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13. ABSTRACT (Maximum 200 Words) Telomeres are repeated double stranded DNA sequences at chromosome ends. Replication of telomeres requires telomerase. Recent data suggest telomere length maintenance might be involved in the origin or progression of cancer and aging. Understanding of telomere replication could offer new targets for drug screen. The conservation of telomerase and telomere structure has made yeast a good system to study the problem. An essential yeast telomere binding protein Cdc13p appears to play a key regulatory role in telomere replication. Identifying Cdc13p interacting proteins could yield important information. A two-hybrid assay was employed. Pollp, the catalytic subunit of DNA polymerase α (Pol α), and Estlp, a telomerase component, were identified. The two-hybrid interactions were also confirmed by biochemical criteria. Disrupting the Cdc13p-Pollp interaction resulted in longer telomeres, while the mutated cells appeared normal in many aspects. The length of the elongated telomeres were in proportion to the extent of the decrease in the interaction. Data suggest that Pol α is involved in telomere replication, and Cdc13p regulates telomere replication by interacting with both telomerase and Pol α . Other Cdc13p interacting proteins were also identified. Further studies are in progress.				
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1. Introduction:

The mechanism of DNA replication used by all known polymerases predicts one problem: at the end of DNA replication, after removal of the last RNA primer, there will be a loss of 7-12 nucleotides at one 5' end of a newly synthesized chromosome. This loss of DNA is compensated for by telomerase, an reverse transcriptase that uses an RNA template to extend the 3' ends. The conventional DNA polymerase can then repair the 5' ends. Since telomerase RNA templates are C-rich, the 3' ends of chromosomes synthesized by telomerase are G-rich and the strands are called G-strands. The other strands, repaired by a DNA polymerase, are called C-strands. The chromosomal ends containing the repeated sequences are called telomeres. In budding yeast, telomeres are about 300 ± 75 bps, consisting of repeated sequences $TG_{1-3}/C_{1-3}A$ (Wang et al., 1989; Wang and Zakian, 1990; Zakian, 1995).

Cdc13p is a budding yeast telomere binding protein *in vivo* and *in vitro* (Bourns et al., 1998; Lin and Zakian, 1996; Nugent et al., 1996). It appears to play a central regulatory role in telomere replication. *cdc13-2* mutant cells have the phenotype as a mutant that lack telomerase activity: telomeres shorten progressively until the cells senesce. However, extracts from these cells have normal telomerase activity *in vitro* (Nugent et al., 1996). This suggests that Cdc13p might recruit telomerase activity *in vivo*. The *cdc13-1* mutant is a temperature sensitive mutation. At the restrictive temperature, cells are arrested due to C-strand specific degradation by an unknown mechanism. This degradation starts from the C-strands of telomere ends and can go up to 20 kb into the chromosomes (Garvik et al., 1995). Cdc13p therefore appears to protect the telomeres from degradation. Studying Cdc13p interacting proteins might yield important information about telomere replication and regulation.

2. Body:

Since joining in Dr. Virginia A. Zakian's lab in 1996, I have focused on studying Cdc13p and its interacting partners. Using a two-hybrid assay, I was able to identify proteins that interact with Cdc13p, which include: DNA polymerase α catalytic subunit Pol1p, the essential telomere replication protein Est1p, Tao3p, Fun12p and others.

a. Cdc13p-Est1p and Cdc13p-Pol1p interaction

The Cdc13p-Pol1p two-hybrid interaction was confirmed by a biochemical assay. Pol1p could be co-immunoprecipitated with Cdc13p when both protein were expressed from their own promoters. These data suggest that the two proteins are in the same complex *in vivo* (Qi and Zakian, 2000).

Single-amino-acid mutations that disrupt the Cdc13p-Pol1p interaction were generated by a random PCR mutagenesis assay. The single-amino-acid mutations obtained in Pol1p clustered together: D236N, E238K, P241T. Each amino acid change disrupted the Cdc13p-Pol1p interaction to a different extent: D236N, 100%; E238K,

50%; and P241, 90%. These mutations were placed back into the cells and the mutated alleles have no general DNA replication defect or growth defect, but do have longer telomeres. The extent of the lengthening of the telomeres is correlates with the severity of the loss of the interaction(Qi and Zakian, 2000).

Cdc13p residues involved in Cdc13-Pol1p interaction were also mapped by the same PCR mediated random mutagenesis. They were throughout the protein. Two of the mutations were placed back into the chromosome. These mutations also showed only a long telomere phenotype(Qi and Zakian, 2000).

TPE(Telomere Position Effect on transcription) can be used as an indicator of telomere structure. The mutations described above have no effect on TPE suggesting that overall telomere structure is normal and the defect is restricted to termini(Qi and Zakian, 2000).

The Cdc13p-Est1p two-hybrid interaction was confirmed by biochemical criteria. Cdc13p can be co pulled down by GST beads in cell extracts over-expressing both Cdc13p and GST tagged Est1p. Over-expression of an Est1p fragment partially suppressed the lethality of *cdc13-1*. These data suggest that Cdc13p interacts with Est1p physically and genetically. *cdc13-2* cells appear to lose the ability to recruit telomerase *in vivo*. However *cdc13-2p* still interacted with Est1p in a two-hybrid assay and a GST-pull-down assay(Qi and Zakian, 2000). We propose that the Cdc13p-Est1p interaction results in a conformation change that is critical for recruiting telomerase activity. The *cdc13-2* phenotype is not due to the loss of the interaction with Est1p, but the inability of the mutant proteins to undergo a conformation change.

The above results suggest: 1. DNA polymerase α is the conventional enzyme involved in C-strand repair during telomere replication. 2. Cdc13p interacts with both the telomerase component Est1p and the C-strand repairing enzyme Pol1p to coordinates telomere synthesis. Two models have been proposed: cooperation and competition, to explain how Cdc13p coordinates G-strand and C- strand synthesis.

b. Cdc13p-Fun12p

Fun12p is a translation initiation factor in budding yeast(Choi et al., 1998). It was found to interact with Cdc13p in a two-hybrid screen. Using a PCR mediated random mutagenesis assay, single-amino-acid mutations in Fun12p that disrupted Cdc13p-Fun12p interaction were isolated. They are I634V, S628F, S566F and F843L. The first two were placed back into the *FUN12* locus on chromosome I. Mutants showed no growth defects and no telomere length defects. The meaning of this interaction remains unknown. Further studies are in progress.

c. Cdc13p-Tao3p

Tao3p function is unknown. Deletion of *tao3* results in cell aggregation. The C-terminus of Tao3p was isolated from a two-hybrid library screen by interacting with

Cdc13p. Deletion of this C-terminus showed the same phenotype as the null: cell aggregation. The reason for this cell aggregation is unknown. To confirm that the aggregation is due to the loss of interaction with Cdc13p, single-amino-acid mutations on Cdc13p will be screened, and protein expression will be measured. The reason of cell aggregation will be studied.

- d. two-hybrid interactions between Cdc13p and Hdf1p, Hdf2p; Pol1p and Hdf1p, Hdf2p

Hdf1p and Hdf2p are the two subunit of the Ku complex in yeast (Boulton and Jackson, 1996). Null mutant of *hdf1* or *hdf2* is viable at 30°C, but senescence quickly at 37°C (Boulton and Jackson, 1998). Telomeres are shorter than wild type, and there are constitutive single-stranded G-tails at telomeres (Gravel et al., 1998). The constitutive G-tails could be due to in C-strand synthesis by Pol α or due to a failure in C-strand protection. I tested the two-hybrid interactions between Cdc13p and Hdf1p, Hdf2p, Pol1p and Hdf1p, Hdf2p. The results were all negative. I also made the double mutation *hdf1::HIS3 pol1-236*, reasoning that if Hdf1p is in the same pathway as Cdc13p in terms of recruiting Pol α activity, the double mutants would have a more severe phenotype. They might senesce faster and have shorter telomeres. These results were also negative. The double mutation senesced slower and had slightly longer telomeres than *hdf1* null. These data suggest that Hdf1p, unlike Cdc13p, is not involved in C-strand synthesis. It is more likely involved in C-strand protection.

- e. Two-hybrid interaction between Cdc13p and Stn1p, Pol1p and Stn1p

Telomere protein Stn1p is a Cdc13p interacting protein (Grandin et al., 1997). The *stn1-13* mutant has a similar phenotype as the *cdc13-1*. These two proteins might potentially form a complex *in vivo*. Therefore, Stn1p might be also involved in recruiting Pol α activity. To test this possibility, full-length *STN1* was cloned into the prey vector in the two-hybrid system. Consistence with C. Nugent's data (Personal communication), it interacts with Cdc13p and Cdc13-1p, but not with Cdc13-2p. Stn1p also interacts with the N-terminal fragment of Pol1p in the two-hybrid assay. I will test if the interaction is independent of Cdc13p and if the interaction is a direct physical interaction.

Key Research Accomplishments

- Identified an interaction between Cdc13p and Pol1p, the catalytic subunit of DNA polymerase α .
- Identified an interaction between Cdc13p and Est1p, an essential protein for telomerase activity.
- Showed that Cdc13p and Pol1p are in the same complex in vivo by immunoprecipitation.
- Showed that yeast cells develop abnormally long telomeres with no noticeable growth defect if they express any one of the mutant *pol1* proteins or *cdc13* mutant proteins.
- Concluded that the Pol1p subunit of DNA polymerase α is responsible for C-strand re-synthesis during telomere replication.
- Suggested that telomere length is regulated by the interactions of Cdc13p with Pol1p and Est1p to balance opposite directions of G-strand and C-strand synthesis.
- Proposed models for regulation: Competition or Coordination.
- Proposed model for Cdc13p recruiting telomerase activity: Cdc13p-Est1p interaction results a conformational change that favors the recruitment of telomerase activity.
- Discovered two-hybrid interactions between Cdc13p and Fun12p, and Cdc13p and B10p.
- Isolated single-amino-acid mutations in Fun12p that disrupted the interaction with Cdc13p.
- Showed that the deletion of the Cdc13p interacting domain in Tao3p gives the same phenotype as *tao3* null cells: cell aggregation .
- Discovered that Hdf1p is not involved in C-strand synthesis. Hypothesized that Hdf1p is involved in C-strand protection.
- Discovered a two-hybrid interaction between Pol1p and Stn1p.

Reportable Outcomes

Publication:

Haiyan Qi and Virginia A. Zakian. "The *Saccharomyces* telomere binding protein Cdc13p interacts with both the catalytic subunit of DNA polymerase α and the telomerase associated Est1 protein" *Gene & Develop.* (2000) **14**(14): 1777-1788.

Products:

Created yeast strains that express mutant forms of Pol1p, Cdc13p, Fun12p and B10p.

Conclusions:

Cdc13p plays an important regulatory role in yeast telomere replication: limits C-strand degradation, recruits telomerase and DNA polymerase α activity, and regulates G-strand and C-strand synthesis in telomere replication.

Pol α is the C-strand synthesis enzyme.

Ku complex is not involved in C-strand re-synthesis, but might be involved in C-strand protection.

