Direct involvement of the TEN domain at the active site of human telomerase

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ABSTRACT

Telomerase is a ribonucleoprotein that adds DNA to the ends of chromosomes. The catalytic protein subunit of telomerase (TERT) contains an N-terminal domain (TEN) that is important for activity and processivity. Here we describe a mutation in the TEN domain of human TERT that results in a greatly increased primer $K_d$, supporting a role for the TEN domain in DNA affinity. Measurement of enzyme kinetic parameters has revealed that this mutant enzyme is also defective in dNTP polymerization, particularly while copying position 51 of the RNA template. The catalytic defect is independent of the presence of binding interactions at the 5'-region of the DNA primer, and is not a defect in translocation rate. These data suggest that the TEN domain is involved in conformational changes required to position the 3'-end of the primer in the active site during nucleotide addition, a function which is distinct from the role of the TEN domain in providing DNA binding affinity.

INTRODUCTION

Telomeres, the protective cap on the end of linear eukaryotic chromosomes, are composed of short G-rich DNA repeats bound by sequence-specific proteins (1). The telomeres of unicellular eukaryotes and of the germ cells of multicellular organisms are maintained by the ribonucleoprotein enzyme telomerase, which was first identified in the ciliated protozoan Tetrahymena thermophila (2). Telomerase activity has been detected in ~90% of human cancers, providing a telomere maintenance mechanism that is necessary for their unlimited growth (3–5).

The catalytic protein subunit of telomerase, telomerase reverse transcriptase [TERT (6)], contains protein motifs that are conserved with other reverse transcriptases (’RT motifs’) and motifs in its amino terminal region that are conserved among TERTs from different species. A defined region of the telomerase RNA subunit is used as the template for addition of DNA repeats onto telomeres (7). A complex consisting of TERT and telomerase RNA is sufficient to reconstitute in vitro telomerase activity from Tetrahymena and human enzymes, measured by processive extension of a DNA oligonucleotide with telomeric DNA repeats (8,9). The repeated cycling of reverse transcription from a short segment of RNA, followed by translocation to the beginning of this sequence, leads to the unique property of telomerase known as ‘repeat addition processivity’.

There is evidence for telomerase from several species that there are multiple sites of interaction with different regions of the DNA primer. For example, the affinity of Tetrahymena telomerase for DNA decreased in a stepwise manner as the primer was reduced in length from 18 to 12 nt then 6 nt (10,11), and two 5'-regions of a 24 nt primer influenced yeast telomerase activity (12). Sequences at the 5'-end of the primer also affected the processivity of endogenous human and Tetrahymena telomerase (13,14). These observations led to the proposal that telomerase contains a DNA-binding site outside the template region, called the ‘anchor site’ (13–15). It was proposed that the anchor site is necessary for repeat addition processivity, by allowing the enzyme to remain bound to the 5'-end of its DNA substrate during translocation. Thus, in order to fully understand the unique mechanism of telomerase, it is necessary to know more about the nature of the anchor site.

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The N-terminal 200 amino acids of TERT [known as region I (16) or the GQ (17), RID1 (18) or TEN (19) domain] have been proposed as a key contributor to the telomerase anchor site. This region of TERT has the ability to form a soluble, protease-resistant, independently-folded structure (17,19–21). Complementation studies with fragments of human TERT indicate that this region (together with an adjacent linker domain) can interact in trans with the rest of the TERT protein (18,22–24). Activity-based assays have provided support for anchor site mutations (25–27). Direct evidence for the involvement of this region in DNA binding was provided by the demonstration that mutations in Tetrahymena TEN led to decreased crosslinking to a primer containing 5-iodo-deoxyuridine (5-iodo-dU) substitutions at its 5'-end (19), and the site of crosslinking to a short primer was mapped to a particular amino acid within TEN (W187) (28). Direct binding assays (using capture with a biotinylated primer or gel shift analysis) have demonstrated that a fragment of human TERT encompassing TEN can bind telomeric DNA primers (24,29,30).

