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My research proposal focuses on characterizing the roles of proteins that are responsible for mediating access of telomerase to the telomere in S. cerevisiae. I have previously shown that fusion of Cdc13 to Est1 results in substantial telomere elongation. Fusions consisting of mutant versions of Cdc13 or Est1 confer similar telomere elongation, indicating that close physical proximity can bypass telomerase-defective mutations in either protein. Fusing Cdc13 directly to the catalytic core of telomerase (Est2) allows stable telomere maintenance in the absence of Est1. This study led to the proposal that Est1 and Cdc13 are co-mediators of telomerase access, and has further suggested that Est1 may have a second role in telomere replication, based on the observation that the Cdc13-Est2 fusion is unable to promote telomere elongation in the absence of Est1. Since Est1 is a terminus-specific single-strand DNA binding protein, this second role for Est1 may be to promote accessibility of the telomeric end to the telomerase active site. To explore this second role, I have isolated a set of alanine-scan mutations in Est1 which retain association with telomerase but display telomere replication defects, and have tentatively identified an Est1 mutant that is deficient in this second function.
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Sara Evans  April 21, 2000
PI - Signature  Date
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INTRODUCTION

Telomeres, the nucleoprotein complexes present at the ends of chromosomes, perform an essential function in maintaining chromosomal stability. Telomeres are replicated by the reverse transcriptase, telomerase. Telomerase has received a great deal of attention in the field of cancer research, since there is a strong correlation between the presence of telomerase activity and cellular immortalization: telomerase activity is absent in most somatic cells (which have a limited replicative capacity) but is present in the majority of tumors. This observation has led to the model that reactivation of telomerase is a critical event for tumor progression. It has been suggested that inhibitors of the telomerase enzyme may be employed for therapeutic use against epithelial cancers, including breast cancer. Experimental results indicate that certain reverse transcriptase inhibitors, which target the catalytic subunit of telomerase, may indeed be a successful approach for cancer treatment. Identifying additional components of telomerase may further characterize other potential anti-telomerase targets. Towards this goal, my research proposal focuses on examining the role of a telomerase-associated protein, Est1, in telomere maintenance in the yeast Saccharomyces cerevisiae. The combined genetic and biochemical approaches that can be used with a yeast model system allows dissection of the molecular details of telomerase function — an important first step in characterizing potential targets in a mammalian system.
RESEARCH SUMMARY

BACKGROUND

Telomeres are replicated by the telomerase reverse transcriptase, which catalyzes the synthesis of telomeric repeats using an internal RNA template. In *S. cerevisiae*, EST1, EST2, EST3 and TLC1 are necessary for telomere replication by telomerase. EST2 and TLC1 encode the reverse transcriptase and RNA subunits, respectively, of the enzyme catalytic core [1, 2]. EST1 and EST3 encode additional components of the yeast telomerase holoenzyme that are required *in vivo* for telomere replication but are dispensable for enzyme catalysis, and therefore function as positive regulators of enzyme function *in vivo* [3] (Hughes et al, submitted). Previous work has shown that Estl has the properties of a single-strand telomere DNA binding protein, with a strong preference for a free single stranded 3’ end [4]. These data have led us to propose that Estl – as a component of the telomerase enzyme complex – functions in delivering or positioning the telomerase enzyme at the end of the chromosome. Presumably, Estl performs this role via its telomeric DNA binding activity, but this proposed telomere accessing activity may be mediated through protein-protein interactions as well.

A fifth gene, *CDC13*, also required for telomere maintenance, encodes a high affinity single-strand telomeric DNA binding protein that is not tightly associated with the telomerase complex [5, 6] (Hughes et al, submitted). The identification of a telomerase-defective allele of *CDC13* (*cdc13es*) indicated that this end binding protein was also required for telomerase function. We therefore proposed that Cdc13, when bound to the terminus, mediated telomerase access to the telomere.

My research proposal focuses on characterizing the role(s) of proteins that are responsible for mediating access of telomerase to the telomere, with a focus on Estl and Cdc13. The results of my research during Year I of this proposal, substantiated the role of these two proteins in conferring telomerase access to the telomere, and led to the proposal that Cdc13 mediates access of telomerase to the chromosome terminus through a direct protein-protein interaction with the Estl protein [6]. This model was based, in part, on the observation that fusion of Estl to Cdc13 resulted in substantial telomere lengthening. This suggested that the affinity of telomerase for the telomere was greatly enhanced by fusing Cdc13 to a telomerase component. Furthermore, fusion of Cdc13 to Estl bypassed telomerase defective versions of either protein. A Cdc13-Est2 fusion also resulted in telomere elongation and, strikingly, allowed cells to propagate in the complete absence of Estl function. However, the Cdc13-Est2 fusion was not able to promote telomere elongation in the absence of Estl, suggesting that Estl has a positive regulatory role in telomere replication in addition to mediating telomerase access.

This body of work left open two questions regarding aspects of Estl and Cdc13 function. First, does Estl have a second role in addition to mediating telomerase access? Second, does Cdc13 mediate telomerase access through direct recruitment of the enzyme? Towards answering these questions, my research in the past year has focused primarily on two projects. Project I reports on a set of experiments to begin investigating the second function of Estl; this is an extension of on-going work using a panel of Estl mutants generated in Year I of this proposal. Project II outlines an experimental approach designed to test the model that Cdc13 mediates telomerase access through direct recruitment of the telomerase complex. This work is still in progress, but preliminary evidence suggests that the presence of Cdc13 alone may not be sufficient to recruit telomerase to the end of the chromosome. A third project describes a series of fusion experiments extending the principles of the Cdc13-telomerase fusion studies. In Project III, I describe results from a series of experiments employing an Est3-DBDCdc13 fusion, which lends *in vivo* support to previous genetic and biochemical observations for the role of Est3 in telomere replication as a component of the telomerase complex. The results from this latter project have been submitted to Current Biology as part of a manuscript entitled “The Est3 protein is a subunit of yeast telomerase”. 

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Project I: Phenotypic analysis of a panel of estl mutants

Rationale

As previously stated, the fusion analysis suggested that Est1 may have a second role in telomere replication, in addition to co-mediating telomerase access. This proposal is based on the observation that fusion of the Est2 subunit of telomerase to Cdc13 partially bypasses the requirement for Est1 in telomere maintenance: telomeres are stably maintained in a strain carrying the Cdc13-Est2 fusion but lacking Est1, but are not elongated to the same degree as observed in the presence of Est1. Since Est1 is a terminus-specific single-strand DNA binding protein, this second proposed role for Est1 may be to promote accessibility of the 3’ end of telomeric DNA to the telomerase active site.

One trivial explanation for the failure of the Cdc13-Est2 fusion to promote telomere elongation in an estl-A strain is that there could be reduced levels of the Cdc13-Est2 telomerase complex in the absence of Est1. To address this possibility, I assayed for the ability of the Cdc13-Est2 fusion to associate with the telomerase RNA in the presence versus absence of Est1. In parallel immunoprecipitation experiments performed in EST1 and estl-A strains, there was less than a two-fold difference in the ability of the Cdc13-Est2 protein to co-immunoprecipitate the TLC1 RNA, arguing that the failure to elongate telomeres in an estl-A strain was not simply due to reduced stability of the Cdc13-Est2 fusion protein levels [6].

In Year I, I generated a panel of Est1 mutants by alanine scanning mutagenesis, and characterized these mutants both genetically for in vivo telomere replication phenotypes and biochemically for their association with the telomerase RNA. One particular class of mutants displayed telomere replication defects, although the mutant Est1 proteins still physically associated with telomerase. To explore the proposed second role of Est1, I have tested whether any of these Est1 mutants are also defective in promoting telomere elongation in the presence of the Cdc13-Est2 fusion.

Results and Discussion

One allele, estl-42, displays the characteristics expected for an Est1 mutant defective in the second function, in that estl-42 strains expressing the Cdc13-Est2 fusion protein do not have elongated telomeres (Fig. 1A). Furthermore, this mutant alone has moderately short telomeres and, interestingly, displays a conditional (temperature dependent) senescence phenotype (data not shown and Fig. 1B). Preliminary biochemical characterization of the Est1-42 protein shows that it retains association with telomerase, however the variability in that initial analysis warrants repeating the experiment (currently in progress).