References

- Boulton, S. J. and Jackson, S. P. (1996). *Nucl. Acids Res.* **24**, 4639-4648.
- Boulton, S. J. and Jackson, S. P. (1998). *Embo J.* **17**, 1819-1828.
- Bourns, B. D., Alexander, M. K., Smith, A. M. and Zakian, V. A. (1998). *Mol. Cell. Biol.* **18**, 5600-5608.
- Choi, S. K., Lee, J. H., Zoll, W. L., Merrick, W. C. and Dever, T. E. (1998). *Science* **280**, 1757-1760.
- Garvik, B., Carson, M. and Hartwell, L. (1995). *Mol. Cell. Biol.* **15**, 6128-38.
- Grandin, N., Reed, S. I. and Charbonneau, M. (1997). *Genes Dev.* **11**, 512-527.
- Gravel, S., Larrivee, M., Labrecque, P. and Wellinger, R. J. (1998). *Science* **280**, 741-744.
- Lin, J. J. and Zakian, V. A. (1996). *Proc. Natl. Acad. Sci. USA* **93**, 13760-13765.
- Nugent, C. I., Hughes, T. R., Lue, N. F. and Lundblad, V. (1996). *Science* **274**, 249-52.
- Qi, H. and Zakian, V. A. (2000). *Genes Dev.*
- Wang, S. S., Pluta, A. F. and Zakian, V. A. (1989). *Prog. Clin. Biol. Res.* **318**, 81-89.
- Wang, S.-S. and Zakian, V. A. (1990). *Mol. Cell. Biol.* **10**, 4415-19.
- Zakian, V. A. (1995). In "Telomeres" (E. H. Blackburn and C. W. Greider, eds), pp. 107-137. Cold Spring Harbor Laboratory Press, Plainview, NY.

The *Saccharomyces* telomere-binding protein Cdc13p interacts with both the catalytic subunit of DNA polymerase α and the telomerase-associated Est1 protein

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Saccharomyces telomeres consist of ~350 bp of C₁₋₃A/TG₁₋₃ DNA. Most of this ~350 bp is replicated by standard, semiconservative DNA replication. After conventional replication, the C₁₋₃A strand is degraded to generate a long single strand TG₁₋₃ tail that can serve as a substrate for telomerase. Cdc13p is a single strand TG₁₋₃ DNA-binding protein that localizes to telomeres in vivo. Genetic data suggest that the Cdc13p has multiple roles in telomere replication. We used two hybrid analysis to demonstrate that Cdc13p interacted with both the catalytic subunit of DNA polymerase α , Pol1p, and the telomerase RNA-associated protein, Est1p. The association of these proteins was confirmed by biochemical analysis using full-length or nearly full-length proteins. Point mutations in either *CDC13* or *POL1* that reduced the Cdc13p–Pol1p interaction resulted in telomerase mediated telomere lengthening. Over-expression of the carboxyl terminus of Est1p partially suppressed the temperature sensitive lethality of a *cdc13-1* strain. We propose that Cdc13p's interaction with Est1p promotes TG₁₋₃ strand lengthening by telomerase and its interaction with Pol1p promotes C₁₋₃A strand resynthesis by DNA polymerase α .

[Key Words: Telomere replication; telomerase; DNA polymerase; *CDC13*; *POL1*; *EST1*]

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In most eukaryotes, telomeres, the physical ends of chromosomes, consist of simple repeated DNA sequences and their associated proteins. Chromosomes in *Saccharomyces cerevisiae* end with ~350 bp of duplex C₁₋₃A/TG₁₋₃ DNA. In organisms where the structure at the very end of the chromosome has been examined, the G-rich strand is extended to form a single-strand tail. Telomere-associated proteins include proteins that directly bind to either duplex or single-stranded telomeric DNA as well as those that are brought to the telomere by protein–protein interactions. The major *Saccharomyces* telomere protein is the duplex C₁₋₃A/TG₁₋₃ binding protein Rap1p (Conrad et al. 1990; Wright et al. 1992; Wright and Zakian 1995). The Cdc13p protein, which binds ss TG₁₋₃ DNA in vitro (Lin and Zakian 1996; Nugent et al. 1996), is also localized to telomeres in vivo (Bourns et al. 1998). Unlike Rap1p, Cdc13p does not bind to internal tracts of C₁₋₃A/TG₁₋₃ DNA (Bourns et al. 1998), suggesting that its association with the telomere is due to its ability to bind ss TG₁₋₃ DNA.

Telomeres are required for the complete replication of linear chromosomes. After replication by a conventional DNA polymerase, removal of the terminal RNA primer

leaves an 8–12 base gap at the 5' end of newly replicated DNA molecules, a gap that can not be repaired by a conventional DNA polymerase. In most eukaryotes, including yeast, the end replication problem is solved by telomerase, a reverse transcriptase that is able to extend the short G-tail left after RNA primer removal, using an integral RNA as its template. There are at least five *Saccharomyces* genes that are required for the telomerase pathway in vivo, *EST1*, *EST2*, *EST3*, *TLC1*, and *CDC13* (for review, see Nugent and Lundblad 1998). Mutation of any one of these genes causes a gradual loss of telomeric DNA or *est* (ever shorter telomere) phenotype, consistent with incomplete replication of chromosome ends. However, the products of only two of these genes, *TLC1*, which encodes the RNA component of telomerase, and *EST2*, the gene for its catalytic component, are essential for telomerase activity in vitro (for review, see Nugent and Lundblad 1998). Est1p is associated with *TLC1* RNA (Lin and Zakian 1995; Steiner et al. 1996) and also binds ss TG₁₋₃ DNA in vitro (Virta-Pearlman et al. 1996; Zhou et al. 2000). *EST1* mutations that reduce Est1p binding to telomerase RNA have an *est* phenotype (Zhou et al. 2000). Given that Cdc13p, an in vivo telomere-binding protein (Bourns et al. 1998), is not required for telomerase activity in vitro (Lingner et al. 1997a), an appealing model is that Cdc13p recruits telomerase to the telo-

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mere. This model is supported by the demonstration that a Cdc13p-Est2p fusion protein bypasses the requirement for both Est1p and Est2p (Evans and Lundblad 1999).

Analysis of telomere replication intermediates revealed an unexpected step in telomere replication. Conventional semiconservative replication of the telomeric C₁₋₃A/TG₁₋₃ repeats occurs late in S phase (Wellinger et al. 1993a). After conventional replication is complete, both ends of individual chromosomes acquire 50–100 base ss TG₁₋₃ tails (Wellinger et al. 1993a, b). Because these G-tails are detected in cells lacking telomerase (Dionne and Wellinger 1996; Wellinger et al. 1996), they must be generated by C-strand degradation. As Cdc13p binds efficiently to both 43 and 270 base TG₁₋₃ tails in vitro (Lin and Zakian 1996), these long G-tails are suitable substrates for Cdc13p; and indeed, genetic data reveal that Cdc13p regulates C-strand degradation. When *cdc13-1* cells are grown at restrictive temperatures, they accumulate ssDNA that extends from the telomere many kilobases towards the center of the chromosome with the C-strand being preferentially degraded (Garvik et al. 1995). C-strand degradation was unanticipated as it exacerbates the end problem of replication by creating a large gap at both ends of individual chromosomes.

It is unclear when or how often telomerase acts on a given chromosome end. As the rate of telomerase lengthening is inversely proportional to telomere length (Marcand et al. 1999), it is possible that only short telomeres are substrates for telomerase. Although telomerase activity can be detected in extracts prepared from cells throughout the cell cycle, formation of new telomeres is restricted to late G₂/M phase (Diede and Gottschling 1999). Regardless of whether or not a given telomere is acted upon by telomerase, its C-strand must be resynthesized as long ss TG₁₋₃ tails are not detected in post S/G₂ phase cells (Wellinger et al. 1993a,b). Presumably, C-strand resynthesis is accomplished by a conventional DNA polymerase, although the polymerase that carries out this step has not been identified, nor is it known how DNA polymerase is recruited to the transient G-tail. After the 8–12 base RNA that is expected to prime C-strand resynthesis is removed, newly replicated telomeres should bear an 8–12 base ss TG₁₋₃ tail. As duplex telomeric DNA bearing a five or 10 base TG₁₋₃ tail is bound by Cdc13p in vitro (Lin and Zakian 1996), newly replicated telomeres are suitable substrates for Cdc13p.

Cdc13p appears to be a key regulatory protein in telomere replication as it functions at two critical steps in yeast telomere metabolism, C-strand degradation and G-strand elongation. To gain insight into how Cdc13p regulates these steps, we identified proteins that interact with Cdc13p in vivo. We found that Cdc13p interacts with the catalytic subunit of DNA polymerase α , Pol1p, and Est1p, by both two hybrid and biochemical criteria. Mutations in either *CDC13* or *POL1* that disrupt the Cdc13p-Pol1p interaction affect telomere length. Overexpression of the portion of Est1p that interacted with Cdc13p by two hybrid criteria partially suppressed the temperature sensitive lethality of a *cdc13-1* strain. The interaction of Cdc13p with Est1p provides additional

support for a model in which Cdc13p promotes the telomerase pathway by recruiting Est1p to the telomere. We propose that Cdc13p's interaction with Pol1p promotes C-strand resynthesis.

Results

Cdc13p interacts with Pol1p by two hybrid criteria

A two hybrid screen (Fields and Song 1989; Gyuris et al. 1993) was used to identify proteins that interact with Cdc13p in vivo. The first 600 amino acids of the 924 amino acid Cdc13p were expressed as a fusion to the LexA DNA binding domain and used as bait in the yeast strain YEM1 α , which carries two reporter genes, *LEU2* and *LacZ* (Fig. 1; the fusion polypeptide encoded by this plasmid is called Cdc13Np). The amino acid changes that confer the *cdc13-1* and *cdc13-2* phenotypes are both contained within this amino-terminal segment of Cdc13p (Lin and Zakian 1996; Nugent et al. 1996), and this region is sufficient to target Cdc13p to telomeres in vivo (Bourns et al. 1998). A library that produces galactose-inducible proteins as fusions of polypeptides encoded by segments of yeast chromosomal DNA and a transcriptional activation domain from *Escherichia coli* (Watt et al. 1995) was introduced into YEM1 α cells carrying the bait plasmid. If a library plasmid produces a fusion protein that interacts with Cdc13Np, it will activate the *LEU2* gene, allowing growth on media lacking leucine. A fusion protein that interacts strongly with Cdc13Np will also activate the *lacZ* gene, generating blue color when cells are assayed for β -galactosidase ac-

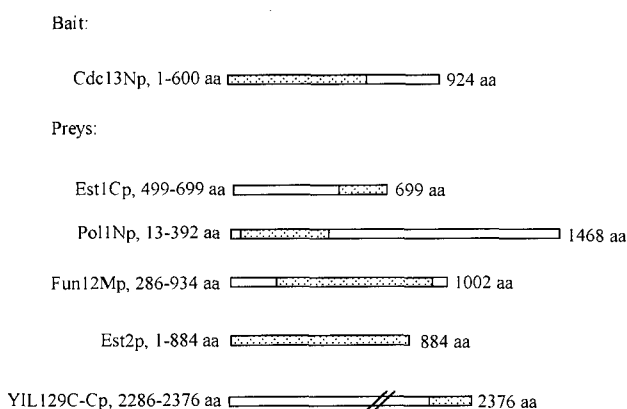


Figure 1. Polypeptides used in the two-hybrid assay. Dotted regions indicated the portion of Cdc13p expressed from the bait plasmid pEG202 and the portions of proteins expressed from the prey vector pJG4-5 (not to scale). The fusion of the amino-terminal region of Cdc13p with the LexA DNA binding domain is referred to as Cdc13Np in the text. Prey proteins are similarly named; e.g., Pol1Np refers to the fusion of 379 amino acids from near the amino-terminus of Pol1p to the activation domain peptide. The amino acid numbers after the name of each protein indicates the specific amino acids expressed from the two-hybrid vector. The numbers to the right of the schematic for each protein indicate the number of amino acids in the full-length protein.

tivity. Colonies expressing interacting fusion proteins were selected by their ability to grow on galactose media lacking leucine (Fig. 2A) and then screened for production of blue color by a filter LacZ assay (Breedon and Nasmyth 1985).