The exact role of the TEN domain in DNA binding, however, remains uncertain. Its affinity for DNA appears to be relatively low compared to other regions of TERT (11); a recent study using a bacterially-expressed TEN fragment of hTERT in gel shift analyses was unable to produce enough protein to quantitatively measure DNA affinity (24), implying a low-affinity interaction. Also, amino acids in TEN critical for repeat addition processivity or activity (L14 in Tetrahymena TERT and Q169 in hTERT, respectively) have been shown to not affect affinity with DNA of the same length as used in the activity assays (24,31). However, the amino acid corresponding to Q169 in Tetrahymena TERT (Q168) is involved in crosslinking to DNA (19); it is not clear if this discrepancy is due to a species difference or a difference in assays used to measure binding.

Given this uncertainty, we set out to quantitatively examine the effect on DNA binding of mutations in the hTERT TEN domain. We also examined the same mutations using activity assays, carrying out a detailed analysis of telomerase kinetic parameters in order to define the reason for reduced activity. Our data suggest that amino acids within the TEN domain are involved in positioning the 3'-end of the primer in the active site during nucleotide addition, a function which is distinct from the role of the TEN domain in providing DNA binding affinity.

**MATERIALS AND METHODS**

**Plasmids and oligonucleotides**

A panel of hTERT mutants with substitutions of the sequence NAAIRS and an N-terminal FLAG tag in the retroviral vector pBabehygro was kindly provided by Dr Christopher Counter, Duke University Medical Centre, Durham, NC (32). To create plasmids for *in vitro* expression, selected mutants were digested with EcoRI/SalI and the FLAG-hTERT fragment was sub-cloned into EcoRI/SalI sites of the pET-28 plasmid (Invitrogen). After carrying out many of the experiments in this study, we discovered two variations in the hTERT sequence in the NAAIRS panel compared to the hTERT sequence in Genbank entry AF015950 (33–35). These hTERT sequence changes (S229T and D516G) were corrected in the wild-type (wt) and mutant +170 expression constructs using the QuikChange II XL site-directed mutagenesis kit (Stratagene). There was no significant difference between these sequence variants in any of the *in vitro* kinetic parameters measured in this study (Supplementary Table S1 and Figure S2).

To construct a plasmid for *in vitro* transcription of hTR, the hTR gene [nucleotides 1–592] was amplified by PCR (36) and inserted into the vector pGEM-T (Promega). The insert of this plasmid was reamplified using primers that covered the 451 nt of hTR and included a T7 promoter (5'-end, for transcription) and a FokI restriction sequence (3'-end, for plasmid linearization) and inserted into the EcoRI/BamHI sites of plasmid pUC19. Linearization of the plasmid with FokI led to decreased crosslinking to DNA (19); it is not clear if this region (together with an adjacent linker domain) can interact in trans with the rest of the TERT protein (18,22–24). Activity-based assays have provided support for anchor site mutations (25–27). Direct evidence for the involvement of this region in DNA binding was provided by the demonstration that mutations in Tetrahymena TEN led to decreased crosslinking to a primer containing 5-iodo-deoxyuridine (5-iodo-dU) substitutions at its 5'-end (19), and the site of crosslinking to a short primer was mapped to a particular amino acid within TEN (W187) (28). Direct binding assays (using capture with a biotinylated primer or gel shift analysis) have demonstrated that a fragment of human TERT encompassing TEN can bind telomeric DNA primers (24,29,30).