The amino acids that are altered in the estl-42 mutant do not lie within the mapped DNA binding domain. However we cannot exclude the possibility that determinants required for DNA binding also lie elsewhere in the protein, and that estl-42 represents a mutant which is defective for DNA binding. Alternatively, since Est1 associates with TLC1 independently of the catalytic subunit, it is possible that Est1 is involved in direct RNA binding, and that the estl-42 mutant is somehow defective in this activity. Further biochemical analysis of the Est1-42 protein should distinguish between these possibilities, and help define the second function of Est1.

Future Directions

The mutants have thus far only been assayed for association with the telomerase RNA. Since this may or may not reflect an association with an active enzyme, I am currently in the process of testing whether telomerase activity can be detected in each mutant Est1 immunoprecipitate. The genetic and biochemical analyses of these estl mutants will form the basis for a manuscript that is
tentatively scheduled to be submitted by the end of the year. In addition, the est1 mutants remain to be tested for their DNA and RNA binding properties in vitro. One ultimate goal of this proposal is to isolate an Est1 mutant that is defective for DNA binding, as it remains unknown what the in vivo consequences of a DNA binding defect are. Thus, if none of the mutants are defective, then I plan to randomly mutagenize the DNA binding domain using PCR-based techniques, and assay these mutants for their in vitro DNA binding properties and in vivo telomere replication phenotypes. High copy suppression genetic screens employing a subset of these est1 mutants have so far been uninformative.

Project II: Does Cdc13 mediate telomerase access by directly recruiting the telomerase enzyme?

Rationale and Experimental strategy

The results of the fusion protein studies lead us to favor a model in which Cdc13 mediates telomerase access through a direct interaction with a component of the enzyme, likely Est1. However, the natural telomere consists of many other protein factors that could potentially aid Cdc13 in its recruitment function. I therefore wanted to pursue the model that Cdc13 is sufficient for telomerase recruitment, by directing Cdc13 to a non-telomeric location and asking if it brings telomerase along with it. The experimental approach involves artificially directing Cdc13 to a double strand break site (created by the site specific HO endonuclease) by virtue of the LexA DNA binding system, and determining if telomerase is recruited to the break by monitoring if telomeric sequences are added to the de novo chromosome end.

The strains I have constructed and employed in this experiment have an HO cut site inserted approximately 20kb from the end of chromosome VIII. The terminal 20kb of this chromosome does not contain any essential genes, and therefore the cell is able to maintain viability in the absence of this DNA, if the end is healed in such a manner as to create a stable chromosome (by telomere addition, for example). In this strain, the HO endonuclease gene has been placed under the inducible galactose promoter. Cleavage by the HO endonuclease creates a four base pair G-rich overhang, which can be elongated by the telomerase enzyme [7]. Prior experiments in yeast have shown that the presence of a telomere "seed" sequence near the chromosome break site will efficiently promote telomere addition [7, 8]. Therefore, as a control to ensure my system was working, I engineered a strain with G13T sequences adjacent to the HO site. This strain displayed equivalent growth on media containing either glucose or galactose, indicating that telomere healing had indeed efficiently occurred, such that there was no decrease in viability when the terminal portion of chromosome VII was lost. I then constructed strains where the G13T sequences were replaced with three tandem Lex sites, and tested for the ability of the Cdc13-LexDBD fusion protein to recruit telomerase by assaying for viability on galactose containing media.

Results and Discussion

The G13T containing strain, as expected, retained 100% viability when plated on media containing galactose (data not shown). However, strains containing the Lex sites (zero or three) - but lacking the G13T sequences - displayed a marked decrease in viability on media containing galactose (0.1% viability); the residual growth presumably arises from colonies where HO cleavage has not happened, or where other events (such as non-homologous end joining) have occurred to heal the chromosome at a low frequency. To assay whether Cdc13 could recruit telomerase, I transformed the strain harboring LexA binding sites with a high copy plasmid expressing an epitope tagged, functional Cdc13-LexADBD fusion protein; this fusion protein complements the cdc13-Δ mutation and interacts with LexA binding sites in vivo (data not shown). Following HO endonuclease induction, no increase in viability was observed on galactose containing media.
Therefore, at least in this preliminary system, it appears that the presence of Cdc13 at a chromosome end is not sufficient to promote telomere elongation. One critical control, currently being tested, is whether a Cdc13-Est1-LexA<sub>DBD</sub> fusion is capable of promoting increased viability.

**Future directions**

One drawback of the current experimental strategy is that the assay for viability may be too stringent; perhaps telomere addition in this experimental system does occur, but is too inefficient to form a functional telomere and permit colony growth. Therefore molecular methods will be used to determine if telomeric sequences are indeed added onto the chromosome end. If it is determined that Cdc13 is not sufficient for telomerase recruitment, this system can easily be manipulated to include other heterologous DNA binding sites at the chromosome break site, which may then be used - in combination with the LexA binding sites - to screen for additional factors required for telomerase recruitment.

A second aspect of the Cdc13-Est1 recruiting model that I am pursuing is whether a physical interaction between Cdc13 and Est1 can be detected; two approaches will be used to examine this. Based on the observation that a Cdc13-telomerase interaction cannot be detected by co-immunoprecipitation, we have proposed the model that a Cdc13-telomerase interaction is weak or transient. However, thus far we have only performed the immunoprecipitation experiments in unsynchronized cultures. Since telomerase is thought to replicate the telomere late in S phase, perhaps a Cdc13-telomerase association can be detected in a population of cells enriched in this stage of the cell cycle. In a second approach examining this interaction, *in vitro* pull-down assays will be used to detect a direct interaction between the Cdc13 and Est1 proteins.

Note: Several fusion protein-related projects were outlined in the Future Directions part of the Year I annual report, and the majority of these projects are currently being conducted by other members of the laboratory.

**Project III: The Est3-DBD<sub>Cdc13</sub> fusion protein**

**Rationale**

Work from our lab has shown that Est1 and Est3 associate with the telomerase core enzyme (Hughes et al, submitted). I have previously shown that telomerase access to the telomere can be greatly augmented by fusing the high affinity DNA binding domain of Cdc13 (DBD<sub>Cdc13</sub>) to the telomerase-associated Est1 protein [6]. In those experiments, the Est1-DBD<sub>Cdc13</sub> fusion, expressed under the EST1 promoter and on a single copy plasmid, conferred substantial telomere lengthening in both wild type and est1-Δ strains. Since Est3, like Est1, is telomerase-associated, one would predict that fusion of the Cdc13 DNA binding domain to Est3 should similarly enhance telomerase access and consequently increase telomere length.

**Results and Discussion**

To test this, I constructed an Est3-DBD<sub>Cdc13</sub> fusion expressed under the EST3 promoter and on a single copy plasmid; this Est3 fusion protein complemented the senescence phenotype of an est3-Δ strain (Fig. 3B), demonstrating that Est3 function had not been abolished in the context of the fusion. Introduction of the Est3-DBD<sub>Cdc13</sub> fusion protein into a wild type strain resulted in substantial telomere elongation, comparable to the lengthening conferred by the Est1-DBD<sub>Cdc13</sub> fusion (Fig. 3A). Telomere elongation was further enhanced in an est3-Δ strain (Fig. 3A, lanes 6 and 7), presumably due to the
absence of the competing wild-type Est3 protein. This effect was dependent on the presence of an intact functional catalytic core, as the Est3-DBDCdc13 fusion failed to bypass the telomere replication defect of an est2-A strain (Fig. 3B). I therefore have shown that telomerase access to the chromosome end can be greatly increased by fusing either of two telomerase-associated proteins, Est1 or Est3, to the Cdc13 DNA binding domain.