From a screening of more than 5×10^6 colonies, we obtained three positive clones. DNA sequence analysis revealed that the plasmids contained amino acids 13–392 from the 1468 amino acid Pol1p, amino acids 286–934 from the 1002 amino acid Fun12p, and amino acids 2286–2376 from the 2376 amino acid YIL129C (Fig. 1; the fusion polypeptides encoded by these prey plasmids were named Pol1Np, Fun12Mp, and YIL129C-Cp). The essential gene *FUN12* encodes a protein that is 27% identical to the *E. coli* IF2 protein and, like its bacterial counterpart, is involved in the initiation of protein synthesis (Choi et al. 1998). Virtually nothing is known about the function of YIL129C. Pol1p is the catalytic subunit of DNA polymerase α , one of three yeast polymerases required for chromosomal DNA replication (Waga and Stillman 1998). Given its central role in DNA replication and the fact that certain *POL1* mutations cause telomere lengthening (Carson and Hartwell 1985; Adams and Holm 1996), we decided to focus on the interaction between Pol1p and Cdc13p.

By the criteria of a two hybrid assay, the interaction of Pol1Np with Cdc13Np was both specific and strong (Fig. 2A). Activation was seen only on galactose medium, not for glucose-grown cells, and required the presence of both the bait and prey plasmids (negative controls 1 and 2; Fig. 2A). The *POL1* polypeptide did not activate in cells expressing other baits, such as LexA::cRafp (a gift from E. Golemis) (Fig. 2A, cRaf). Cells expressing the interacting Pol1Np and Cdc13Np polypeptides grew as well on plates lacking leucine as the strongly interacting proteins Rpb4p and Rpb7p, two subunits of RNA polymerase II (Khazak et al. 1995) (Fig. 2A, positive control), and produced a similar amount of β -galactosidase by a LacZ filter assay (data not shown).

Full-length Cdc13p and Pol1p interact by biochemical criteria

Immunoprecipitation (IP) was used to determine if the interaction between Cdc13Np and Pol1Np detected by two-hybrid analysis reflected an *in vivo* association of full-length proteins (Fig. 2B). Extracts were prepared from cells carrying the endogenous copies of both genes expressed from their own promoters with the only modification being the introduction of three MYC epitopes at the Cdc13p carboxyl terminus. Cells with the MYC₃-tagged *CDC13* allele had a normal growth rate and no detectable change in telomeric DNA (data not shown). Monoclonal anti-MYC antibody was used to immunoprecipitate MYC₃-tagged Cdc13p. The precipitate was analyzed by Western blotting using both a monoclonal anti-Pol1p (a generous gift from Dr. P. Plevani) and anti-MYC antibodies. MYC₃-Cdc13p was readily detected in the anti-MYC immunoprecipitate (Fig. 2B). Cdc13p appeared as a doublet, because of its existing in

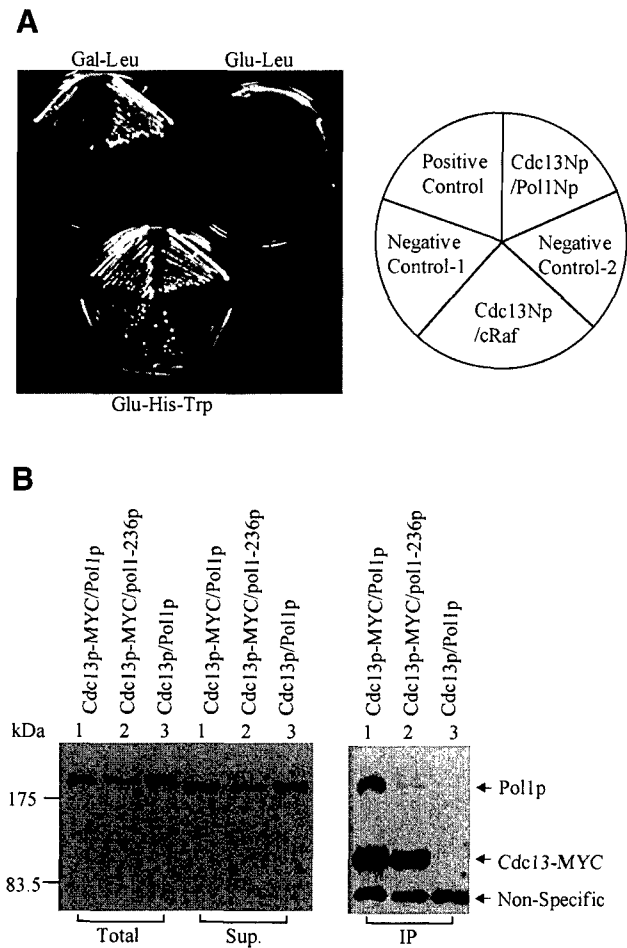


Figure 2. Cdc13p interacted with Pol1p, the catalytic subunit of DNA polymerase α . (A) Cells expressing prey and/or bait proteins were streaked on galactose plates lacking leucine (Gal-Leu). Galactose induces expression of prey proteins and growth in the absence of leucine requires interaction of bait and prey polypeptides. Alternatively, cells were streaked on glucose plates lacking leucine (Glu-Leu) where prey proteins are not expressed or glucose plates lacking histidine and tryptophan (Glu-His-Trp), which selects for maintenance of both the prey and bait plasmids but not for interaction of their products. The positive control was cells expressing two subunits of RNA polymerase II, Rpb4p and Rpb7p. The negative controls were cells expressing Cdc13Np and carrying the empty prey vector pJG4-5 (Negative Control-1), cells expressing Pol1Np and carrying the empty bait vector pEG202 (Negative Control-2), and cells expressing Cdc13Np and a LexA::cRafp fusion protein (gift from E. Golemis) (Cdc13Np/cRaf). (B) Extracts from cells carrying an MYC₃-tagged Cdc13p were immunoprecipitated by monoclonal anti-MYC antibody immobilized on protein-A and protein-G beads. The extract prior to immunoprecipitation (Total), the supernatant from the immunoprecipitate (Sup.), and the immunoprecipitate (IP) were analyzed by Western blotting using both anti-MYC and anti-Pol1p monoclonal antibodies. Although Cdc13p was not visible in the total cell extract in this gel, it was detectable when more protein was loaded. Extracts were prepared from cells containing 3XMYC (lanes 1,2) or untagged (lane 3) Cdc13p. Cells had either the wild-type *POL1* (lanes 1,3) or the *pol1-236* allele (lane 2). The nonspecific band was detected by the anti-MYC serum.

different phosphorylated states in vivo (A. Taggart and V.A. Zakian, unpubl.) (Fig. 2B, panel IP). Pol1p was detectable in the anti-MYC immunoprecipitate (Fig. 2B, lane 1 IP) but only if the extract was prepared from a strain with an MYC₃-tagged Cdc13p (Fig. 2B, cf. lanes 1 and 3, IP). We conclude that full-length Pol1p and Cdc13p interact in vivo.

Identification of Pol1Np mutations that reduce its ability to interact with Cdc13Np

To identify point mutations in Pol1Np that disrupted the Cdc13p–Pol1p interaction, we used PCR to mutagenize the 379 amino acid fragment of Pol1p identified in the two-hybrid screen (Leung et al. 1989; Zhou et al. 1991). The mutagenized fragments were then assessed for their ability to interact with Cdc13Np in the two-hybrid assay. Transformants that produced white colonies in the lacZ filter assay were chosen for further analysis. Out of ~10⁶ colonies, three mutants gave white or pale blue colored colonies in the LacZ filter assay and produced full-length polypeptides by Western analysis (Fig. 3B). As each of these mutants produced normal amounts of Pol1Np, their failure to interact in the two hybrid assay was not attributable to failure in protein expression. DNA sequencing showed that each of the three had a single amino acid substitution: aspartic acid to asparagine at residue 236 (D236N), glutamic acid to lysine at residue 238 (E238K), and proline to threonine at residue 241 (P241T), respectively (Fig. 3A). Mutations were named *pol1-236*, 238, and 241 according to the amino acid residue that was mutated.

To estimate the extent of loss of the Pol1Np–Cdc13Np two-hybrid interaction in each of the *pol1* mutants, a quantitative liquid lacZ assay was performed (Guarente 1983). Using this assay, interaction with Cdc13Np was

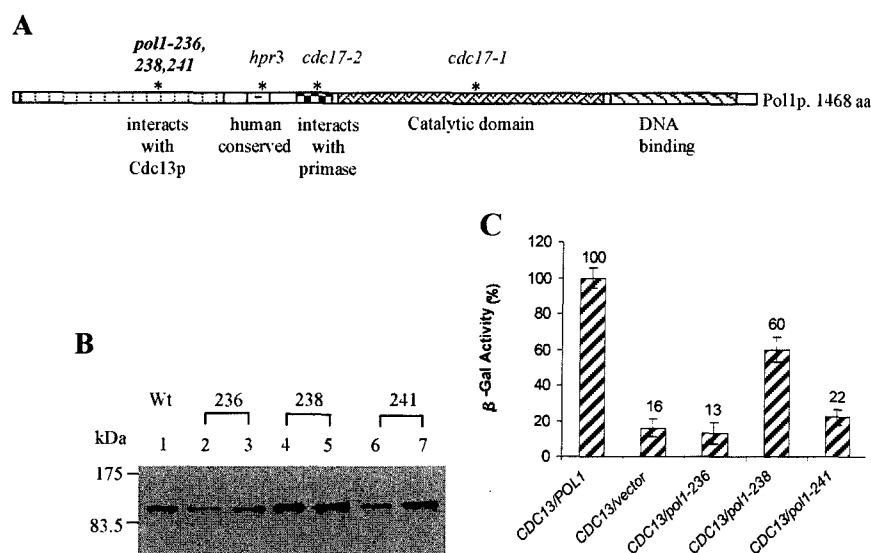
essentially abolished in *pol1-236* (D236N), reduced ~50% in *pol1-238* (E238K) and reduced ~90% in *pol1-241* (P241T) (Fig. 3C).

Mutations in Pol1p that disrupt the interaction with Cdc13p result in longer telomeres

To determine the importance of the Cdc13p–Pol1p interaction, each of the three *POL1* mutations that reduced or eliminated this interaction by two-hybrid criteria was introduced into the chromosomal copy of *POL1*, using the integration plasmid pCM1 (Lucchini et al. 1988). In these *pol1* haploid isolates, the mutant allele was the only copy of *POL1*, and Pol1p was expressed from its own promoter. Strains carrying the *pol1-236*, 238, or 241 alleles grew as well as wild-type cells at 23°C, 30°C and 37°C (data not shown). Thus, none of the mutant alleles eliminated an essential function of Pol1p.

