing glucose or on plates containing glycerol. While all four strains produced colonies on FOA-glucose plates, only the strain with wild-type EXO5 produced colonies on FOA-glycerol plates (Fig. 2D). This analysis shows that the nuclease activity of Exo5 is essential for respiration proficiency and therefore indicates an essential role for Exo5 exonuclease activity in the maintenance of the mitochondrial genome.

### TABLE 1. DNA substrate specificity for Exo5

<table>
<thead>
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<th>Expt</th>
<th>DNA</th>
<th>Sequence</th>
<th>% Activity</th>
</tr>
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<tr>
<td>1</td>
<td>v31</td>
<td>5'-GCCCAT---------</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>c41</td>
<td>3'-CGGGTA---------</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>v31</td>
<td>5'-GCCCAT---------</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>c42</td>
<td>3'-GGGTA---------</td>
<td>0.05</td>
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<tr>
<td></td>
<td>v31</td>
<td>5'-GCCCAT---------</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>c43</td>
<td>3'-GGTA---------</td>
<td>2.6</td>
</tr>
<tr>
<td></td>
<td>v31</td>
<td>5'-GCCCAT---------</td>
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</tr>
<tr>
<td>5</td>
<td>c44</td>
<td>3'-TA---------</td>
<td>78</td>
</tr>
<tr>
<td></td>
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<td>6</td>
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<td></td>
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<tr>
<td>7</td>
<td>c48</td>
<td>3'-TGCCGTATA------</td>
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\(^a\) Standard assay mixtures contained 100 fmol single-stranded 5'-labeled oligonucleotide v31 (experiment 1) or 100 fmol 5'-labeled oligonucleotide v31 hybridized to 200 fmol of the indicated oligonucleotide and either 10, 100, or 1,000 fmol of Exo5 for 1, 3, or 10 min (see Materials and Methods). One hundred percent activity corresponds to 0.14 dinucleotide released per second per Exo5 molecule.

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**FIG. 4.** Substrate specificity of Exo5. (A) 5'-labeled substrates. 5'-\(^32\)P-labeled v31 was either single stranded (a) or hybridized to a twofold excess of oligonucleotide c50 (b), oligonucleotide c49 (c), or oligonucleotide c41 (d). (B) 3'-labeled substrates. 3'-\(^32\)P-v31-32P-dA was either single stranded (a) or hybridized to a twofold excess of oligonucleotide cT50 (b), oligonucleotide c49 (c), or oligonucleotide cT41 (d). An asterisk indicates the position of the label. All of the assay mixtures contained 10 nM labeled substrate, 200 mM NaCl, and 10 nM Exo5 for the indicated times at 30°C. Assay mixtures were analyzed on 7 M urea-20% polyacrylamide gels. PDE, partial digestion with snake venom phosphodiesterase, a 3' exonuclease; RecJ, partial digestion with E. coli RecJ, a 5' exonuclease. A dotted guide line is added in panel B. The values on the left of each panel are oligomer sizes.
Analysis of petite phenotypes in exo5Δ yeast. Yeast strains can become petite because of rearrangement or partial deletions of the mitochondrial genome (rho−) or because of the deletion of all mitochondrial DNA (rho0). The latter phenotype is, for example, observed in mip1Δ mutant strains defective for the mitochondrial DNA polymerase. In petites that arise from gross deletions in the genome (rho−), the DNA region retained is rapidly amplified, resulting in a total mitochondrial genome. Control strains were wild type (rho0) strains of yeast (15, 22).

We generated petites by eviction of a complementing EXO5 plasmid from an exo5Δ mutant strain. The colonies arising from 11 independent events were picked and grown on YPD for 30 generations by serial dilution. Total DNA was prepared, and the mitochondrial genome was examined by PCR with sets of primers that map to different regions of the mitochondrial genome. Control strains were wild type (rho−) and mip1Δ mutant (rho0). The primer sets map to the AI1 gene (coordinates 16.3 to 17.4 kb on the 86-kb circular mitochondrial genome) or to the COX2 gene (coordinates 73.8 to 75.2 kb).

We also used a primer set (ori-in) that amplifies the various ORI sequences that are located throughout the genome. They show a high degree of sequence identity (~80%), and the A/T-rich cores of all ORI sequences are bordered by identical G/C-rich sequences.