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**In vitro translation and reconstitution of human telomerase**

Human telomerase RNA was transcribed *in vitro* from the plasmid pUC-hTR as described earlier (37). The RNA was gel purified and quantitated by absorbance at 260 nm before use in *in vitro* translation reactions using the FLAG-hTERT-pET-28 plasmids described above. Translations were carried out using the TnT Quick for PCR rabbit reticulocyte lysate system (Promega). Recombinant enzyme reconstitution reactions (400 μl) contained 320 μl TnT Quick mastermix, 8 μg of plasmid encoding the protein, 32 μl of [35S]methionine (Perkin-Elmer; 1175 Ci/mmol, 10 μCi/μl) and 150 nM telomerase RNA and were incubated at 30°C for 1 h. Reconstituted enzyme was immunopurified with anti-FLAG M2 affinity agarose beads (Sigma-Aldrich) in Protein LoBind tubes (Eppendorf). Beads (200 μl 1:1 slurry) were initially washed four times in 880 μl Tris–IP buffer [10 mM Tris–Cl, pH 7.5, 100 mM KCl, 1 mM MgCl2, 10% glycerol and 0.1 mM dithiothreitol (DTT)], centrifuging at 1000 g for 2 min at 4°C between washes. The translation reaction was diluted 1:10 in Tris–IP buffer. The washed anti-FLAG beads were added and the sample rotated at 12 rpm at 4°C for 2 h. The beads were washed four times in 4 ml of hTel buffer (50 mM Tris–Cl, pH 8.0, 1 mM MgCl2, 100 mM KCl, 5 mM DTT, 1 mM spermidine and 10% glycerol) before telomerase was eluted from the beads by competition with a 3× FLAG peptide (Sigma-Aldrich). The beads were resuspended in 120 μl of an elution mix (0.75 mg/ml FLAG peptide, 0.5 mg/ml BSA in hTel buffer) and rotated at 4°C for 1 h, centrifuged at 1000 g for 2 min at 4°C and the eluate removed to a fresh LoBind tube.
Table 1. Oligonucleotides used in this study

| Name   | Sequence                              |
|--------|---------------------------------------|
| 18GGG  | TTAGGGTTAGGGTTAGGG                   |
| 18GGT  | TAGGGTTAGGGTTAGGG                   |
| 18GTT  | AAGGGTTAGGGTTAGGG                   |
| 18TGA  | GGGTTAGGGTTAGGG                    |
| 18AGG  | GTAGGGTTAGGGTTAGG                   |
| 18ACT  | GTTAGGGTTAGGGTTAGG                  |
| 9GGG   | GGTAGGG                               |
| 9TTA   | GGTAGG                                |
| 5GTT   | AGGTG                                 |
| Bio-18GGG | Biotin-CTAGACCTGTCACTATTAGGGTTAGG   |
| Bio-18TAG | Biotin-CTAGACCTGTCACTAGGTAGTAG       |
| Bio-12GGG | Biotin-CTAGACCTGTCACTATTAGGGTTAGG   |
| Bio-PBR | Biotin-AGCCACTATCGACTAGCGCATCAT     |
| Telo4  | CGGTGGAAGGCGGCAGGCGAGGC             |

All sequences are listed 5′→3′.

The yield of active enzyme was determined by dot blotting using a probe against hTR. Aliquots of the immunopurified enzyme in formamide loading buffer (90% deionized formamide, 0.1% bromophenol blue, 0.1% xylene cyanol, in 1× TBE (89 mM Tris, 89 mM Boric acid, 2 mM EDTA)) were heated to 70°C for 10 min and chilled on ice for 2 min. The samples were dotted onto Hybond N+ membrane (GE Healthcare). The membrane was air dried for 1 h at RT, cross-linked for 10 min and chilled on ice for 2 min. The samples were transferred into Illustra Microprep columns (GE Healthcare) attached to a vacuum manifold and were eluted with 50 mM Tris–Cl, pH 8.0, 1 mM MgCl₂, 10% glycerol, 0.5 mM dTTP, 0.5 mM dATP, 4.6 μM nonradioactive dGTP and 0.33 μM [α-32P]dGTP at 20 μCi/ml, 6000 Ci/mmol (PerkinElmer Life Sciences). The indicated DNA primers were added to the plugged column and rotated at RT for 1 h. The microprep column was unplugged and centrifuged at 1000g for 1 min at RT to collect the enzyme into a Protein LoBind tube (Eppendorf). Purified enzyme was snap frozen in liquid nitrogen and stored at −80°C. The yield of active enzyme was determined by dot blotting using a probe against hTR, as described above.