I further tested the ability of the Est3-DBDCdc13 fusion to confer enzyme access in the absence of the telomerase-mediating functions of Cdc13 and Est1. Two previously described mutations, cdc13-2nd and est1-47, appear to specifically compromise the ability of telomerase to access the end of the chromosome. This conclusion is based in part on previous observations showing that (i) the cdc13-2nd mutant phenotype can be bypassed in a strain carrying the Est1-DBDCdc13 fusion, and (ii) the telomere replication defect of the Est1-47 protein can be bypassed by fusion of Est1-47 to the DBDCdc13. Fusion of Est3 to DBDCdc13 was similarly effective in bypassing the senescence phenotype of both the cdc13-2nd and est1-47 mutations (Fig. 3C). This indicates that fusing the DBDCdc13 to a telomerase-associated protein can alleviate the requirement for the telomerase accessing functions of either Cdc13 or Est1. However, although the Est3-DBDCdc13 fusion bypassed the telomerase-accessing function of Est1, it was not capable of rescuing the telomere replication defect of an est1-A null strain (data not shown). An Est1-DBDCdc13 fusion is similarly unable to rescue an est3-A strain (data not shown). Thus, although enzyme access can be increased when either Est1 or Est3 is fused to DBDCdc13, neither fusion is sufficient to bypass the complete function of the other protein. Therefore, the roles of these two telomerase-associated proteins do not overlap. More specifically, these results demonstrate that Est3 does not function in the telomerase recruitment role previously proposed for Est1. These results have been submitted for publication in Current Biology.

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Figure 1. The est1-42 mutant does not promote telomere elongation by the Cdc13-Est2 fusion protein. Genomic Southern blot hybridized with a telomere specific probe. Plasmids expressing the EST1 or est1-42 allele (or empty vector) were cotransformed with a plasmid expressing the Cdc13-Est2 fusion into an est1-Δ cdc13-Δ/pCDC13 strain. Genomic DNA was prepared ~25 generations of growth following eviction of the CDC13 plasmid.

Figure 2. Conditional senescence phenotype of the est1-42 mutant. Each plate represents an assembled set of streakouts, grown for a number of generations as depicted in the schematic to the right. The conditional senescent mutant displays near wild type growth at 30°C, but is senescent at the higher temperature.
Figure 3. The Est3-DBD_{Cdc13} fusion increases access of telomerase to the chromosome terminus. (A) Genomic Southern blot performed as in Fig. 1; DNA was prepared ~ 50 generations of growth following plasmid transformation. EST+ control (lane 1); EST+/ Est1-DBD_{Cdc13} (lanes 2, 3); EST+/ Est3-DBD_{Cdc13} (lanes 4, 5); est3-Δ/ Est3-DBD_{Cdc13} (lanes 6, 7); all plasmids were single copy CEN vectors and expressed under the native EST gene promoter. (B) Growth after approximately 50 generations of est2-Δ or est3-Δ strains harboring either vector or single copy plasmids expressing the appropriate wild type Est protein or the Est3-DBD_{Cdc13} fusion. (C) Growth after ~ 50 generations of an est3-Δ cdc13^{est} strain or an est3-Δ est1-47 strain with plasmids expressing the wild type Est3 protein or the Est3-DBD_{Cdc13} fusion. These strains expressing the Est3-DBD_{Cdc13} fusion have been propagated a further 75 generations with no signs of senescence.
SUMMARY APPENDICES

Key research accomplishments:

- Screened a panel of Est1 mutants for those that do not promote telomere elongation by the Cdc13-Est2 fusion, and identified one candidate mutant that may be used to further define the auxiliary role for Est1.

- Devised an experimental system to test the hypothesis that Cdc13 mediates telomerase access by recruiting the enzyme to the end of the chromosome, and tentatively assessed that Cdc13 is not sufficient for this recruiting role. This system may be expanded to screen for factors that are required in conjunction with Cdc13 for telomerase recruitment.

- Provided in vivo support for the role of Est3 as a subunit of telomerase through the use of Est3-DBD<sub>Cdc13</sub> fusions.

Reportable outcomes:

A. Manuscripts:


2. Hughes TR, Evans SK, Weilbaecher RG, Lundblad V. The Est3 protein is a subunit of yeast telomerase. (submitted to Current Biology).

B. Abstracts and Presentations:

Analysis of Est1 and Cdc13: Co-mediators of telomerase access
Hal Weintraub Graduate Student Award Presentation, May 5-6, 2000
Platform Presentation

Analysis of the Roles of EST1 and CDC13 in Telomere Replication
Era of Hope Department of Defense Breast Cancer Research Program Meeting, June 8-12, 2000
Poster Presentation

C. Copies of the above cited manuscripts and abstracts are attached.
REFERENCES


ANALYSIS OF THE ROLES OF EST1 AND CDC13 IN YEAST TELOMERE REPLICATION

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Telomerase is a reverse transcriptase that specifically replicates the end of the chromosome, thereby counterbalancing the terminal sequence loss that occurs due to semi-conservative replication. In *Saccharomyces cerevisiae*, TLC1 and EST2 encode the templating RNA and reverse transcriptase subunits of telomerase, respectively, and are required both in vivo for telomere replication and in vitro for telomerase enzyme activity. In contrast, EST1, EST3 and CDC13 are required in vivo for telomerase function but are dispensable for catalytic activity in vitro. Both Est1p and Est3p are stably associated with the telomerase enzyme. Est1 is a single-stranded telomeric DNA-binding protein, suggesting that Est1 promotes accessibility of the chromosome terminus to the active site of telomerase. Cdc13 is a high-affinity single-stranded telomeric DNA-binding protein that has an essential function at the telomere, presumably in protecting the end of the chromosome, as well as a role in telomere replication. The prior identification of a telomerase-defective allele (cdc13es') of a telomere end-binding protein suggested a regulatory role for Cdc13 in telomere replication, perhaps in mediating the process by which telomerase interacts with chromosome termini. Thus, Est1 and Cdc13p display biochemical and genetic properties that are consistent with those expected for proteins that mediate telomerase access.

To define more clearly the mechanistic function of these proteins at the telomere, I have pursued two experimental approaches. The first has been a detailed site-directed mutational analysis of the Est1 protein, which has yielded three particularly informative classes of mutants. One particular allele, estl-47, displays a telomere replication defect, although the mutant Est1-47 protein still retains association with telomerase. One possibility for this mutant phenotype is that the Est1-47 protein is defective in its telomerase accessing function. In a second approach, I have examined the consequences of physically tethering telomerase components to the telomere as protein fusions with Cdc13p (or, alternatively, with the high affinity DNA-binding domain of Cdc13p). Strikingly, tethering Est1p, Est2p or Est3p in this manner confers a substantial telomere lengthening phenotype, suggesting that telomerase access to the terminus has been greatly increased. Furthermore, fusion proteins consisting of mutant versions of Cdc13p (Cdc13es') or Est1p (Est1-47) confer similar telomere elongation, indicating that close physical proximity can bypass telomerase-defective mutations in either protein. Fusing Cdc13p directly to the catalytic subunit of telomerase (Est2p) allows stable telomere maintenance in the complete absence of Est1p, consistent with a role for Est1p in mediating telomerase access. These experiments support a model in which Cdc13p mediates telomerase access by a direct interaction with the enzyme. Furthermore, these data indicate that Est1p is a comediator of this accessing function.

The U.S. Army Medical Research and Material Command under DAMD 17-98-1-8020 supported this work.
Telomerase is a reverse transcriptase that specifically replicates the end of the chromosome, thereby counterbalancing the terminal sequence loss that occurs due to semi-conservative replication. In *Saccharomyces cerevisiae*, TLC1 and EST2 encode the templating RNA and reverse transcriptase subunits of telomerase, respectively, and are required both *in vivo* for telomere replication and *in vitro* for telomerase enzyme activity. In contrast, Est1 is required *in vivo* for telomerase function but is dispensable for catalytic activity *in vitro*. Previous work in the Lundblad laboratory has shown that Est1 is a single-stranded telomeric DNA-binding protein that is stably associated with the telomerase enzyme. These findings suggest that Est1 promotes accessibility of the chromosome terminus to the active site of telomerase.

To define more clearly the mechanistic function of Est1, I have pursued two experimental approaches. The first has been a detailed site-directed mutational analysis of the protein, which has yielded three distinct classes of mutants. Detailed biochemical and genetic analysis of these mutants is one of the remaining goals of my thesis. One particular allele, est1-47, displays a telomere replication defect, although the mutant Estl-47 protein still retains association with telomerase. One possibility for this mutant phenotype is that the Estl-47 protein is defective in its telomerase accessing function.