To determine if telomere length is altered in the *pol1* mutant strains, DNA was prepared from wild-type and mutant cells, digested with *XhoI*, and analyzed by Southern blotting using a telomere probe (Fig. 4A). *XhoI* digestion generates a ~1.3-kb terminal fragment from Y'-bearing telomeres, which make up about two-thirds of the telomeres in yeast, and multiple larger sized bands from X-bearing telomeres. Each of the *pol1* mutants caused an increase in the length of X' and Y' telomeres (Fig. 4A). Cells with the *pol1-236* allele, the allele that essentially eliminated the Pol1Np–Cdc13Np interaction by two-hybrid criteria (Fig. 3C), had telomeres that averaged 150-bp longer than wild type (Fig. 4A, lanes 2,3). The *pol1-238* cells had telomeres that were ~40 bp longer than wild type (Fig. 4A, lanes 5,6) and an ~50% reduction in the Pol1Np–Cdc13Np interaction (Fig. 3C). The *pol1-241* cells had telomeres that were ~90 bp longer than wild type (Fig. 4A, lanes 8,9) and an ~90% reduction in the

Figure 3. Analysis of *pol1* alleles that disrupt interaction of Pol1p with Cdc13p. (A) Positions of functional domains and locations of various mutations within Pol1p are indicated. Mutations in bold disrupted Pol1p interaction with Cdc13p. Positions or regions containing mutations characterized in other labs are also shown: *hpr3* (G439E), *cdc17-2* (G637D), *cdc17-1* (G904D) (Pizzagalli et al. 1988; Lucchini et al. 1990). (B) The wild-type (lane 1) or mutant (lanes 2–7) proteins were expressed from the pJG4-5 vector under control of a *GAL1* promoter. The allele number of the expressed protein is indicated above the lanes. Proteins expressed from pJG4-5 are fused to a transcriptional activation domain and an HA epitope. Cell extracts were analyzed from two different isolates for each mutation using Western blotting with an anti-HA antibody. (C) Extracts were produced from strains expressing both Cdc13Np and either the pJG4-5 vector alone or pJG4-5 with the wild-type *POL1* segment or one of the point mutants, as indicated below. Data are the average of three independent β -galactosidase measurements. Error bars, s.d.



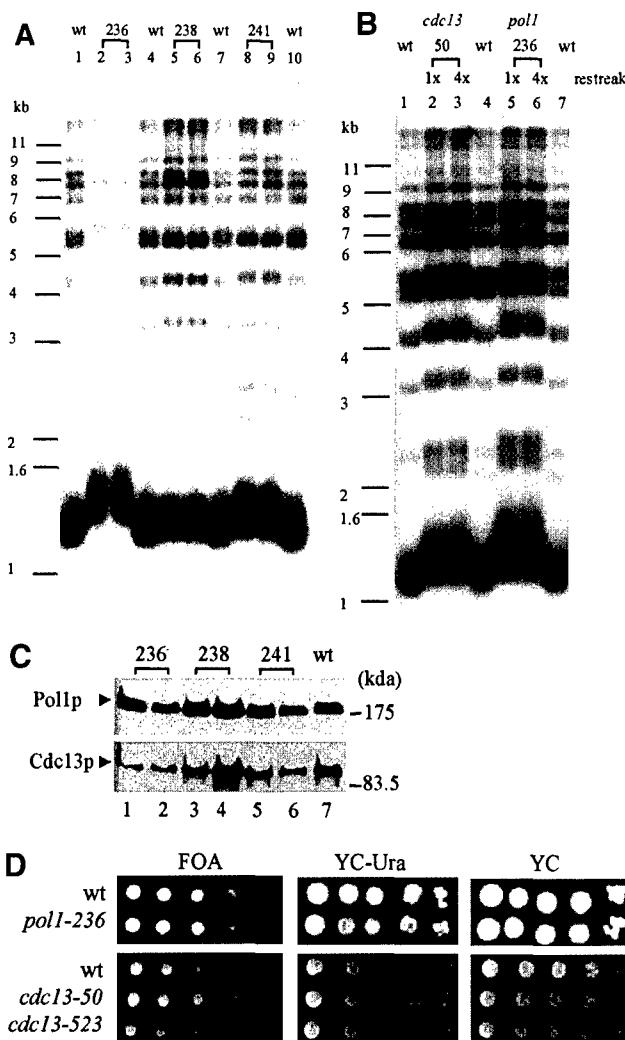


Figure 4. The *pol1* alleles that disrupt Pol1p-Cdc13p interaction cause telomere lengthening. (A) Genomic DNA was isolated from wild type and two independent isolates for each of the three *pol1* alleles (allele numbers are above the lanes), digested with *Xho*I and analyzed by Southern hybridization using a telomeric probe. (Lanes 2, 3) *pol1-236* cells; (lanes 5, 6) *pol1-238* cells; (lanes 8, 9) *pol1-241* cells. (Lanes 1, 4, 7, 10) DNA from an otherwise isogenic wild-type strain. (B) Cells of the indicated genotype were streaked on complete media right after meiosis and allowed to form colonies. DNA was prepared from colonies after the first restreak (~55 divisions postgermination) and after four restreaks (~130 generations). DNA was analyzed as described for panel A. DNA is from wild type (lanes 1, 4, 7), *cdc13-50* cells (lane 2, 55 divisions; lane 3, 155 divisions), and *pol1-236* cells (lane 5, 55 divisions; lane 6, 155 divisions). (C) Western analysis was carried out using DNA from *POL1* or *pol1-236*, -238, and -241 strains expressing 3XMYC-Cdc13p (mutant allele indicated above the lanes; extracts from two independent mutant isolates are shown for each mutant allele). Extracts were analyzed by SDS-PAGE and Western blotting using the anti-Pol1p serum (top) or anti-myc serum (lower panel). The minor differences in protein levels were not reproducible. (D) To measure telomere position effect, ten-fold serial dilutions of otherwise isogenic strains that were either wild type (wt) or contained the indicated mutations and having *URA3* next to the left telomere of chromosome VII were spotted onto plates containing complete medium plus FOA, complete medium lacking uracil (YC-uracil), or complete medium (YC).

Pol1p-Cdc13p interaction (Fig. 3C). The increase in telomere length seen in these mutants was not progressive, as telomeres were equally long at ~55 and ~130 divisions after sporulation (Fig. 4B). This telomere lengthening was telomerase-mediated as lengthening was not seen in a *tlc1 pol1-236* double mutant strain (data not shown). As Western analysis showed that each of the mutant alleles produced essentially wild-type levels of both Pol1p and Cdc13p (Fig. 4C), the differences in telomere lengths are unlikely to be attributable differences in the abundance of either protein.

Telomere position effect, (TPE), refers to the fact that genes placed near a yeast telomere are transcriptionally repressed (Gottschling et al. 1990). Certain *POL1* alleles that affect the catalytic activity of polymerase α cause telomere lengthening, loss of TPE, and altered G-tail metabolism (Adams-Martin et al. 2000). To monitor TPE, the *URA3* gene was placed next to the left telomere of chromosome VII in the *pol1-236* strain and the fraction of cells able to grow on FOA medium determined. As the *pol1-236* strain had the same fraction of FOA^R cells as an isogenic wild-type control (Fig. 4D), TPE was not affected by the *pol1-236* mutation. Long TG₁₋₃ tails are

detected on yeast telomeres at the end of S phase (Wellinger et al. 1993b) but not in DNA from log phase cultures except in strains that are defective in their processing, as occurs in *cdc13-1* cells growing at high temperatures (Gravel et al. 1998) or *hdf1* strains that lack the Ku heterodimeric complex (Gravel et al. 1998; Polotnianka et al. 1998). Using the nondenaturing hybridization method described in Dionne and Wellinger (1996), G-tails were not detected at *pol1-236* telomeres whereas G-tails were readily detected on *cdc13-1* and *hdf1*Δ telomeres (data not shown). Thus, by the limits of this assay, a mutation that disrupted Cdc13p-Pol1p interaction did not affect G-tail structure.

The *pol1-236* mutation eliminated interaction of Pol1p with Cdc13p, as monitored by the two-hybrid assay (Fig. 3C). To determine if this mutation eliminated the interaction of full-length Pol1p and full-length Cdc13p, we carried out the same experiment used to detect the interaction of these proteins in wild-type cells (Fig. 2B). A protein extract was prepared from *pol1-236* cells expressing a MYC₃-tagged Cdc13p. The extract was immunoprecipitated with anti-MYC antibodies and analyzed by Western blotting using both anti-MYC and anti-

Pol1p antibodies. The amount of Pol1-236p in the immunoprecipitate was reduced substantially but not eliminated (Fig. 2B, IP panel, lane 2). The residual interaction between Cdc13p and Pol1-236p suggests that the Pol1p-Cdc13p interaction is mediated by more than one region in Pol1p and/or that other subunits of polymerase α contribute to its interaction with Cdc13p.

Mutations in CDC13 that disrupt the Cdc13Np-Pol1Np interaction cause long telomeres

To identify mutations in *CDC13* that disrupt the Cdc13Np-Pol1Np interaction, the 600 amino acid fragment of Cdc13p used in the initial two-hybrid screen (Fig. 1) was mutagenized by PCR. The mutagenized segments were reintroduced into the yeast two hybrid strain, and mutants that failed to interact with Pol1Np were identified by their generating white colonies in the lacZ filter assay. Of the 14 candidate mutations that pro-

duced full-length protein, DNA sequencing revealed that 12 contained a single amino acid mutation, and two contained multiple point mutations. The single mutations were spread throughout the 600 amino acids of Cdc13Np (Fig. 5A). As with the *pol1* mutations, these alleles were named according to the mutated amino acid residue. Eight of the 12 mutant polypeptides were expressed at wild-type levels, and six of these still interacted with Fun12Mp in the two hybrid assay, one of three polypeptides identified as interacting with wild-type Cdc13Np (summarized in Fig. 5B). Mutants *cdc13-50* (K50Q), *cdc13-124* (C124R), *cdc13-129* (L129S), *cdc13-228* (S228P), *cdc13-392* (L392P) and *cdc13-523* (I523V) were defective specifically in their ability to interact with Pol1Np, and these defects were not due to reduced protein expression.