**Immunopurification of overexpressed or endogenous telomerase**

Lysates of telomerase-transfected WI-38 VA13/2RA or 293T cells (10^7 cells in 1 ml; described above) or untransfected 293 cells (5×10^7 cells in 1 ml) were incubated with 20 μg sheep polyclonal anti-hTERT antibody (40) for 1 h at 4°C. Protein G agarose beads (50 μl) were pre-blocked in buffer A+BSA (0.5 mg/ml) for 30 min at RT, added to the sample and rotated for 1 h at 4°C. The remaining steps of the protocol were conducted in a 4°C coldroom with ice-cold buffers. Samples were transferred into Illustra Microprep columns (GE Healthcare) attached to a vacuum manifold and were washed with 10 ml of buffer A followed by 5 ml of hTel buffer (50 mM Tris–Cl, pH 8.0, 1 mM MgCl₂, 100 mM KCl, 5 mM DTT, 1 mM spermidine, 10% glycerol). To elute the purified protein, 300 μl of hTel buffer containing 50 μM peptide antigen (40) was added to the plugged column and rotated at RT for 1 h. The microprep column was unplugged and centrifuged at 1000g for 1 min at RT to collect the enzyme into a Protein LoBind tube (Eppendorf). Purified enzyme was snap frozen in liquid nitrogen and stored at −80°C.

Telomerase activity assays

Telomerase activity was measured by incubating 10 μl of immunopurified telomerase (recombinant, endogenous or overexpressed) in a 20 μl reaction including 50 mM Tris–Cl, pH 8.0, 1 mM MgCl₂, 50 mM KCl, 5 mM DTT, 1 mM spermidine, 10% glycerol, 0.5 mM dTTP, 0.5 mM dATP, 4.6 μM nonradioactive dGTP and 0.33 μM [α-32P]dGTP at 20 μCi/ml, 6000 Ci/mmol (PerkinElmer Life Sciences). The indicated DNA primers were added to the reactions at concentrations of 10 nM to 5 μM. Standard primer concentrations were 1–2 μM for 18-nt primers, 5 μM for 9-nt primers and 50 μM for 5 nt primers. The reactions were incubated at 30°C for 1–6 h; the linear phase of enzyme activity extends for ~10h (data not shown). Reactions were terminated by addition of 100 μl of TES buffer (50 mM Tris–Cl, pH 8.3, 20 mM EDTA, 0.2% SDS; 1000 cpm of a [32P] end-labelled 100-nt oligonucleotide (Sigma) and grown in a 37°C humidified incubator containing 5% CO₂.

WI-38 VA13/2RA and 293T cells were co-transfected with plasmids encoding hTERT under a CMV promoter (38) and hTR under a U3 promoter (39) using Fugene-6 transfection reagent (Roche) and incubated at 37°C for 6 h. After 72 h at 32°C the cells were harvested and lysed in 1 ml of buffer A per 10^7 cells (20 mM HEPEs-KOH, pH 7.9, 300 mM KCl, 2 mM MgCl₂, 10% glycerol, 1 mM EDTA, 0.1% Triton X-100, 1 mM DTT and 1 mM PMSF). The cell suspensions were rotated at 4°C for 1 h and centrifuged at 18,000g at 4°C for 20 min to clarify the lysate. The supernatant was transferred to a Protein LoBind microfuge tube (Eppendorf), snap frozen in liquid nitrogen and stored at −80°C.