In a second approach, I have examined the consequences of physically tethering telomerase to the telomere as a protein fusion with Cdc13. Cdc13 is a high-affinity single-stranded telomeric DNA-binding protein that has an essential function at the telomere, presumably in protecting the end of the chromosome, as well as a role in telomere replication. The prior identification of a telomerase-defective allele (*cdc13*mut) of a telomere end-binding protein suggested a regulatory role for Cdc13 in telomere replication, perhaps in mediating the process by which telomerase interacts with chromosome termini. Strikingly, a Cdc13-Est1 fusion confers a substantial telomere lengthening phenotype, suggesting that tethering Est1 to Cdc13 greatly increases access of telomerase to the terminus. Fusion proteins consisting of mutant versions of Cdc13 (*CDC13*mut) or Est1 (Est1-47) confer similar telomere elongation, indicating that close physical proximity can bypass telomerase-defective mutations in either protein. Fusing Cdc13 directly to the catalytic subunit of telomerase (Est2) allows stable telomere maintenance in the complete absence of Est1, which is consistent with a role for Est1 in mediating telomerase access. These experiments support a model in which Cdc13 mediates telomerase access by a direct interaction with the enzyme (Evans and Lundblad, 1999). Furthermore, these data indicate that Est1 is a comediator of this accessing function; experiments are currently in progress to determine if Est1 is the direct binding partner of Cdc13.

In a separate set of experiments, I have shown that increased expression of EST1, EST2, and TLC1 also causes significant telomere elongation (Hughes et al., submitted). In conjunction with the fusion experiments mentioned above, these studies suggest that telomere length homeostasis is maintained in part by restricting the access of telomerase to chromosome termini, but this limiting situation can be overcome either by directly tethering telomerase to the telomere, or by increasing the levels of functional telomerase in the cell.
Est1 and Cdc13 as Comediators of Telomerase Access

Sara K. Evans and Victoria Lundblad*
Est1 and Cdc13 as Comediators of Telomerase Access

Sara K. Evans and Victoria Lundblad*

Cdc13 and Est1 are single-strand telomeric DNA binding proteins that contribute to telomere replication in the yeast Saccharomyces cerevisiae. Here it is shown that fusion of Cdc13 to the telomerase-associated Est1 protein results in greatly elongated telomeres. Fusion proteins consisting of mutant versions of Cdc13 or Est1 confer similar telomere elongation, indicating that close physical proximity can bypass telomerase-defective mutations in either protein. Fusing Cdc13 directly to the catalytic core of telomerase allows stable telomere maintenance in the absence of Est1, consistent with a role for Est1 in mediating telomerase access. Telomere length homeostasis therefore is maintained in part by restricting access of telomerase to chromosome termini, but this limiting situation can be overcome by directly tethering telomerase to the telomere.

In most species, telomeres are composed of G-rich repetitive sequences that are elongated by telomerase (1). Several factors govern the balance between sequence addition and loss to maintain telomeres at a stable length, including positive and negative regulation of telomerase access to the chromosome terminus (2-4). In S. cerevisiae, five genes are required for the telomerase pathway (4-7). TLC1 and EST2 encode the RNA and reverse transcriptase subunits of telomerase, respectively, and as expected for subunits that are essential for catalysis, telomerase activity is absent in extracts from strains defective in EST2 or TLC1(7-9). In contrast, mutations in EST1, EST3, and CDC13 do not eliminate enzyme activity in vitro (9, 10), despite the fact that strains carrying mutations in any of these three genes have the same severe telomere replication defect as est2Δ or tlc1Δ strains (6, 10).

Both Cdc13 and Est1 bind single-strand telomeric DNA (4, 11, 12), although they make separate contributions to telomere replication and stability. Est1 is required solely for the telomerase pathway (11), whereas Cdc13 has an essential function at the telomere, presumably in protecting the end of the chromosome (13), as well as a role in telomere replication (4). This latter activity was revealed by a telomerase-defective allele of Cdc13, called cdc13<sup>est4</sup> [originally named est4 (6)], leading to the proposal that Cdc13, like Est1, mediates telomerase access (4). The two proteins also display different biochemical properties. Est1, but not Cdc13, requires a free single-strand 3' terminus for DNA binding and binds telomeric DNA with a 500-fold reduced affinity compared with Cdc13 (4, 11). In addition, Est1 is associated with telomerase, whereas Cdc13 does not exhibit a detectable interaction with the enzyme (14).

These results suggest that telomerase is recruited to the telomere due to a direct (but weak) protein-protein interaction between Cdc13 and the enzyme, and the telomere shortening in the cdc13<sup>est4</sup> mutant strain is due to a further reduction in this interaction. This model predicts that increasing the association between Cdc13 and telomerase would increase telomere length. To test this, we examined the consequences of fusing Cdc13 to the telomerase-associated protein Est1 (15). Introduction of the gene encoding this Cdc13-Est1 fusion, present on a single-copy plasmid and expressed by the CDC13 promoter, into a CDC13<sup>+</sup> strain resulted in substantial telomere elongation (Fig. 1A, lanes 2
and 3). This effect was dependent on functional telomerase, because telomeres were not elongated when the fusion was introduced into an est2Δ strain (Fig. 1A, lanes 14 to 16). Telomere elongation was even more pronounced in a cdc13Δ strain, in which telomere length increased by ~800 base pairs after ~100 generations of growth (Fig. 1A, lanes 4 and 5); continued propagation resulted in even further telomere lengthening (16).

Several experiments indicated that both Cdc13 and Est1 retained function in the context of the fusion. First, the essential function of CDC13 was fully complemented by the fusion protein (Fig. 1B). The Cdc13-Est1 fusion also complemented the senescence phenotype of an est1Δ strain (Fig. 1C). This was not due to a general bypass of the telomerase pathway, because this same fusion did not complement est2Δ or est3Δ strains (Fig. 1C). The complementation behavior of the fusion protein indicates that Est1 and Cdc13 normally function in temporal and physical proximity in their respective roles in telomere replication. Telomerase also has been shown to coimmunoprecipitate with Est1 but not with Cdc13 (Fig. 1D) (14). The Cdc13-Est1 fusion protein coimmunoprecipitated the RNA subunit of the telomerase complex (Fig. 1D) and enzyme activity (17), indicating that telomerase is associated with the fusion protein.

These results suggest that the proposed recruitment function of Cdc13 can be enhanced by fusing it to a telomerase component, and predict that the telomerase-defective cdc13Δ mutation would be bypassed in a fusion. Consistent with this prediction, a fusion containing the mutant Cdc13Δ protein behaved indistinguishably from the wild-type (WT) fusion: Telomere elongation occurred to the same degree in both CDC13+ and cdc13Δ strains (Fig. 1A, compare lanes 6 to 9 and 2 to 5), and no senescence was observed when the Cdc13Δ-Est1 fusion was introduced into a cdc13Δ strain (17). Similar results were observed in a reciprocal experiment with a mutant allele of EST1 (est1-47); this mutation disrupts telomere replication