To determine the phenotypes of *cdc13* alleles that reduced the Cdc13Np-Pol1Np interaction, we introduced two of the mutations, *cdc13-50* and *cdc13-523*, on a cen-

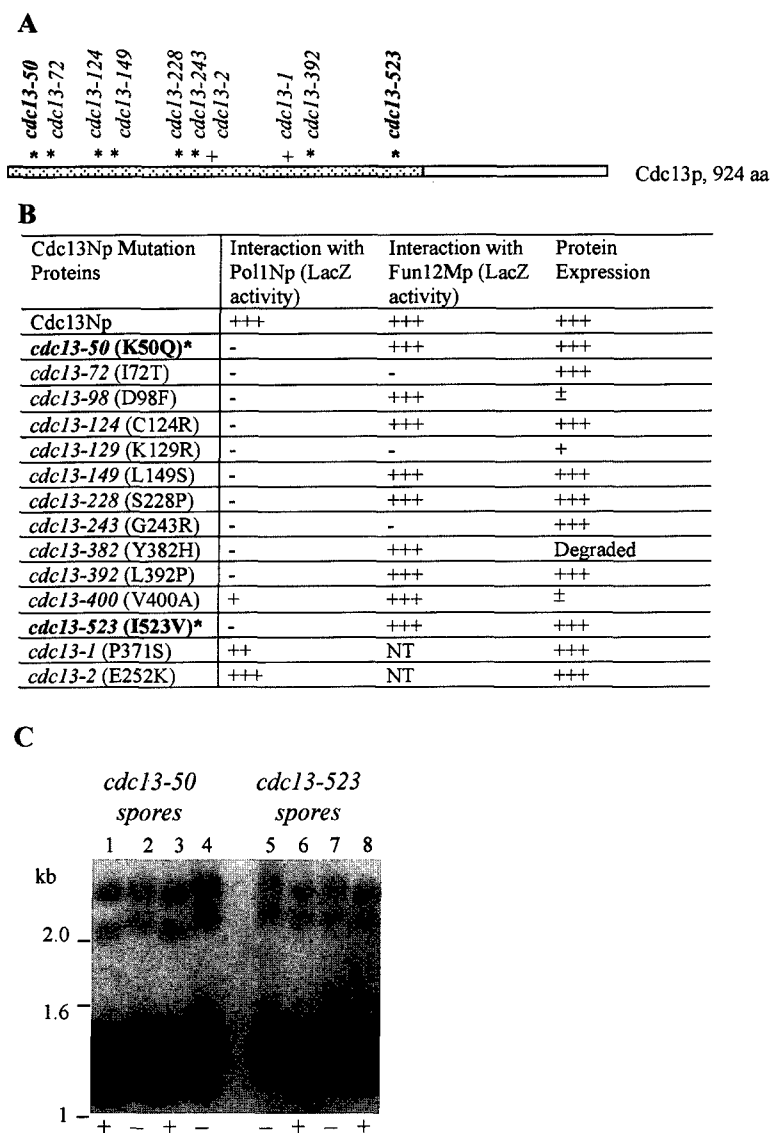


Figure 5. Alleles of *CDC13* that disrupt interactions with *POL1*. (A) The portion of Cdc13p that was used as bait in the two-hybrid screen is represented by the dotted region. Asterisks mark the sites of single amino acid substitutions that reduced the interaction of Cdc13Np with Pol1Np by two hybrid criteria. The alleles in bold are those used for phenotypic analyses. The locations of the *cdc13-1* and *cdc13-2* alleles are also indicated (+) (Lin and Zakian 1996; Nugent et al. 1996). (B) Alleles for *cdc13* polypeptides that lost interaction with Pol1Np are named by the number of the mutated residue. The amino acid change in each allele is noted in parentheses. Each mutant allele was checked for its ability to interact with both Pol1Np and Fun12Mp in the two hybrid assay using the LacZ filter assay. Western blotting was used to determine if strains carrying the mutant allele made wild-type levels of Cdc13p. (NT) Not tested (+++) wild-type levels of interaction or protein expression, (-) no interaction or protein expression; (±, +, and ++) intermediate levels of interaction or protein expression. (C) *Xho*I digested DNA from wild-type strain or *cdc13Δ* strains carrying a centromere plasmid with mutant *cdc13* alleles (*cdc13-50* and *cdc13-523*) that disrupted interaction of Cdc13Np with Pol1Np was analyzed by Southern blotting. Only the lower portion of the gels is shown. (+) Wild type (-) mutant *cdc13* alleles.

tromere plasmid into a *cdc13Δ* strain such that the mutant allele was the only copy of *CDC13* in the strain. These alleles were chosen because they produced stable protein and had little or no interaction with Pol1Np by two hybrid criteria, yet continued to interact with Fun12Mp (Fig. 5B). The growth of *cdc13-50* and *cdc13-523* cells was comparable to that of wild-type cells at 23°C, 30°C, and 37°C (data not shown), but *cdc13-50* (Fig. 5C, lanes 2,4) and *cdc13-523* (Fig. 5C, lanes 5,7) cells had telomeres that averaged 50 bp longer than wild type (Fig. 5C, lanes 1,3,6,8). As with the *pol1* alleles, this lengthening was not progressive (Fig. 4B, *cdc13-50*; cf. lanes 2 and 3). Cells carrying the *cdc13-50* and *cdc13-523* alleles had wild-type levels of TPE (Fig. 4D).

Cdc13Np interacts with Est1Cp by two-hybrid, genetic, and biochemical criteria

Telomere-bound Cdc13p has been proposed to function by recruiting telomerase to the telomere (Evans and Lundblad 1999; Zhou et al. 2000). As both Est1p (Lin and Zakian 1995; Steiner et al. 1996) and Est2p (Counter et al. 1997; Lingner et al. 1997a,b) are associated with *TLC1* telomerase RNA, we tested the ability of Cdc13Np to interact with both Est1p and Est2p in the two-hybrid assay. Full-length Est2p or the carboxy-terminal region of Est1p (Est1Cp) were cloned into the prey vector (Fig. 1). These plasmids were introduced into the two hybrid reporter strain expressing Cdc13Np. Cells expressing the Est2p fusion protein did not interact with Cdc13Np in the two-hybrid assay (Fig. 6A), although Western analysis showed that the Est2p fusion protein was expressed (data not shown). In contrast, Est1Cp, which contained the carboxy-terminal 200 amino acids of the 699 amino acid Est1p, did interact with Cdc13Np as demonstrated by the growth of cells expressing Est1Cp and Cdc13Np on plates lacking leucine (Fig. 6A,B). However, this association was weaker than the Cdc13Np-Pol1Np interaction, as expressing Cdc13Np and Est1Cp did not activate the LacZ reporter gene (data not shown). By the two hybrid assay, Est1Cp also interacted with mutant proteins Cdc13-1Np and Cdc13-2Np (Fig. 6B).

Using the biochemical approach that detected interaction of Cdc13p and Pol1p (Fig. 2B), we were unable to detect association of endogenous Cdc13p and Est1p (data not shown), perhaps because both proteins are rare and/or because their interaction within cells occurs only in a transient manner. To demonstrate the interaction biochemically, we prepared extracts from cells that overexpressed both proteins. GST-Est1p was expressed from the centromere plasmid pKT/*EST1* (Mitchell et al. 1993) using the inducible *GAL1* promoter, and HA-tagged Cdc13p was expressed from the *PGK1* promoter on the 2 μm-based plasmid pTHA/*CDC13* (Lin and Zakian 1996). Although the GST-Est1p lacked the first 31 amino acids of Est1p, it complemented an *est1* strain (Y. Yamashita, unpubl.). Extracts were incubated with glutathione sepharose (GS) beads. After extensive washing, the material bound to the beads was eluted and analyzed by Western blotting for the presence of GST-Est1p and HA-Cdc13p

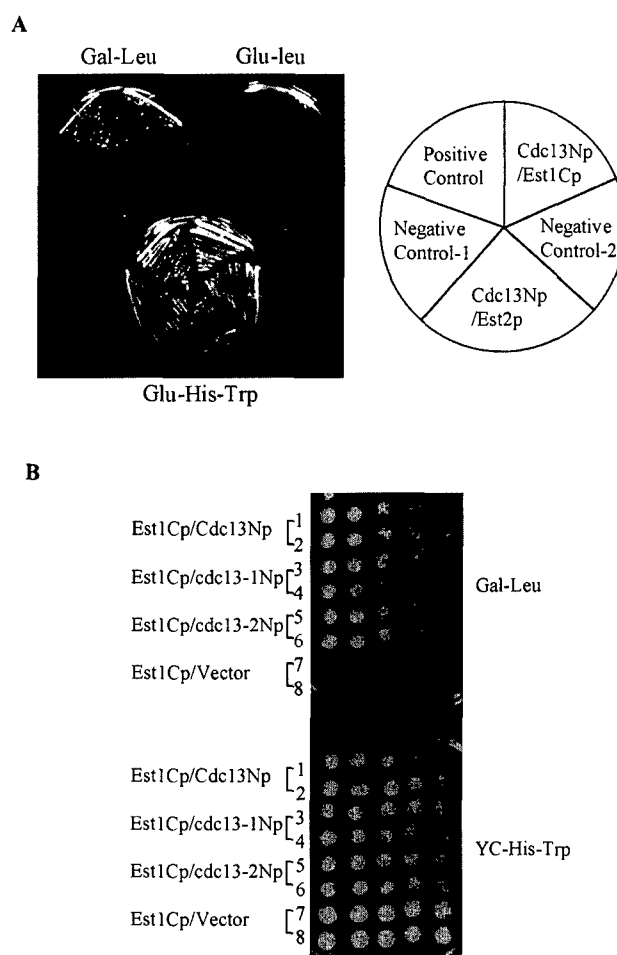
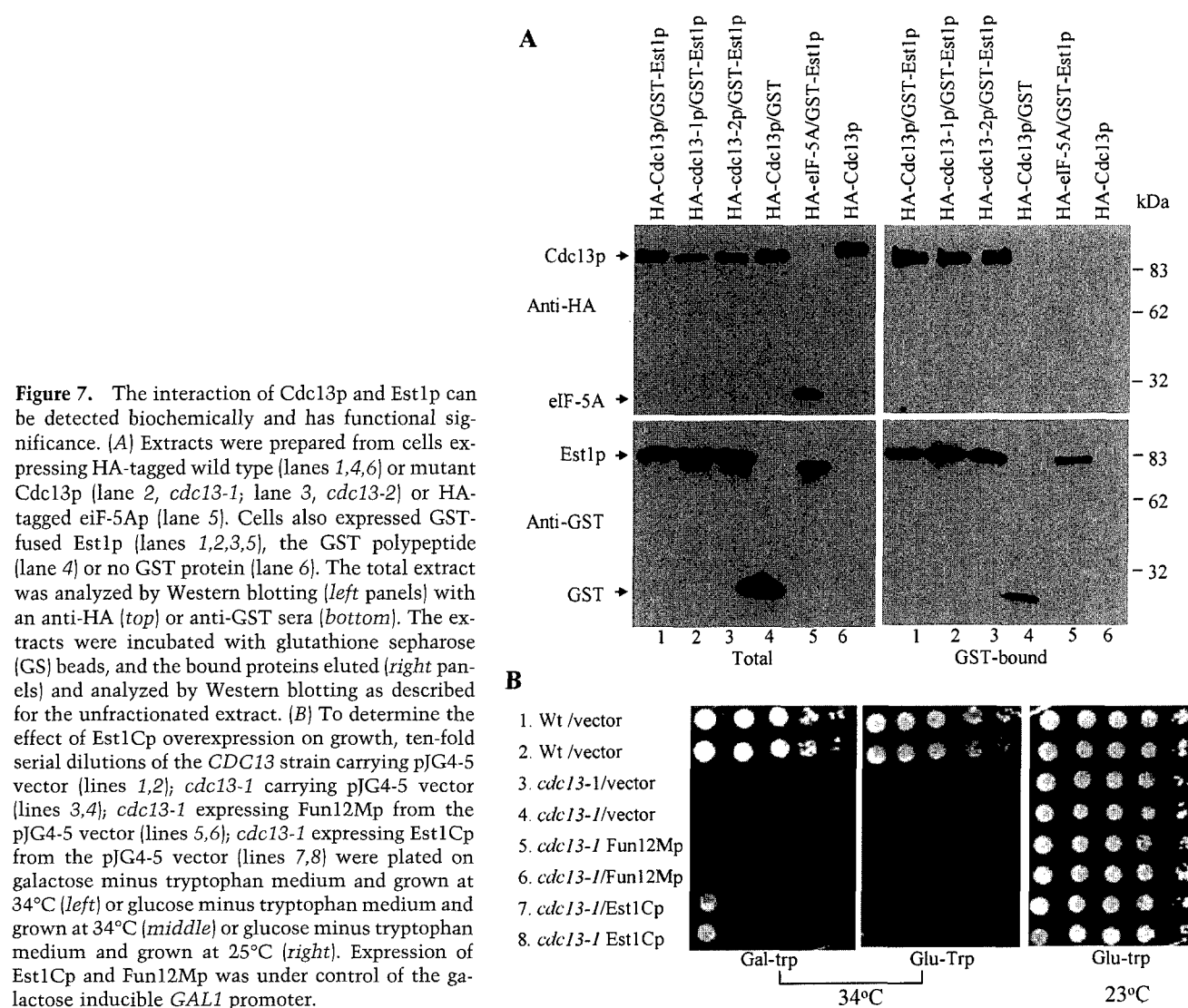