Cell culture and plasmid transfection

WI-38 VA13/2RA human foetal fibroblasts (ATCC) and human embryonic kidney (HEK) cell line 293T were cultured in Dulbecco’s Modified Eagles Medium (Invitrogen) supplemented with 10% foetal calf serum (Hyclone) and grown in a 37°C humidified incubator containing 5% CO₂.

WI-38 VA13/2RA and 293T cells were co-transfected with plasmids encoding hTERT under a CMV promoter (38) and hTR under a U3 promoter (39) using Fugene-6 transfection reagent (Roche) and incubated at 37°C for 6 h. After 72 h at 32°C the cells were harvested and lysed in 1 ml of buffer A per 10^7 cells (20 mM HEPEs-KOH, pH 7.9, 300 mM KCl, 2 mM MgCl₂, 10% glycerol, 1 mM EDTA, 0.1% Triton X-100, 1 mM DTT and 1 mM PMSF). The cell suspensions were rotated at 4°C for 1 h and centrifuged at 18,000g at 4°C for 20 min to clarify the lysate. The supernatant was transferred to a Protein LoBind microfuge tube (Eppendorf), snap frozen in liquid nitrogen and stored at −80°C.
was added as a recovery and loading control. The reaction products were phenol/chloroform extracted and ethanol precipitated. The pellet was resuspended in 5 μl of TE buffer and 5 μl of formamide loading buffer (90% deionised formamide, 0.1% bromophenol blue, 0.1% xylene cyanol in 1× TBE). Immediately prior to gel loading, samples were heated to 70°C for 5 min and chilled on ice for 2 min before centrifuging at 16 000 g at 4°C for 2 min. Half the reaction was electrophoresed on a denaturing 10% polyacrylamide/8 M urea sequencing gel for 2 h at 80 W. The gel was dried at 80°C for 30 min, exposed to a phosphorimager screen, scanned with a Typhoon 9410 Workstation and quantitated with ImageQuant software.

For experiments using a single nucleotide, the concentrations of nucleotide were 47.5 μM non-radioactive dTTP and 2.5μM [α-32P] dTTP at 10 mCi/ml, 800 Ci/mmol (for ‘T-only’ assays), 47.5 μM non-radioactive dATP and 2.5μM [α-32P] dATP at 10 mCi/ml, 800 Ci/mmol (for ‘A-only’ assays) and 10 μM non-radioactive dGTP and 0.66 μM [α-32P] dGTP at 20 mCi/ml, 6000 Ci/mmol (for ‘G-only’ reactions). These concentrations are above the approximate K_m (for ‘G-only’ reactions). These concentrations are above K_m for dTTP, 0.66 M for dGTP and 10 M for radioactive dGTP. Experiments measuring the affinity of dGTP included 0.5 M dTTP, 0.5 mM dATP and the indicated concentration of dGTP at a constant specific activity of 200 Ci/mmol.

The intensity of extension products at each substrate concentration was summed and normalized against the intensity of the 32P-labelled 100-nt oligonucleotide loading control to give relative activities; those in Figure 1c were carried out at 1 μM substrate. These data were then expressed as a percentage of the reaction with maximal activity and plotted against oligonucleotide (substrate) concentration [S]. The curve was fitted to the equation \( y = \frac{y_{\text{max}} \times [S]}{K_m + [S]} \), to yield the Michaelis–Menten affinity constant K_m. To calculate relative processivity, the intensity of the first 3–5 repeats was adjusted for specific activity (i.e. number of incorporated radiolabelled nucleotides per repeat) and plotted versus repeat number on a log-linear graph. Processivity was defined as −0.693/s, where s is the slope of the resulting straight line.

For calculating k_obs, gels such as that in Figure 6b included a standard curve of known amounts (cpm) of a 32P-labelled 100-nt oligonucleotide. The cpm of each telomerase product band were converted to Ci (since 1 cpm = 2.2 × 10^{-12} Ci for 32P), then to fmol based on the specific activity of the radiolabelled nucleotides, and normalized to the fmol of enzyme used in the assay to give k_obs.