Fig. 1. The Cdc13-Est1 fusion confers telomere elongation. (A) Telomere Southern (DNA) blots were performed as in (6). The bracket indicates a heterogeneous telomeric band that represents about two-thirds of the telomeres in this strain background. Cultures for lanes 1 to 13 were grown for ~100 generations before DNA preparation. Lanes 1, 3, 4, CDC13+ EST1 control strain; lanes 2 to 5, 10, 15, and 16, pVL1091 (expressing the Cdc13-Est1 fusion); lane 6 to 9, pVL1092 (cdc13Δ-Est1); lane 12, pVL1098 (Cdc13Δ-Est1-47); the relevant genotypes of the strains are indicated. Plasmids were introduced into strains deleted for CDC13 for this and subsequent figures by first transforming into a cdc13Δ/pVL438 (CDC13+ URA3−) strain followed by subsequent eviction of pVL438 by plating on media containing 5-fluoro-orotic acid. Molecular sizes are indicated on the left (in kilobase pairs). (B) cdc13Δ strains containing pVL648 (CDC13+), pVL1091, or pVL676 (cdc13Δ) (13) were grown at 23°C, and equivalent numbers of cells, as serial 10-fold dilutions, were plated at 25° and 36°C. (C) Growth after ~50 generations of est1Δ, est2Δ, or est3Δ strains, with single-copy plasmids bearing the CDC13-EST1 fusion gene or the appropriate WT EST gene (each under their native promoter). (D) Immunoprecipitation from extracts prepared from strains expressing proteins with a triple hemagglutinin epitope (HA3) introduced at the NH2-terminus: HA-Est1 (pVL1086), HA-Cdc13 (pVL841), or HA-Cdc13-Est1 (pVL1102), followed by detection of the telomerase RNA (TLC1) levels in the extract (E) and immunoprecipitates (P) by Northern (RNA) blotting (27).
(Fig. 1C), although the mutant Est1 protein still physically associates with telomerase (18), suggesting a defect in the same telomerase-accessing function that is altered by the cdc13
sup strain. This defect was bypassed when the Est1-47 protein was fused to Cdc13. In fact, the double-mutant fusion protein (Cdc13
sup fused to Est1-47) conferred the same degree of telomere elongation as the WT fusion protein, even in a strain deleted for both est1-Δ and cdc13-Δ (Fig. 1A, lane 12); moreover, this strain did not exhibit senescence (17). The ability of the double-mutant fusion to complement a cdc13-Δ est1-Δ strain indicates that the fusion is acting as a dimeric molecule that bridges telomerase and the telomere.

One alternative interpretation of our data is that telomere elongation is due to perturbation of chromatin structure, rather than to increased access of telomerase to the telomere. In particular, telomere lengthening could be a secondary consequence of altered Cdc13 function, because mutations in CDC13 have been identified that increase telomere length (19, 20). However, these recessive alleles of CDC13 have a set of genetic and biochemical features that distinguish them from the gain-of-function properties of the Cdc13-Est1 fusion (17, 20). In addition, fusion of several unrelated protein sequences, or an inactive telomerase subunit (see below), to the COOH-terminus of either the WT Cdc13 protein or the mutant Cdc13
sup protein does not increase telomere length (17, 21). The most direct argument against this alternative interpretation is the result of an experiment in which we examined the behavior of a fusion in which only the high-affinity DNA binding domain of Cdc13 (DBD$_{Cdc13}$) was fused to Est1 (15). This experiment was based on our previous demonstration that DBD$_{Cdc13}$ can be expressed as a stable, functional subdomain (22) and therefore could be used as an alternative means of directing Est1 to the telomere with high efficiency, while leaving the full-length Cdc13 protein intact. As predicted, the Est1-DBD$_{Cdc13}$ fusion conferred extensive telomere lengthening in a CDC13
sup strain (Fig. 2A) and bypassed senescence of a cdc13-Δ strain (Fig. 2B). Furthermore, fusion of the defective Est1-47 protein to DBD$_{Cdc13}$ bypassed both est1-Δ and cdc13
sup mutations (Fig. 2B), even though the est1-47 allele fails to complement either mutation (Fig. 1C) (17). Expression of either Est1 or DBD$_{Cdc13}$ had no effect on telomere length or viability in CDC13
sup or cdc13
sup strains, and the Est1-DBD$_{Cdc13}$ fusion failed to rescue the inviability of a cdc13-Δ strain (Fig. 2A) (17). Thus, the telomere lengthening properties of these fusions are likely to be a consequence of delivery of telomerase to the telomere, rather than a perturbation of Cdc13 function.

We next fused Cdc13 directly to Est2, the catalytic subunit of telomerase (8). The Cdc13-Est2 fusion (15) resulted in telomere lengthening to levels comparable to that of the Cdc13-Est1 fusion (Fig. 3A, lanes 3 and 4). The fusion complemented cdc13-Δ and est2-Δ null mutations, and telomere elongation occurred to the same degree in est2-Δ and EST2
sup strains (17). A Cdc13-Est2$_D70A$ fusion, containing an Asp to Ala mutation at position 670 in the active site of Est2 (8), did not confer extensive telomere elongation but instead maintained telomere length at WT levels in a cdc13-Δ EST2
sup strain (Fig. 3A, lanes 5 and 6) (23), showing that telomere elongation is only observed when a catalytically active version of telomerase is tethered to the telomere. Strikingly, the Cdc13-Est2 fusion allowed cell growth in the complete absence of Est1 function, because an est1-Δ strain carrying this fusion was viable for more than 250 generations (Fig. 3B) (24).

Long-term propagation in the absence of Est1 was not due to a previously described alternative pathway that can maintain telomeres in the absence of telomerase function (6, 25): Telomeres in an est1-Δ strain carrying the Cdc13-Est2 fusion were stably maintained at a length slightly below that of WT telomere length (Fig. 3C), with none of the striking changes in telomere structure that characterize the alternative pathway (6, 25). The ability of the Cdc13-Est2 fusion to maintain an est1-Δ strain required tethering of a functional telomerase, because an est1-Δ strain carrying the Cdc13-Est2$_D70A$ fusion exhibited senescence (Fig. 3B). This supports the hypothesis that a critical function of the Est1 protein is to mediate access of telomerase to the telomere. Notably, neither the Cdc13-Est1 fusion nor the Cdc13-Est2 fusion bypassed the requirement for Est3 (Fig. 1C) (17), showing that Est1 and Est3 perform functionally distinct roles in telomere replication.

Our results are consistent with a model in which Cdc13 mediates telomerase access by a direct interaction with the enzyme (Fig. 4). Furthermore, these data indicate that Est1 is a co-factor of this "accessing" function, potentially as a direct binding partner of Cdc13, although we cannot rule out the possibility of...
additional intervening protein or proteins. These experiments may have also uncovered an additional role for Est1 in telomerase function, as the Cdc13-Est2 fusion was not capable of promoting extensive telomere elongation in the absence of Est1 (Fig. 3C). Because Est1 is a terminus-specific DNA binding protein (11), we speculate that this second role may be to promote accessibility of the 3’ terminus to the active site of telomerase.

References and Notes
15. In-frame fusion proteins were constructed as follows: The Cdc13-Est1 fusion (pV11091) fused Cdc13 and Est1, the Cdc13-Est2 (pV11072) and Cdc13-Est2 (pV11111) proteins fused Cdc13 and Est2. All three constructs were expressed from the genomic CDC13 promoter and derived from the single-copy CEN plasmid pRS415. The Est1-DBD-Cdc13 and Est1-DBD-Cdc13 fusions (pV1120 and pV1121, respectively), expressed in single copy from the EST1 promoter, fused the Cdc13 DNA binding domain (22) to the COOH terminus of Est1, to generate Est1 -4kb - Cdc13. Telomere elongation by any of these fusion proteins is not due to increased protein expression in the context of the fusion, because overexpression of Est1, Est2, or Cdc13, or expression of Est1 by the CDC13 promoter, has little or no effect on telomere length (8, 11, 17, 19). We have also observed up to a ~4-kb increase in the length of a single telomere (chromosome III). Strains with greatly elongated telomeres do not exhibit any discernable growth defect but were not examined for more subtle defects (such as alterations in cell cycle progression). We have not investigated yet whether the increase in telomere length in strains carrying Cdc13-telomerase fusions occurs at a constant rate, or if there is some influence of cis-inhibition in elongation, as previously observed (3).
16. The est1-47 mutation is one of a panel of alanine scan mutations in EST1 (17); the Est1-47 mutant protein still retains association with telomerase (at 20% of WT levels), as assessed by coimmunoprecipitation with the TLC1 RNA (27).
The Est3 protein is a subunit of yeast telomerase

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EST1, EST2, EST3 and TLC1 function in a single pathway for telomere replication in the yeast Saccharomyces cerevisiae, as would be expected if these genes all encode component of the same complex. Previous work has shown that Est2p, the reverse transcriptase protein subunit, and TLC1, the templating RNA, are subunits of the catalytic core of yeast telomerase. We show here that Est3p is a stable component of the telomerase holoenzyme and furthermore, association of Est3p with the enzyme requires an intact catalytic core. As predicted for a telomerase subunit, fusion of Est3p to the high affinity Cdc13p telomeric DNA binding domain greatly increases access of telomerase to the telomere. Like Est3p, Est1p is also tightly associated with telomerase, although in contrast to Est3p, Est1p is capable of forming a stable TLC1-containing complex even in the absence of Est2p or Est3p. Yeast telomerase therefore contains a minimum of three Est proteins for which there is both in vivo and in vitro evidence for their role in telomere replication as subunits of the telomerase complex.
Results and Discussion

Maintenance of chromosome termini, or telomeres, requires the action of the telomerase reverse transcriptase, which counterbalances the terminal sequence loss that occurs due to incomplete semi-conservative replication. Telomere length is normally maintained at a stable set point for a given organism, indicating that there is a regulated balance between sequence loss and addition [1]. One way of modulating the extent of telomere addition is by regulating access of telomerase to the end of the chromosome through both positive and negative mechanisms [2-6]. Telomerase-associated proteins may function in such regulation. Thus, identifying and characterizing such factors is a logical starting point for developing a more complete understanding of the mechanisms of telomere length regulation.