Figure 6. Cdc13Np interacts with Est1Cp in a two-hybrid assay. (A) Two hybrid analysis demonstrates that Cdc13Np interacted with Est1Cp but not with full-length Est2p. Positive and negative controls are the same as in Figure 2A. (B) Ten-fold serial dilutions of each strain were plated onto media selecting for the two-hybrid interaction (Gal - Leu) or selecting for the two plasmids (YC-His-Trp). The terminal 600 amino acids from *CDC13*, *cdc13-1*, or *cdc13-2* were tested for their ability to interact with Est1Cp.

(Fig. 7A). HA-Cdc13p was bound to the beads in extracts from cells expressing GST-Est1p (Fig. 7A, upper right panel, lane 1) but not from cells expressing just the GST protein (Fig. 7, upper right panel, lane 4) or no GST protein at all (Fig. 7, lane 6). The telomere-unrelated protein HA-eIF-5Ap did not interact with GST-Est1p (Fig. 7, lane 5). HA-Cdc13-1p (Fig. 7, lane 2) and HA-Cdc13-2p (Fig. 7, lane 3) also interacted with GST-Est1p in this assay.

To assess if the Cdc13Np-Est1Cp interaction has in vivo significance, we asked if high-level expression of Est1Cp could rescue the temperature-sensitive lethality of *cdc13-1* cells (Fig. 7B). The maximum permissive temperature for cells carrying the *cdc13-1* allele was 28°C although the isogenic wild-type strain grew well at 37°C. Ten-fold serial dilutions of wild-type or *cdc13-1* cells were spotted on galactose (gal) or glucose (glu) plates lacking tryptophan (galactose induces expression of the



fusion proteins; no tryptophan selects for the prey plasmid) and the plates incubated at different temperatures (Fig. 7B). The *cdc13-1* cells carried either prey vector alone, the prey vector expressing Fun12Mp, or the prey vector expressing Est1Cp. All strains grew on glucose plates at 25° (Fig. 7B, right panel); wild-type cells grew on both glucose (Fig. 7B, middle panel) and galactose (Fig. 7B, left panel) plates at 34°. However, *cdc13-1* cells did not grow at 34° unless they expressed Est1Cp: Cells with Est1Cp did not grow at 34° on glucose medium where Est1Cp was not expressed (Fig. 7B, middle panel) nor on galactose plates when expressing Fun12Mp (Fig. 7B, left panel). Thus, high-level expression specifically of Est1Cp suppressed the temperature sensitivity of *cdc13-1* cells, suggesting that the Est1Cp–Cdc13Np interaction was significant. The suppression of the *cdc13-1* growth defect was partial: *cdc13-1* cells expressing Est1Cp did not grow to the same dilution at 34° as the same strain at 25°. Also, the expression of Est1Cp did not allow *cdc13-1* cells to grow at 37° (data not shown).

Discussion

Two-hybrid analysis demonstrated that the amino-terminal two-thirds of Cdc13p interacts with the amino terminus of Pol1p, the catalytic subunit of DNA polymerase α (Fig. 2A). This interaction was confirmed by demonstrating that full-length Cdc13p and Pol1p could be coimmunoprecipitated (Fig. 2B). The interaction could be direct or indirect, mediated by one or more as yet unidentified proteins. Whether direct or not, this interaction is satisfying because polymerase α , by virtue of its primase association, is the only eukaryotic polymerase that can initiate DNA replication (Waga and Stillman 1998) and is hence the best candidate for the polymerase that resynthesizes C-strand DNA. *POL1* mutations that reduced this interaction mapped to a negatively charged, five amino acid patch within the amino-terminal 379 amino acid segment of Pol1p, identified by two-hybrid experiments (Fig. 3A). Although the catalytic subunit of DNA polymerase α is conserved

among eukaryotes, its amino-terminal portion is not conserved and no function had been mapped previously to this region (Fig. 3A).

Mutations in either *POL1* (Fig. 4A) or *CDC13* (Fig. 5C) that reduced the Pol1p-Cdc13p interaction caused telomerase-mediated telomere lengthening. For the *POL1* mutations, this *TLC1*-dependent length increase was greatest in the mutant with the most severe effect on the Pol1p-Cdc13p interaction (Figs. 3C and 4A). These genetic results argue strongly that the Pol1p-Cdc13p interaction detected by both two-hybrid (Fig. 2A) and biochemical (Fig. 2B) criteria is important for telomere maintenance. Others have shown that certain *pol1* alleles show telomerase-mediated telomere lengthening (Carson and Hartwell 1985; Adams and Holm 1996). However, in these cases the mutations are temperature sensitive, affect the catalytic activity of Pol1p, and have global effects on both DNA replication and cell viability. In contrast, the *pol1* mutations studied here occurred in a region that is thought to be dispensable for catalysis, and these mutations had no effect on cell growth or TPE (Fig. 4D). Because these mutations disrupted interaction with Cdc13p (Figs. 2B and 3C), an in vivo telomere binding protein (Bourns et al. 1998), their effects on telomere length were likely direct. Based on these data, we propose that telomere-bound Cdc13p recruits polymerase α to the telomere and that this interaction promotes C-strand resynthesis.

If the Pol1p-Cdc13p interaction is essential for C-strand resynthesis, mutations like *pol1-236* that eliminated this interaction by two-hybrid criteria (Fig. 3C) should be lethal due to loss of telomeric DNA and/or activation of DNA damage checkpoints by long ss G-tails (Lee et al. 1998). However, biochemical experiments demonstrated that there is residual interaction between Cdc13p and Pol1p even in the *pol1-236* strain (Fig. 2B), suggesting that other parts of Pol1p might also interact with Cdc13p. Alternatively or in addition, other telomere proteins (such as Stn1p) whose loss has a similar effect on C-strand degradation as the *cdc13-1* mutation (Grandin et al. 1997), or the Ku heterodimer, whose loss results in G-tails throughout the cell cycle (Gravel et al. 1998; Polotnianka et al. 1998), might be partially redundant with Cdc13p and help recruit Pol1p to the telomere.

We also report that the amino terminus of Cdc13p interacted with the carboxyl third of Est1p by two-hybrid analysis (Fig. 6A). Biochemical experiments confirmed this interaction, demonstrating that full-length Cdc13p interacted in vivo with a close to full-length, functional Est1p (Fig. 7A). Again this interaction could be direct or mediated by another protein. As overexpression of the carboxy-terminal 200 amino acid segment of Est1p partially suppressed the temperature sensitive lethality of *cdc13-1* cells whereas overexpression of another Cdc13p interacting polypeptide, Fun12Mp, did not (Fig. 7B), the Cdc13p-Est1p interaction is likely to be important for the essential function of Cdc13p.

Because Est1p is associated with telomerase RNA (Lin and Zakian 1995; Steiner et al. 1996), a Cdc13p-Est1p

interaction could recruit telomerase to telomeres, thereby explaining the senescence phenotype of *cdc13-2* cells. That Cdc13p functions to recruit telomerase is also supported strongly by the demonstration that expression of a Cdc13p-Est1p or a Cdc13p-Est2p fusion protein supplants the need for Est1p in telomere maintenance (Evans and Lundblad 1999). However, as these fusion proteins might bypass the normal role of Est1p, these data do not address whether Cdc13p and Est1p interact in vivo, whereas our data provide strong support for this possibility.

If Cdc13p recruits telomerase by virtue of its ability to interact with Est1p, the phenotype of *cdc13-2* cells could be explained if Cdc13-2p were unable to interact with Est1p. However, Est1p interacted with Cdc13-2p as assayed by both two-hybrid (Fig. 6B) and biochemical (Fig. 7A) criteria. One explanation for this result is trivial: Est1p might be able to interact with Cdc13-2p when both proteins are overproduced as they were in both the two-hybrid and biochemical assays, but not when the proteins are expressed at physiological levels. This possibility predicts that overexpression of Est1p would suppress the telomere defects of a *cdc13-2* strain. However, overexpression of a GST-EST1p fusion protein that contained all but the most amino-terminal 31 amino acids of Est1p, and which complemented an *est1* strain or overexpression of the carboxy-terminal portion of Est1p, did not suppress the telomere length defect of *cdc13-2* cells (data not shown). Alternatively, interaction of Cdc13p with Est1p might trigger a conformational change in Cdc13p, such as phosphorylation, that makes it better able to promote the telomerase pathway: Cdc13-2p might be able to interact with Est1p but not undergo the subsequent conformational change.

In the ciliate *Euplotes*, aphidicolin, an inhibitor of conventional DNA polymerases, causes changes in the length of both the C- and G-strands of telomeric DNA. This result led to the proposal that G-strand lengthening and C-strand synthesis are coordinately regulated (Fan and Price 1997). In *Saccharomyces*, addition of duplex telomeric DNA to a double-strand break requires not only telomerase but also DNA primase and two of the three conventional DNA polymerases, polymerase α and polymerase β (Diede and Gottschling 1999). These data also support a model of coordinate regulation of C- and G-strand synthesis. Cdc13p, by virtue of its ability to interact with both the conventional replication apparatus (Fig. 2A) and a telomerase component (Fig. 6A), could play a critical role in this coordination.

Alternatively, there might be competition between C-strand resynthesis by polymerase α and G-strand lengthening by telomerase. In support of this model, when the interaction between Cdc13p and Pol1p was weakened, telomerase-mediated lengthening of telomeres increased (Fig. 4A). By two hybrid criteria, Cdc13p interacted more strongly with Pol1p than it did with Est1p. Also, polymerase α is much more abundant than telomerase. Based on these considerations, a competition model predicts that recruitment by Cdc13p of polymerase α to ss TG₁₋₃

tails is much more likely than recruitment of telomerase. As short telomeres are preferentially lengthened by telomerase [Marcand et al. 1999], telomere length may influence this competition.