ORI sequences are found as amplified repeats in [rho−] strains of yeast (15, 22).

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Amplification of a particular ORI sequence in petites generally produces head-to-tail repeats of the ORI sequence with variable neighboring DNA. These neighboring sequences should be identifiable by PCR analysis with a primer set complementary to the ori-in primers (ori-out, Fig. 6A). Indeed, PCR with the ori-out primers showed prominent products for each isolate, confirming that the generation of petites had occurred through head-to-tail amplification of an ORI sequence-containing region. Remarkably, the ori-out PCR consistently identified products with a very distributive size distribution (Fig. 6B). This diffuse distribution could result either from a variability in the repeat lengths generated during establishment of the petite genome or from a general genomic instability in the exo5Δ petite cells during continued growth (Fig. 6B).

A Southern blot analysis was carried out with total mitochondrial DNA isolated from these clones and digested with restriction endonuclease EcoRV by using ori3 and ori5 DNAs as hybridization probes. This analysis confirmed the pattern of strong mitochondrial repeat length variability (Fig. 6C). Some isolates also showed a
reduction in hybridization strength, particularly isolates 2 and 11, which is presumably due to partial loss of the mitochondrial genome in subpopulations of cells. Furthermore, as expected, isolates 4 and 7, which were $\rho^0$, showed no hybridization signal.

We reasoned that in the absence of Exo5, each mitochondrial replication cycle might be associated with a high probability of genome rearrangements and duplications, leading to a dynamic variability in the sizes of the amp icons by PCR and by Southern analysis. We sequenced the ori-out PCR products from the nine clones shown in Fig. 6B (right panel). Overlapping sequences were identified in all of the clones, prohibiting accurate sequence assignment. However, for several isolates, a specific sequence predominated and could be mapped to regions directly to the left and right of either ori3 or ori5. Importantly, this analysis showed that the sequences flanking ori3 or ori5 contained additional internal repeat sequences between 50 and 200 nt in length, suggesting that recombination frequently occurred in the exo5Δ clones (Fig. 6D). We confirmed the high-instability phenotype of exo5Δ mutants by carrying out an experiment in which two independently isolated exo5Δ clones, one with ori3 amplified and one with ori5 amplified, were serially propagated and expanded through successive growth cycles, and individual clones were analyzed by PCR. The two clones produced after loss of EXO5 persistently altered their ori-out PCR patterns upon serial propagation (Fig. 6E).

We next tested whether the reintroduction of EXO5 into the petite clones would cause stabilization of the petite genomes.
FIG. 7. exo5Δ mutant strains are not damage sensitive. All strains were mip1Δ ([rho0]) and, in addition, had the indicated genotype. Serial 10-fold dilutions of late-log-phase cells, from 10⁵ to 10⁷ cells per spot, were spotted onto YPD plates or YPD plates containing the indicated concentrations of hydroxyurea (HU, mM) or camptothecin (CPT, μg/ml). Some YPD plates were irradiated with the indicated fluency of UV254. Plates were grown for 3 days at 30°C and photographed. WT, wild type.

An EXO5-complementing plasmid was transformed back into randomly selected petite isolates, and the transformants were analyzed by PCR analysis with the ori-in and ori-out primers (Fig. 6F). Remarkably, the ori-out PCR product distribution had narrowed drastically, indicating that a stable repeat pattern had been established in the Exo5-containing transformants. Sequence analysis of the repeat unit identified unique wild-type sequences that border ori3 or ori5 on the left and right. The junction between the upstream and downstream ORI sequences varied from clone to clone. However, invariably, this junction was formed by a sequence 7 to 13 nt in length that was present in both the upstream and downstream mitochondrial genomic sequences. This observation strongly suggests that junction formation had occurred through recombination between these small identical sequence motifs.

Petite mutants can exhibit yotic suppressiveness, a phenomenon in which, upon mating with a wild-type [rho+] strain, the wild-type mitochondrial genome is excluded from diploids; i.e., the diploids are also [rho–] (13). Hypersuppressiveness is a consequence of the large number of ORI sequence repeats in the [rho+] strain. Both RNA-primed replication and recombination mechanisms contribute to hypersuppressiveness (22, 24). The exo5Δ clones lacking mitochondrial DNA (clones 4 and 7 in Fig. 6B) showed no hypersuppressiveness upon mating with a [rho+] strain; i.e., all diploids were wild type [rho+]. In contrast, the other clones showed high degrees of hypersuppressiveness, from 80% to 100%; i.e., up to 100% of the diploids were [rho–], in agreement with similar findings by others (22, 24).