DNA binding assay to measure primer binding to hTERT

Ultralink Immobilized NeutrAvidin Protein Plus beads (Thermo Scientific) (700 μl) were washed four times in 1 ml wash buffer 75 (50 mM Tris–Cl, pH 8.0, 1 mM MgCl_2, 10% glycerol, 75 mM KCl, 5 mM DTT, 1 mM spermidine), centrifuging at 1000 g for 1 min at 4°C between washes. The beads were incubated twice with 1 ml of blocking buffer (50 mM Tris–Cl, pH 8.0, 1 mM MgCl_2, 10% glycerol, 75 mM KCl, 5 mM DTT, 1 mM spermidine, 0.75 mg/ml BSA, 0.15 mg/ml glycogen) for 30 min at RT with agitation and resuspended in 350 μl blocking buffer (for 293T telomerase) or hTel buffer (for recombinant telomerase). Immunopurified overexpressed telomerase from 293T cells (50 μl) or immunopurified recombinant telomerase (10 μl) was added to a 100 μl reaction including hTel buffer (50 mM Tris–Cl, pH 8.0, 1 mM MgCl_2, 50 mM KCl, 5 mM DTT, 1 mM spermidine, 10% glycerol) and oligonucleotide at concentrations ranging from 0 to 10 μM. Similar results were obtained with telomerase expressed in 293T cells or WI-38 VA13/2RA cells (data not shown). The reactions were incubated at 30°C for 30 min (293T telomerase) or 60 min (recombinant telomerase) followed by the addition of 50 μl of blocked NeutrAvidin beads and agitation at 4°C for 2 h. The samples were washed three times in 300 μl of wash buffer 50 (50 mM Tris–Cl, pH 8.0, 1 mM MgCl_2, 10% glycerol, 50 mM KCl, 5 mM DTT, 1 mM spermidine) with centrifugation at 1000 g for 2 min at 4°C between washes, before the bead pellets were resuspended in 100 μl of formamide loading buffer (see above) containing 0.5 μM biotin. Immediately prior to dot blotting, the samples were heated to 70°C for 10 min, chilled on ice for 2 min and centrifuged at 14 000 g for 1 min at 4°C. Half of the reaction was dotted onto Hybond N+ membrane alongside in vitro transcribed hTR standards loaded in an equal volume of formamide loading buffer. The membrane was probed with 32P end-labelled Tel4 probe against hTR, as described above. The blot was exposed to a phosphorimager screen for 72 h and analysed using ImageQuant TL software. The signal from the sample blank (containing no oligonucleotide) representing background binding to the beads was subtracted from all others. The resulting value was expressed as a percentage of the maximal intensity at the highest primer concentration and plotted against oligonucleotide (substrate) concentration [S].

The curve was fitted to the following equation: \( y = \frac{(B_{\text{max}} \times [S])/(K_d + [S])}{y_{\text{max}} \times [S]} \), where B_{max} is the maximal level of binding, to yield the dissociation constant K_d.

**Glycerol density gradient centrifugation and fractionation**

Glycerol gradient stock solutions containing either 10 or 40% glycerol in 300 mM KCl, 2 mM MgCl_2, 0.1% Triton X-100, 1 mM DTT and 20 mM HEPES-KOH pH 7.9 were combined using a two-chamber gradient mixer to pour 11 ml of of 10–40% linear glycerol density gradients in Ultra-Clear 13.2 ml centrifuge tubes (14 x 89 mm; Beckman Coulter). Recombinant wild type or +170 mutant telomerase (500 μl), synthesized in rabbit reticulocyte lysate as described above, was applied to the top of the gradient. Centrifugation was performed with a SW-41 Ti swinging-bucket rotor (Beckmann Coulter) at 230000 g at 4°C for 24 h. Following centrifugation, glycerol gradients were collected by puncturing the base of each tube with a 30-gauge needle; 23 0.5 ml fractions were collected by gravity elution and stored at −80°C. Telomerase activity was measured using 27 μl of each glycerol
gradient fraction in a 40 μl reaction using 1 μM of primer Bio-18GGG, as described above.