Materials and methods

The yeast two-hybrid strain YEM1 α (MATa *his3-11 trp1-1 LEU2::pLexAop-LEU2 ura3-1::URA3-pLexAop-Gal1-lacZ*) was constructed by E. Monson in our lab using reagents described in Gyuris et al. (1993). The protease-deficient yeast strain BJ2168 [Jones 1991] was used to detect Est1p-Cdc13p interaction. YPH499, YPH500, and YPH501 [Sikorski and Hieter 1989] were used for analysis of *pol1* and *cdc13* mutant phenotypes. *CDC13* deletion in YPH501 was created by J.J. Lin [Lin and Zakian 1996]. Cdc13p was tagged at its carboxyl terminus with three MYC epitopes [Schneider et al. 1995] in strain VPS106 [V. Schulz and V. Zakian 1994] and used for immunoprecipitation. *CDC13* was also tagged with nine MYC epitopes at its carboxyl terminus in strain YPH499 [A. Taggart and V.A. Zakian, unpubl.] as in Zachariae et al. [1996]. Cells containing either 3-myc-tagged or 9-myc-tagged *CDC13* had normal cell growth and telomere length.

The two-hybrid assay was done essentially as described [Gyuris et al. 1993] using components generously provided by Dr. R. Brent and colleagues. The prey library [pJG4-5/X] [Gyuris et al. 1993; Golemis et al. 1994] was transformed into YEM1 α strain containing the *CDC13N* bait plasmid [pEG202/*CDC13-B*, constructed by J.-J. Lin] (Fig. 1). HA₃-tagged *CDC13* from pTHA/*CDC13* [Lin and Zakian 1996] was inserted into *Bam*HI-*Sal*I digested pVZ1, then *Bgl*II digested, followed by self-ligation to delete the *Bgl*II fragment within *CDC13*. This introduces a frameshift at the ligation site such that only the first 600 amino acids of Cdc13p are produced. The HA-tagged amino-terminal region of *CDC13* was cloned into pEG202 generating a LexA DNA binding domain-Cdc13Np fusion protein expressed from the *ADH1* promoter. The library proteins were also HA tagged and expressed from a *GAL1* promoter. Transformants were harvested and replated to 3% galactose YC plates minus leucine. Leu⁺ colonies were transferred to nitrocellulose filters [Schleicher & Schuell] for the colony *lacZ* filter assay [Breedon and Nasmith 1985]. Liquid *lacZ* assays were done as in Miller [1972] and Guarente [1983]. Leu⁺ LacZ⁺ colonies were purified and confirmed by rescuing the plasmids into *E. coli* and retransforming them into the two-hybrid yeast strain. Prey plasmids that retested positively were sequenced with the amplitaq FS dye terminator cycle sequencing kit [ABI].

The plasmid pJG4-5/*EST1C* was constructed by Y. Yamashita and contained amino acids 499–699 of Est1p fused in-frame with the B42 trans-activation domain and an HA epitope tag in the prey vector pJG4-5. Full-length *EST2* was PCR amplified from YPH499 genomic DNA and cloned into pJG4-5 to create pJG4-5/*EST2* (made by S.-C. Teng). This plasmid complements an *est2 Δ* strain. The *cdc13-1* mutation P371S creates an additional *Eco*RI site [Lin and Zakian 1996] and *cdc13-2* mutation E252K eliminates the first *Eco*RI site in *CDC13* [Nugent et al. 1996]. pEG202/*cdc13-1B* and pTHA/*cdc13-1* were created by site-directed mutagenesis [QuikChange Site-Directed Mutagenesis Kit, Stratagene] from pEG202/*CDC13N* and pTHA/*CDC13*, respectively. pEG202/*cdc13-2B* and pTHA/*cdc13-2* were created from pEG202/*CDC13N* and pTHA/*CDC13*, respectively. The mutated plasmids were introduced into *E. coli*, purified, and then sequenced to confirm that the correct mutation was made. Plasmids producing LexA::cRafp or LexA::Krev1p fusion protein were gifts from E. Golemis.

Immunoprecipitation to detect Cdc13p-Pol1p interaction was done in yeast strain VPS106 containing 3XMYC-*CDC13* as in Garrett et al. [1991], except that cells were lysed in a homogenizer (Avestin's High Pressure Homogenizer EmulsiFlexC-5) in 40 mM Tris (pH 8.0), 75 mM NaCl, and 20 mM KOAc, 1 mM 2-mercaptoethanol, 0.01% NP-40, 1 mM phenylmethylsulfonyl fluoride (PMSF), 10 μ g/ml pepstatin A, 5 μ g/ml leupeptin, 10 μ g/ml aprotinin, 10 mM benzamidine, 250 μ g/ml soybean trypsin inhibitor, and 10% glycerol. Monoclonal anti-MYC 9E10 [Princeton University] was used for the immunoprecipitation. Monoclonal anti-Pol1p antibody (from P. Plevani), anti-MYC antibody, and ECL (Amersham Life Science) were used to visualize the proteins.

The GST (glutathione S transferase) assay was conducted essentially as described in Mitchell et al. [1993]. pEGKT/*EST1*, a plasmid that produces a GST::Est1p fusion protein under the control of a *GAL1* promoter, was constructed and generously provided by Y. Yamashita. In brief, the *Hinc*II fragment from *EST1*, which contains amino acids 31–699 of Est1p, was cloned into pEGKT [Mitchell et al. 1993]. Expression of GST::Est1p from this plasmid complements an *est1 Δ* mutant (data not shown). Full-length Cdc13p tagged with 3 $\times$ HA at its amino terminus was expressed from a *PGK1* promoter on plasmid pTHA/*CDC13* [Lin and Zakian 1996]. The two proteins were coexpressed in strain BJ2168. A similarly HA-tagged version of the yeast protein eIF-5A [Kang and Hershey 1994] was constructed by J.-J. Lin and used as a negative control. Raffinose-grown log phase cells were transferred to 3% galactose medium and incubated at room temperature overnight to induce expression of *GAL1*-regulated genes. Cells were lysed in 20 mM Tris (pH 8.0), 200 mM NaCl, 1 mM EDTA, 1 mM DTT, 0.01% NP-40, 10% glycerol with protease inhibitors plus an equal volume of glass beads by vortexing in mini-beadbeater-8 [Biospec Products]. NP-40 and Triton X-100 were added to the soluble protein fractions to final concentrations of 0.5% and 0.1%, respectively. GST-Sepharose 4B beads (Pharmacia Biotech) were then added. After incubation at 4°C, beads were collected by centrifugation and then washed once with the lysis buffer containing 1% NP-40 0.1% Triton X-100 and once with lysis buffer containing 450 mM NaCl. The beads were treated with 0.1 mg/ml DNase I in lysis buffer plus 1 mM MgCl₂, then washed with lysis buffer containing 450 mM NaCl and 350 mM KOAc. Proteins were visualized by ECL Western using anti-HA antibody 12CA5 [Princeton University] and rabbit anti-GST antibodies [from Jin-Qiu Zhou].

PCR mutagenesis was used to generate *POL1* and *CDC13* mutations [Grandin et al. 1997]. Two DNA oligonucleotides that were complementary to the sequences from the prey vector pJG4-5 (for *POL1* mutations) or to the bait vector pEG202 (for *CDC13*) were used. A modified random PCR mutagenesis [Leung et al. 1989; Zhou et al. 1991] was done in a reaction mixture containing 100 μ M dNTP, 200 μ M oligonucleotides, 1 $\times$ PCR buffer with 3 mM MgCl₂, 1–10 ng/100 μ l plasmid DNA template, and 1 unit Taq polymerase [Roche Molecular Biochemicals]. PCR products were cotransformed into the two-hybrid strain containing bait or prey plasmids with *Xho*I linearized pJG4-5 or pEG202 vector. Pools of the PCR mutagenized *POL1* segments were introduced by transformation into the yeast strain containing both the two-hybrid reporter genes and expressing Cdc13Np [Grandin et al. 1997]. Transformants were selected by their ability to grow on media lacking histidine and tryptophan, thus selecting for both the prey and bait plasmids but not for the two-hybrid interaction. Transformants were replica plated to galactose plates then transferred to nitrocellulose membranes for LacZ assays.

The *pol1-236*, *pol1-238*, and *pol1-241* mutant alleles were in-

roduced into the *POL1* integration plasmid pCM1 (Lucchini et al. 1988) using the QuickChange site-directed Mutagenesis Kit (Stratagene). Silent mutations that created an *AccI* site in *pol1-236* (T₇₀₈→C and G₇₁₀→T in addition to the D236N mutation, G₇₀₆→A) or an additional *EcoRI* site in *pol1-238* and *241* (A₇₁₃→G in addition to the E238K mutation, G₇₁₁→A or P241T, C₇₂₀→A) were also introduced to allow identification of mutant alleles by restriction enzyme digestion. The mutagenized plasmids were linearized by *Bst*XI digestion and integrated into the *POL1* locus of diploid strain YPH501 as in Lucchini et al. (1988). Transformants were selected by their Ura⁺ phenotype and verified by Southern. The diploid cells were sporulated; spores containing the mutant *POL1* alleles were identified by Ura⁺ phenotype. Cells that excised *URA3*, leaving behind an intact but mutated *POL1* gene, were selected on plates containing 5-fluoro orotic acid.

Wild-type *CDC13* with its own promoter and terminator sequences was released from YEP24/*CDC13* using *ApaI* digestion (Garvik et al. 1995) and cloned into the *ApaI*-digested *CEN* plasmid pRS314. The pRS314/*CDC13* was mutagenized using the QuickChange mutagenesis kit. To facilitate identification of mutations, an *AvrII* site was created near K50Q in pRS/*cdc13-50* and the first *HindIII* site of *CDC13* was eliminated at I523V in pRS314/*cdc13-523* without changing additional amino acid residues. The mutated plasmids were introduced into a derivative of YPH 501 that had one copy of *CDC13* and one copy of *cdc13::HIS3* (Lin and Zakian 1996). After sporulation and dissection, haploid *cdc13::HIS3* spores carrying the mutated *cdc13* plasmid were selected by their His⁺ Trp⁺ phenotype.