EXO5 deletion shows no nuclear phenotypes. We explored the possibility that Exo5 may also reside in the yeast nucleus at low levels, even though it was only detected in the mitochondria (18). However, our genetic analysis identifying a potential nuclear function for Exo5 was uniformly negative. To eliminate phenotypes due to mitochondrial dysfunction, we carried out all of our studies in a [rho+] mip1Δ mutant background. We compared the growth and DNA damage survival of exo5Δ mip1Δ double mutants with mip1Δ mutants as a control. In addition, we combined these mutants with deletions of the additional nuclease genes RAD27, EXO1, MUS81, and MRE11. The damage sensitivities of the triple mutants were compared with those of the relevant double mutants (Fig. 7). FEN1, encoded by RAD27, is primarily involved in Okazaki fragment maturation and base excision repair, Exo1 is involved in mismatch repair and double-strand break repair, Mus81 is involved in the processing of stalled replication forks, and Mre11 is involved in double-strand break repair (reviewed in references 2, 6, and 34). Serial dilutions were plated on YPD plates containing hydroxyurea or camptothecin, or plates were UV irradiated. No significant differences in damage sensitivity were observed between a given mip1Δ nuclease double mutant and the exo5Δ mip1Δ nuclease triple mutant. Therefore, we conclude that either EXO5 is not involved in nuclear DNA maintenance or a very efficient redundancy with other nucleases exists in the DNA damage repair pathways tested.

DISCUSSION

Exo5 has the remarkable ability to slide over RNA and less efficiently over dsDNA to initiate cutting at a position 2 nt downstream of the RNA-DNA or dsDNA-RNA junction. The ability to differentiate between RNA and DNA but cut ssDNA flaps efficiently through sliding may be important for the maturation of intermediates during mitochondrial DNA replication. Since Exo5 does not degrade dsDNA or dsRNA-DNA, the generation of such single-stranded flaps would have to be accomplished either through strand displacement synthesis by the mitochondrial DNA polymerase γ or through the participation of a 3′-5′ helicase. The first is an important Okazaki maturation mechanism in the nucleus, where substrates for cutting by FEN1 and by Dna2 are generated through strand displacement synthesis by polymerase δ (reviewed in reference 2). Alternatively, the single-stranded flap could be generated through helicase action. The Hml1 helicase has the correct directionality (3′-5′) to generate substrates for Exo5, and it is also essential for maintenance of the wild-type mitochondrial genome but not for that of [rho–] petites (21, 26). Unfortunately, very little is known about the mechanism of replication of the yeast mitochondrial genome. Replication may occur through several mechanisms, including initiation by RNA polymerase and by double-strand break-induced recombinational mechanisms that can mature into rolling-circle replication (22, 28, 35). Sufficient replicative ability must remain in an exo5Δ mutant strain to allow replication of the ORI repeat sequences, since these cells are [rho+] and not [rho0]. However, the observations that 2 out of the 11 petite isolates studied lost their mitochondrial DNA and 2 more were severely depleted for mitochondrial DNA indicate that even maintenance of the [rho+] genome is severely compromised (Fig. 6B and C).

An alternative function for Exo5 could be in recombination. We think this is unlikely because of the extreme repeat variability displayed in the exo5Δ mutant strains that can only be a consequence of very active recombination. All of the [rho+] clones that we generated by evocation of EXO5 had amplified either ori3 or ori5, together with variable-length sequences surrounding either ORI sequence. These regions showed high variability during propagation, indicative of very active
recombination (Fig. 6B, C, and E). In addition, sequences inside the repeat unit were subject to additional partial duplication events, suggesting a hyperrecombination phenotype in the mutants (Fig. 6D).

The generation of conditional mutations in EXO5 should be of invaluable help in delineating its contribution to mitochondrial genome maintenance. As Exo5 is conserved in other organisms, including humans (Fig. 2A), resolving the contribution of yeast EXO5 may have important implications for understanding genome stability in higher organisms.

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