In the budding yeast *S. cerevisiae*, the reverse transcriptase catalytic subunit and the templating RNA are encoded by the *EST2* and *TLC1* genes, respectively [7,8]. Strains deleted for either of these two genes exhibit in vivo telomere replication defects [8,9], and extracts prepared from these strains are deficient for telomerase activity [7,10]. Mutations in *EST1, EST3* or *CDC13* also eliminate telomere replication in vivo [5,9,11-13] but are dispensable for in vitro telomerase catalytic activity [10,14]. However, this does not rule out the possibility that one or more of these three proteins performs its role as a component of the telomerase ribonucleoprotein particle (RNP). Consistent with this premise, Est1p is a single strand telomeric DNA binding protein [15] that has been shown to associate with telomerase, both when over-expressed [16-18] and in single copy (Evans and Lundblad 1999; this work). Cdc13 also binds single strand telomeric DNA [5,19] but is not tightly associated with telomerase (Evans and Lundblad 1999; this work), suggesting that it instead is bound to chromosome termini [20]. We have recently proposed that Est1p and Cdc13p, as components of telomerase and telomeric chromatin, respectively, cooperate to directly recruit telomerase to the end of the chromosome [6].
In contrast to Estlp and Cdc13p, Est3p has not yet been biochemically characterized and thus its specific role in telomere replication is unclear. To address whether Est3p is a component of the telomerase enzyme, we examined whether it is associated with the TLC1 RNA and with enzyme activity, in parallel with Estlp, Est2p and Cdc13p. In order to avoid artifactual associations that might arise from over-expression, we constructed a set of strains containing epitope-tagged versions of each gene integrated into the respective endogenous locus under the control of the native promoter (Table 1). The identical HA3 epitope was introduced into a region of each protein such that the activity of the tagged protein was not overtly affected, as assessed by examining the telomere length of these strains. Figure 1 shows that little or no alterations in telomere length were observed in strains bearing integrated versions of HA3-EST1, HA3-EST2, EST3-HA3 or HA3-CDC13.

Immunoprecipitates from extracts prepared from these four individually tagged strains were assayed for the presence of the relevant recombinant protein, the telomerase RNA subunit (TLC1) and telomerase activity (Fig. 2). HA3-Est1, HA3-Est2, and HA3-Cdc13 proteins could be detected in the immunoprecipitates (Fig. 2A, top panel), but were not visible in the starting extracts (data not shown), due to the low levels of each of these proteins when expressed under their native promoter. The Est3-HA3 protein co-migrates with the antibody heavy chain and therefore was masked in this particular western blot (however, see Fig. 3A for an alternative means of Est3p detection). The TLC1 RNA subunit was readily detectable in Estlp, Est2p and Est3p immunoprecipitates by northern analysis, but was not present in immunoprecipitates prepared from an untagged strain (Fig. 2A, middle panel). The interactions with TLC1 were specific for the telomerase RNP, since the U1 small nuclear RNP (snRNP) RNA was not present in each immunoprecipitate (Fig. 2A, bottom panel). In multiple repeats of this experiment in which the two RNA species were monitored quantitatively, typically 1-5% of the
telomerase RNA present in the starting extract was immunoprecipitated, whereas the more abundant U1 snRNP RNA was recovered at an efficiency of $\leq 0.02\%$. This interaction between the Est proteins and the TLC1 RNA also reflected an association with an active telomerase complex, since enzyme activity was identified in each of the Est protein immunoprecipitates (Fig. 2B). Two related activity assays were performed using the primer substrate (5' -TGTGGTGTGTGGG-3'); both assays, as well as the primer, have previously been used to monitor telomerase activity from *S. cerevisiae* [7,10,14]. In the first assay, enzyme extension products were assayed in the presence of the chain-terminating ddGTP nucleotide, which results in extension of this primer by two nucleotides. This concentrates the signal in the product labeled "+2" and thus provides a sensitive measure for detection of enzyme activity [7,14]. Telomerase activity was readily observed in each Est protein immunoprecipitate using this assay (Fig. 2B, lanes 1-5); activity was RNase A-sensitive and primer-dependent in each case (data not shown). In several repetitions of this experiment, the levels of enzyme activity, which correlated directly with the levels of TLC1 that were recovered, varied by no more than 2-fold when compared among the three Est protein immunoprecipitates (data not shown), as would be predicted if these three proteins, each containing the identical HA$_3$ epitope, were subunits of the same complex. Enzyme activity was also monitored using a non-terminating reaction (Fig. 2B, lanes 6-10) that generates a ladder of reaction products representing the incomplete extension of a primer across the TLC1 template [10,21]. Telomerase activity (which was RNase A-sensitive; data not shown) was again detected in HA$_3$-Est1p and Est3p-HA$_3$ immunoprecipitates at roughly the same levels as observed in HA$_3$-Est2p immunoprecipitates (compare lanes 6 and 7 with lane 10). In both sets of assays, immunoprecipitates were subjected to relatively stringent wash conditions (0.4 M NaCl) prior to enzyme assays. Therefore, Est3p is tightly associated with an active telomerase complex, demonstrating that Est3p plays its role in telomere replication as a subunit of telomerase. These experiments also show that Est1p is a
stable component of the enzyme, confirming and extending previous observations that showed that Est1, when over-expressed, was associated with the TLC1 RNA [16-18].

In contrast, neither the TLC1 RNA subunit (Fig. 2A) nor enzyme activity (Fig. 2B) were detectable in Cdc13p immunoprecipitates performed under the same high stringency conditions. Even under less stringent conditions, telomerase activity was not detected in Cdc13p immunoprecipitates (data not shown). Thus, Cdc13p is not tightly associated with telomerase, although these observations do not exclude the possibility of a transient interaction.

To determine the requirements for interaction of Est3p with telomerase, we examined whether Est3p association with a TLC1-containing complex could be retained in the absence of either Est1p or Est2p. A GST-pulldown assay was used to determine levels of the Est3 protein in the deletion strains used in this experiment, since Est3p comigrated with the immunoglobulin heavy chain when immunoprecipitated. Deletion of EST1 or EST2 did not affect the level of Est3-HA3-GST protein recovered from whole cell extracts on glutathione beads (Fig. 3A, lower panel). In addition, the steady state levels of TLC1 RNA, which were normalized to the levels of U1 RNA, were also unchanged in extracts prepared from strains that lacked either Est1p, Est2p or Est3p (Fig. 3B). When immunoprecipitated, Est3p retained association with a TLC1-containing complex in a strain deleted for EST1 (Fig. 3A, lanes 3 and 4). However, the ability of Est3p to co-immunoprecipitate with TLC1 was abolished in the absence of the Est2p subunit (Fig. 3A, lanes 5 and 6), indicating that an intact catalytic core was a requirement for the interaction of Est3p with the telomerase complex. Whether this reflects a direct interaction between the Est2 and Est3 proteins has not yet been determined; two hybrid tests intended to detect such an interaction have been so far negative (D.K. Morris and V.L., unpublished data).