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References

- Adams, A.K. and Holm, C. 1996. Specific DNA replication mutations affect telomere length in *Saccharomyces cerevisiae*. *Mol. Cell. Biol.* **16**: 4614-4620.
- Adams-Martin, A., Dionne, I., Wellinger, R.J., and Holm, C. 2000. The function of DNA polymerase alpha at telomeric G tails is important for telomere homeostasis. *Mol. Cell. Biol.* **20**: 786-796.
- Bourns, B.D., Alexander, M.K., Smith, A.M., and Zakian, V.A. 1998. Sir proteins, Rif proteins and Cdc13p bind *Saccharomyces cerevisiae* telomeres in vivo. *Mol. Cell. Biol.* **18**: 5600-5608.
- Breedon, L. and Nasmyth, K. 1985. Regulation of the yeast HO gene. *Cold Spring Harbor Symp. Quant. Biol.* **50**: 643-650.
- Carson, M.J. and Hartwell, L. 1985. CDC 17: An essential gene that prevents telomere elongation in yeast. *Cell* **42**: 249-257.
- Choi, S.K., Lee, J.H., Zoll, W.L., Merrick, W.C., and Dever, T.E. 1998. Promotion of met-tRNA^{Met} binding to ribosomes by yIF2, a bacterial IF2 homolog in yeast. *Science* **280**: 1757-1760.
- Conrad, M.N., Wright, J.H., Wolf, A.J., and Zakian, V.A. 1990. RAP1 protein interacts with yeast telomeres in vivo: Overproduction alters telomere structure and decreases chromosome stability. *Cell* **63**: 739-750.
- Counter, C.M., Meyerson, M., Eaton, E.N., and Weinberg, R.A. 1997. The catalytic subunit of yeast telomerase. *Proc. Natl. Acad. Sci.* **94**: 9202-9207.
- Diede, S.J. and Gottschling, D.E. 1999. Telomerase-mediated telomere addition in vivo requires DNA primase and DNA polymerases alpha and delta. *Cell* **99**: 723-733.
- Dionne, I. and Wellinger, R.J. 1996. Cell cycle-regulated generation of single-stranded G-rich DNA in the absence of telomerase. *Proc. Natl. Acad. Sci.* **93**: 13902-13907.
- Evans, S.K. and Lundblad, V. 1999. Est1 and Cdc13 as comediators of telomerase access. *Science* **286**: 117-120.
- Fan, X. and Price, C.M. 1997. Coordinate regulation of G- and C-strand length during new telomere synthesis. *Mol. Biol. Cell* **8**: 2145-2155.
- Fields, S. and Song, O.-K. 1989. A novel genetic system to detect protein-protein interactions. *Nature* **340**: 245-246.
- Garrett, S., Menold, M.M., and Broach, J.R. 1991. The *Saccharomyces cerevisiae* YAK1 gene encodes a protein kinase that is induced by arrest early in the cell cycle. *Mol. Cell. Biol.* **11**: 4045-4052.
- Garvik, B., Carson, M., and Hartwell, L. 1995. Single-stranded DNA arising at telomeres in *cdc13* mutants may constitute a specific signal for the RAD9 checkpoint. *Mol. Cell. Biol.* **15**: 6128-6138.
- Golemis, E.A., Gyuris, J., and Brent, R. 1994. Interaction trap/two-hybrid system to identify interacting proteins. In *Current Protocols in Molecular Biology* (ed. F.M. Ausubel, R. Brent, R. Kingston, D. Moore, J. Seidman, J.A. Smith, and K. Struhl), pp. 20.0.1-20.1.28. John Wiley & Sons, New York.
- Gottschling, D.E., Aparicio, O.M., Billington, B.L., and Zakian, V.A. 1990. Position effect at *S. cerevisiae* telomeres: Reversible repression of Pol II transcription. *Cell* **63**: 751-762.
- Grandin, N., Reed, S.I., and Charbonneau, M. 1997. Stn1, a new *Saccharomyces cerevisiae* protein, is implicated in telomere size regulation in association with Cdc13. *Genes & Dev.* **11**: 512-527.
- Gravel, S., Larrivee, M., Labrecque, P., and Wellinger, R.J. 1998. Yeast Ku as a regulator of chromosomal DNA end structure. *Science* **280**: 741-744.
- Guarente, L. 1983. Yeast promoters and lacZ fusions designed to study expression of cloned genes in yeast. *Methods Enzymol.* **101C**: 181-191.
- Gyuris, J., Golemis, E., Chertkov, H., and Brent, R. 1993. Cdi1, a human G1 and S phase protein phosphatase that associates with Cdk2. *Cell* **75**: 791-803.
- Jones, E. 1991. Tackling the Protease Problem in *Saccharomyces cerevisiae*. In *Guide to Yeast Genetics and Molecular Biology* (ed. C. Guthrie and G. Fink), pp. 933. Academic Press, San Diego, California.
- Kang, H.A. and Hershey, J.W. 1994. Effect of initiation factor eIF-5A depletion on protein synthesis and proliferation of *Saccharomyces cerevisiae*. *J. Biol. Chem.* **269**: 3934-3940.
- Khazak, V., Sadhale, P.P., Woychik, N.A., Brent, R., and Golemis, E.A. 1995. Human RNA polymerase II subunit hSRP7 functions in yeast and influences stress survival and cell morphology. *Mol. Biol. Cell* **6**: 759-775.
- Lee, S.E., Moore, J.K., Holmes, A., Umez, K., Kolodner, R.D., and Haber, J.E. 1998. *Saccharomyces* Ku70, mre11/rad50 and RPA proteins regulate adaptation to G2/M arrest after DNA damage. *Cell* **94**: 399-409.

- Leung, D., Chen, E., and Goeddel, D. 1989. A method for random mutagenesis of a defined DNA segment using a modified polymerase chain reaction. *Technique* **1**: 11-15.
- Lin, J.-J. and Zakian, V.A. 1995. An *in vitro* assay for *Saccharomyces* telomerase requires *EST1*. *Cell* **81**: 1127-1135.
- . 1996. The *Saccharomyces CDC13* protein is a single-strand TG₁₋₃ telomeric DNA binding protein *in vitro* that affects telomere behavior *in vivo*. *Proc. Natl. Acad. Sci.* **93**: 13760-13765.
- Lingner, J., Cech, T.R., Hughes, T.R., and Lundblad, V. 1997a. Three Ever Shorter Telomere (EST) genes are dispensable for *in vitro* yeast telomerase activity. *Proc. Natl. Acad. Sci.* **94**: 11190-11195.
- Lingner, J., Hughes, T.R., Shevchenko, A., Mann, M., Lundblad, V., and Cech, T.R. 1997b. Reverse transcriptase motifs in the catalytic subunit of telomerase. *Science* **276**: 561-567.
- Lucchini, G., Mazza, C., Scacheri, E., and Plevani, P. 1988. Genetic mapping of the *Saccharomyces cerevisiae* DNA polymerase I gene and characterization of a pol¹ temperature-sensitive mutant altered in DNA primase-polymerase complex stability. *Mol. Gen. Genet.* **212**: 459-465.
- Lucchini, G., Falconi, M.M., Pizzagalli, A., Aguilera, A., Klein, H.L., and Plevani, P. 1990. Nucleotide sequence and characterization of temperature-sensitive pol¹ mutants of *Saccharomyces cerevisiae*. *Gene* **90**: 99-104.
- Marcand, S., Brevet, V., and Gilson, E. 1999. Progressive cis-inhibition of telomerase upon telomere elongation. *EMBO J.* **18**: 3509-3519.
- Miller, J.H. 1972. Replica plating. In *Experiments in Molecular Genetics* (ed. J.H. Miller), pp. 56-59. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY.
- Mitchell, D.A., Marshall, T.K., and Deschenes, R.J. 1993. Vectors for the inducible overexpression of glutathione S-transferase fusion proteins in yeast. *Yeast* **9**: 715-723.
- Nugent, C.I. and Lundblad, V. 1998. The telomerase reverse transcriptase: Components and regulation. *Genes & Dev.* **12**: 1073-1085.
- Nugent, C.I., Hughes, T.R., Lue, N.F., and Lundblad, V. 1996. Cdc13p: A single-strand telomeric DNA-binding protein with a dual role in yeast telomere maintenance. *Science* **274**: 249-252.
- Pizzagalli, A., Valsasnini, P., Plevani, P., and Lucchini, G. 1988. DNA polymerase I gene of *Saccharomyces cerevisiae*: Nucleotide sequence, mapping of a temperature-sensitive mutation, and protein homology with other DNA polymerases. *Proc. Natl. Acad. Sci.* **85**: 3772-3776.
- Polotnianska, R.M., Li, J., and Lustig, A.J. 1998. The yeast Ku heterodimer is essential for protection of the telomere against nucleolytic and recombinational activities. *Curr. Biol.* **8**: 831-834.
- Schneider, B.L., Seufert, W., Steiner, B., Yang, Q.H., and Futcher, A.B. 1995. Use of polymerase chain reaction epitope tagging for protein tagging in *Saccharomyces cerevisiae*. *Yeast* **11**: 1265-1274.
- Schulz, V.P. and Zakian, V.A. 1994. The *Saccharomyces PIF1* DNA helicase inhibits telomere elongation and *de novo* telomere formation. *Cell* **76**: 145-155.
- Sikorski, R.S. and Hieter, P. 1989. A system of shuttle vectors and yeast host strains designed for efficient manipulation of DNA in *Saccharomyces cerevisiae*. *Genetics* **122**: 19-27.
- Steiner, B.R., Hidaka, K., and Futcher, B. 1996. Association of the Est1 protein with telomerase activity in yeast. *Proc. Natl. Acad. Sci.* **93**: 2817-2821.
- Virta-Pearlman, V., Morris, D.K., and Lundblad, V. 1996. Est1 has the properties of a single-stranded telomere end-binding protein. *Genes & Dev.* **10**: 3094-3104.
- Waga, S. and Stillman, B. 1998. The DNA replication fork in eukaryotic cells. *Annu. Rev. Biochem.* **67**: 721-751.
- Watt, P.M., Louis, E.J., Borts, R.H., and Hickson, I.D. 1995. Sgs1: A eukaryotic homolog of *E. coli* RecQ that interacts with topoisomerase II *in vivo* and is required for faithful chromosome segregation. *Cell* **81**: 253-60.
- Wellinger, R.J., Wolf, A.J., and Zakian, V.A. 1993a. Origin activation and formation of single-strand TG₁₋₃ tails occur sequentially in late S phase on a yeast linear plasmid. *Mol. Cell. Biol.* **13**: 4057-4065.
- . 1993b. *Saccharomyces* telomeres acquire single-strand TG₁₋₃ tails late in S phase. *Cell* **72**: 51-60.
- Wellinger, R.J., Ethier, K., Labrecque, P., and Zakian, V.A. 1996. Evidence for a new step in telomere maintenance. *Cell* **85**: 423-433.
- Wright, J.H. and Zakian, V.A. 1995. Protein-DNA interactions in soluble telosomes from *Saccharomyces cerevisiae*. *Nucleic Acids Res.* **23**: 1454-1460.
- Wright, J.H., Gottschling, D.E., and Zakian, V.A. 1992. *Saccharomyces* telomeres assume a non-nucleosomal chromatin structure. *Genes & Dev.* **6**: 197-210.
- Zachariae, W., Shin, T.H., Galova, M., Obermaier, B., and Nasmyth, K. 1996. Identification of subunits of the anaphase-promoting complex of *Saccharomyces cerevisiae*. *Science* **274**: 1201-1204.
- Zhou, J., Hidaka, K., and Futcher, B. 2000. The Est1 subunit of yeast telomerase binds the Tlc1 telomerase RNA. *Mol. Cell. Biol.* **20**: 1947-1955.
- Zhou, Y.H., Zhang, X.P., and Eubright, R.H. 1991. Random mutagenesis of gene-sized DNA molecules by use of PCR with Taq DNA polymerase. *Nucleic Acids Res.* **19**: 6052.



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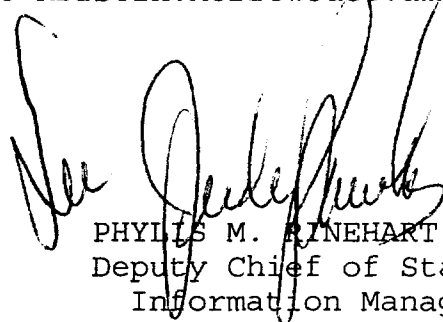
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