We similarly investigated the requirement for the association of Est1p with telomerase. For these experiments, we constructed a strain containing differentially tagged versions of Est1p
and Est2p (HA3-Est1p and myc3-Est2p) integrated into the genome and expressed under the respective native EST1 or EST2 promoters. Est3p is not required for the interaction of Est1p with the enzyme (Fig. 3C, lane 7), indicating that Est1p and Est3p associate with telomerase independently of each other. Unlike Est3p, however, the Est1 protein was capable of forming a TLC1-containing complex even in the absence of the catalytic Est2p subunit (Fig. 3C, lane 5), indicating that Est1p and Est2p interact with the TLC1 RNA independently of one another. Similar results for Est2p- and Est3p-independent association of Est1p with a TLC1 complex have been reported, using an over-expressed LexA-Est1 protein fusion [18].

We have previously shown that telomerase access to the telomere can be greatly augmented by fusing the high affinity DNA binding domain of Cdc13p (DBD<sub>Cdc13</sub>) to the telomerase-associated protein Est1p [6]. In those experiments, the Est1-DBD<sub>Cdc13</sub> fusion, expressed under the EST1 promoter and on a single copy plasmid, conferred substantial telomere lengthening in both wild type and est1-A strain [6]. The above experiments demonstrating that Est3p, like Est1p, is telomerase-associated predict that fusion of the Cdc13 DNA binding domain to Est3p should similarly enhance telomerase access and consequently increase telomere length. To test this, we constructed an Est3-DBD<sub>Cdc13</sub> fusion expressed under the EST3 promoter and on a single copy plasmid; this Est3 fusion protein complemented the senescence phenotype of an est3-A strain (Fig. 4B), demonstrating that EST3 function had not been abolished in the context of the fusion. Introduction of the Est3-DBD<sub>Cdc13</sub> fusion protein into a wild type strain resulted in substantial telomere elongation, comparable to the lengthening conferred by the Est1-DBD<sub>Cdc13</sub> fusion (Fig. 4A). Telomere elongation was further enhanced in an est3-A strain (Fig. 4A, lanes 6 and 7), presumably due to the absence of the competing wild-type Est3 protein. This effect was dependent on the presence of an intact functional catalytic core, as the Est3-DBD<sub>Cdc13</sub> fusion failed to bypass the telomere replication defect of an est2-A strain (Fig. 4B). We therefore have shown that telomerase access to the chromosome end can be greatly increased by fusing either of two telomerase-associated proteins,
Est1p or Est3p, to the Cdc13 DNA binding domain. This is not a non-specific effect of the DBD_{Cdc13} but instead is a specific consequence of fusing the DBD_{Cdc13} to a telomerase subunit, since expression of either the DBD_{Cdc13} alone, or as a fusion to two other proteins implicated in telomere length maintenance (Stn1 and Pol1; [22,23]), does not result in telomere elongation (S.K.E., A. Chandra, E. Pennock and V.L., data not shown).

We further tested the ability of the Est3-DBD_{Cdc13} fusion to confer enzyme access in the absence of the telomerase-mediating functions of Cdc13p and Est1p. Two previously described mutations, cdc13-2est and est1-47, appear to specifically compromise the ability of telomerase to access the end of the chromosome [6]. This conclusion is based in part on previous observations showing that (i) the cdc13-2est mutant phenotype can be bypassed in a strain carrying the Est1-DBD_{Cdc13} fusion, and (ii) the telomere replication defect of the Est1-47 protein can be bypassed by fusion of Est1-47p to the DBD_{Cdc13} [6]. Fusion of Est3p to DBD_{Cdc13} was similarly effective in bypassing the senescence phenotype of both the cdc13-2est and est1-47 mutations (Fig. 4C). This indicates that fusing the DBD_{Cdc13} to a telomerase-associated protein can alleviate the requirement for the telomerase accessing functions of either Cdc13 or Est1. However, although the Est3-DBD_{Cdc13} fusion bypassed the telomerase-accessing function of Est1p, it was not capable of rescuing the telomere replication defect of an est1-Δ null strain (data not shown). An Est1-DBD_{Cdc13} fusion is similarly unable to rescue an est3-Δ strain (data not shown). Thus, although enzyme access can be increased when either Est1p or Est3p is fused to DBD_{Cdc13}, neither fusion is sufficient to bypass the complete function of the other protein. Therefore, the roles of these two telomerase-associated proteins do not overlap. More specifically, these results demonstrate that Est3p does not function in the telomerase recruitment role previously proposed for Est1p.

The demonstration that the Est3 protein is a component of telomerase completes the prediction for the functions of EST1, EST2, EST3 and TLC1 based on genetic observations.
Previous epistasis analysis had shown that these four genes function in a single pathway for telomere replication, as would be expected if these genes each encode subunits of the same enzyme complex [9,14]. The work presented here, combined with previous observations, shows that all four genes encode components of the telomerase enzyme. The polymerase core is generally considered to be the minimum subset of components required to perform the basic catalytic action which reflects that of the enzyme in vivo, whereas the holoenzyme has either additional properties or augmented activity, such as the ability to respond appropriately to different substrates. The association of the Est1 and Est3 proteins with the telomerase catalytic core, and the requirement for Est1p and Est3p \textit{in vivo} to mediate telomerase function, mirrors the behavior of holoenzyme components of other polymerases. A role for the Est1 protein subunit has been previously proposed, as a bridging protein between telomerase and the telomere. Additional genetic and biochemical investigation should help reveal the precise biochemical activity of the Est3 telomerase protein subunit.
Materials and methods

Yeast strains and plasmids

AVL78 (MATa leu2 trp1 ura3-52 prb prc pep4-3) and isogenic derivatives, as described below, were used for all experiments in Figures 1, 2 and 3. Replacement of native chromosomal sequence with the epitope-tagged version was achieved by a pop-in/pop-out strategy, as described previously. Integrating constructs (derived from YIpplac211) are as follows: pVL816 (HA3-EST1), pVL846 (HA3-EST2), pVL845 (myc3-EST2), pVL901 (EST3fsc-HA3-GST; EST3fsc is a frameshift corrected version of EST3 [12]), pVL983 (EST3fsc-HA3-HMK-His6), pVL842 (HA3-CDC13). The HA3 tag was the same for all these constructs [(YPYDVPDYA)2RSMYPYDVPDYA] and was inserted in frame at the amino terminus of Est1p (between aa 1 and 2), Est2p (between aa 11 and 12) or Cdc13p (between aa 1 and 2) or at the carboxy terminus of Est3p (after the terminal aa). The myc3 tag [(EQKLISEEDL)3] was inserted at the amino terminus of Est2p (between aa 11 and 12). In each case, insertion of the DNA sequence encoding the epitope tag introduced a few additional amino acids, which are not indicated. Deletion derivatives for the experiments in Fig. 3 were made by one-step gene disruption directly into tagged haploid strains, and cultures were grown from freshly disrupted strains; these deletion strains were not permanently stored due to their senescent characteristics. EST1 was disrupted with a kanamycin cassette derived by PCR of plasmid pFA6a.kanMX2 [24] which removed 89% of the EST1 coding region. EST2 was disrupted with pVL363 (est2-Δ1::URA3, which removed 76% of the coding sequence) or by PCR-mediated gene deletion (est2-Δ2::kan, derived from pFA6a.kanMX2, which removed the entire EST2 coding region, 51 bases of upstream sequence, and 49 bases of downstream sequence). EST3 was disrupted with pVL418 (est3-Δ2::URA3, which removed 63% of the coding sequence).

Haploid strains for Figure 4 were generated from dissection of diploid strains DVL172 (to generate est2-Δ), DVL322 (to generate est3-Δ and est1-Δ est3-Δ) and DVL323 (to generate...
est3-Δcdc13-2^ext). Plasmids used in these experiments expressed each gene on a single copy plasmid under the appropriate EST gene native promoter: pVL499 (Est1), pVL1293 (Est1-47), pVL1048 (Est2), pVL1005 (Est3fsc-HA3-HMK-His6), pVL1120 (Est1-DBDCdc13), pVL1292 (Est3fsc-HA3-DBDCdc13). pVL1292 was constructed by inserting the DBD of Cdc13p (aa 2-19, 452-693) in frame at the end of Est3fsc-HA3.

Immunoprecipitations

For western and northern analysis, extracts were prepared by lysis with glass beads essentially as previously described [14], with the exception that extracts were prepared in TMG (10 mM Tris-HCl pH 8.0, 1 mM MgCl₂, 5% glycerol) plus 300 mM NaCl, 0.5% Tween-20, 1 mM PMSF, 1 u/ml RNase inhibitor (RNasin, Promega), and 0.1 mM DTT, and immunoprecipitates were washed three times in TMG plus 300 mM NaCl and once in TMG. For telomerase activity assays, cultures were grown at 30°C in YPD and harvested at an OD₆₀₀ of 1.0. All subsequent steps were performed at 4°C. The cell pellet was washed once in water, once in TMG plus 300 mM NaCl, and resuspended in 2 pellet volumes TMG plus 300 mM NaCl, 0.5 mM DTT, and 0.25 mM PMSF. Cells were frozen in liquid nitrogen and homogenized with a mortar and pestle. RNasin (40 units / ml) and Tween-20 (0.5% final concentration) were added to the crude extract before clearing twice by centrifugation for 15 minutes at 14,000 rpm. ~2.5 mg extract in 0.5 ml was incubated for 45 min with 1 μl anti-HA high affinity antibody (Boehringer Mannheim). 20 μl settled volume Protein G Plus / Protein A Agarose was added and incubated for an additional 90 min. After gentle pulse centrifugation, the beads were washed 3 times with TMG plus 400 mM NaCl, 0.5% Tween-20, 0.5 mM DTT, 40 units/ml RNasin, one time with TMG plus 50 mM NaCl, 0.5 mM DTT, 40 units/ml RNasin, and finally resuspended with an equal volume of the latter buffer. Telomerase activity recovered by this method retains at least 50% activity following a freeze-thaw cycle with storage at -70°C.

Telomerase activity was assayed by incubating 5 μl of the agarose bead slurry containing HA-
immune complexes with 5 μl of a 2x reaction mixture containing 5 μM substrate oligonucleotide [5'-TGTGGTGTTGTTGGG-3'], 80 mM Tris-Cl (pH 8.0), 100 mM NaCl, 10% glycerol, 5 mM MgCl₂, 1 mM spermidine, 1 mM DTT, and either 200 μM TTP and 1.5 μl [α³²-P] dGTP (3000 Ci/mmol), or 800 μM ddGTP and 1.5 μl [α³²-P] TTP (3000 Ci/mmol). The reaction proceeded for 20 minutes at 30°C and was stopped by the addition of EDTA to 25 mM. Reaction products were extracted with phenol/chloroform, ethanol precipitated and resolved in a 15% acrylamide / 7 M urea sequencing gel.

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**Figure 1.** Addition of epitope tags to EST1, EST2, EST3, and CDC13 has a minimal effect on telomere length. Genomic DNA was prepared from strains carrying integrated, epitope tagged versions of each gene as indicated, and subjected to Southern blotting and hybridization with a telomere-specific probe as described previously [9]. Telomeric restriction fragments are indicated by arrows and a bracket. Each pair of lanes is derived from cells grown approximately 70 and 100 generations following replacement of the wild-type allele. Strains are as follows: untagged (AVL78; lanes 1, 2, 11, 12); HA3-Est1 (TVL288; lanes 3, 4); HA3-Est2 (TVL292; lanes 5, 6); Est3-HA3-GST (TVL293; lanes 7, 8); HA3-Cdc13 (TVL290; lanes 9, 10).

**Figure 2.** Est1, Est2 and Est3 proteins associate with an active telomerase enzyme complex. (A) Extracts from strains, each bearing an HA3 epitope tag on a single gene as indicated, were immunoprecipitated and analyzed by western blotting to detect HA tagged protein (top) and northern blotting to detect TLC1 RNA (middle) or U1 RNA (bottom). HA3-Est1 (TVL288; lane 2); HA3-Est2 (TVL292; lane 3); Est3-HA3-GST (TVL293; lane 4); HA3-Cdc13 (TVL290; lane 5); untagged (AVL78; lane 6). TLC1 and U1 RNA levels in crude extracts prepared from an untagged strain, prior to immunoprecipitation, are shown in lane 1; the amount of extract represents 4% of the amount loaded in the IP lanes. Lanes 2 - 6 are equivalently loaded, as assessed by total protein. The Est3-HA3-GST polypeptide co-migrates with the antibody heavy chain (which is not shown) and is not detectable on this particular blot. (B) Immunoprecipitates from each tagged strain were assayed for telomerase activity using either a chain terminating reaction in the presence of ddGTP (lanes 1 - 5; the band representing the +2 product is indicated) or non-chain terminating reaction (lanes 6 - 10). The strains used in this experiment are identical to those used in (A), with the exception of the Est3-HA3 strain (TVL307).

**Figure 3.** The association of Est3p with the telomerase RNP requires an intact catalytic core. (A, top) Extracts prepared from an Est3-HA3-GST strain (TVL293) or deletion derivatives of TVL293 were immunoprecipitated, and TLC1 RNA was detected by northern blotting. TVL293
(lanes 1, 2); *estl*-Δ derivative of TVL293 (lanes 3, 4); *est2*-Δ derivative of TVL293 (lanes 5, 6); untagged (AVL78; lanes 7, 8). (Bottom) Extracts were purified on glutathione sepharose and analyzed by western blotting with an antibody to detect the HA3-tagged Est3 protein.

(B) Extracts prepared from *estl*-Δ (lane 2), *est2*-Δ (lane 3), *est3*-Δ (lane 4) and *tlc1*-Δ strains (lane 5), in parallel with an ESTf strain (lane 1), were analyzed by northern blotting to detect TLC1 RNA and U1 RNA, followed by quantitation of the relative RNA levels by PhosphorImager analysis. The abundance of TLC1 RNA was reduced by no more than threefold in the three EST deletion strains, relative to the ratio of the two RNAs in a wild type strain.

(C) Extracts from TVL300 (containing HA3-Est1p and myc3-Est2p) or deletion derivatives of strain TVL300 were immunoprecipitated with either anti-myc or anti-HA antibodies and analyzed by northern blotting to detect TLC1 RNA. TVL300 (lanes 1, 2); *estl*-Δ derivative of TVL300 (lanes 3, 4); *est2*-Δ derivative of TVL300 (lanes 5, 6); *est3*-Δ derivative of TVL300 (lanes 7, 8); *tlc1*-Δ derivative of TVL300 (lanes 9, 10); untagged (AVL78; lanes 11, 12). The variations in TLC1 signal in HA versus myc immunoprecipitations are due, at least in part, to the fact that the myc3 tag is a more efficiently immunoprecipitated epitope, as myc3-Est2p co-immunoprecipitates more telomerase RNA than HA3-Est2p (data not shown).

**Figure 4.** The Est3-DBD<sub>Cdc13</sub> fusion increases access of telomerase to the chromosome terminus. (A) Genomic Southern blot performed as in Fig. 1; DNA was prepared ~ 50 generations of growth following plasmid transformation. EST<sup>+</sup> control (lane 1); EST<sup>+</sup>/pVL1120 (Est1-DBD<sub>Cdc13</sub>; lanes 2, 3); EST<sup>+</sup>/pVL1292 (Est3-DBD<sub>Cdc13</sub>; lanes 4, 5); *est3*-Δ/pVL1292 (Est3-DBD<sub>Cdc13</sub>; lanes 6, 7); all plasmids were single copy CEN vectors and expressed under the native EST gene promoter. (B) Growth after approximately 50 generations of *est2*-Δ or *est3*-Δ strains harboring either vector or single copy plasmids expressing the appropriate wild type Est protein or the Est3-DBD<sub>Cdc13</sub> fusion. (C) Growth after ~ 50 generations of an *est3*-Δ cdc13<sup>est</sup> strain or an *est3*-Δ *estl*-47 strain with plasmids expressing the wild type Est3 protein or
the Est3-DBD$_{Cdc13}$ fusion. These strains expressing the Est3-DBD$_{Cdc13}$ fusion have been propagated a further 75 generations with no signs of senescence.
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