**Northern blotting**

Purified telomerase (from 5 × 10⁶ telomerase-transfected WI-38 VA13/2RA cells) in 300 μl buffer A was digested with 33 μl Proteinase K buffer (100 mM Tris–Cl, pH 7.5, 75 mM EDTA, 6% SDS) and 2 μl proteinase K (40 μg) for 2 h at 45°C. The samples were spiked with 5000 cpm of a 32P-labeled 100-nt oligonucleotide as a recovery and loading control, phenol/chloroform extracted and ethanol precipitated. The pellet was resuspended in 10 μl of MilliQ water and 10 μl formamide loading buffer (90% deionized formamide, 0.1% bromophenol blue, 0.1% xylene cyanol in 1 × TBE). Samples were heated at 70°C for 5 min, cooled on ice for 2 min and electrophoresed on a 1.5-mm-thick 4% polyacrylamide, 8 M urea, 1 × TBE gel for 1 h at 25 W in 1 × TBE buffer. The gel was transferred onto Hybond XL membrane (GE Healthcare) by electroblotting for 2 h at 1.5 A in 0.5 × TBE buffer cooled to 4°C with a recirculating water bath. The membrane was probed with 32P end-labeled Telo4 probe directed against hTR, as described above.

**Western blotting**

Purified telomerase (from 5 × 10⁶ telomerase-transfected WI-38 VA13/2RA or 293T cells) was added to an equal volume of 5x Laemmli’s sample buffer (250 mM Tris–Cl, pH 6.8, 10% SDS, 0.01% bromophenol blue, 50% glycerol and 1.8 M β-mercaptoethanol) and heated to 95°C for 5 min. The samples were electrophoresed on an 8% polyacrylamide/SDS-PAGE gel in Laemmli’s running buffer (25 mM Tris, 0.1% SDS, 192 mM glycine) for 2 h at 100 V and transferred to Hybond enhanced chemiluminescent (ECL) nitrocellulose (GE healthcare) by electroblotting for 2 h at 1.5 A in 0.5 × TBE buffer cooled to 4°C with a recirculating water bath. The membrane was washed for 1 h at RT in phosphate buffered saline (PBS) before incubation with the secondary antibody (rabbit anti-goat immunoglobulin horseradish peroxidase; DAKO) diluted 1:10 000 in 1% PBS/0.05% Tween-20. The membrane was rinsed with PBS before incubation with Amersham ECL plus Western blotting detection reagent (GE Healthcare) for 5 min at RT. The blot was exposed to a LASS-4000 Luminescent Image Analyser (FujiFilm) and quantitated using Multigauge software.

**Protein homology modelling and nucleic acid docking**

An homology model of the human hTERT TEN domain was determined by threading the TEN domain sequence of hTERT onto chain A of the *Tetrahymena* TEN domain crystal structure [PDB id:2B2A (19)] using the Bioinformatics Online Toolkit (http://toolkit.tuebingen.mpg.de). A 3D homology model was then constructed with MODELLER (http://www.salilab.org/modeller) and the top-ranked model minimized using the Tripos force field until a gradient consensus was reached (21 003 iterations, energy = −615.496 kcal/mol) followed by minimization using the Kollman-all atom force field, with the addition of AMBER charges until a gradient consensus was reached (16 957 iterations, energy = −2648.889 kcal/mol). The stereochemical quality of the model was checked with PROCHECK V3.5, with 96.9% of residues in an acceptable region of a Ramachandran plot and all other stereochemical statistics deemed reasonable (41).

The DNA primer 5′-TTA-3′ was constructed in Sybyl-X-1.1 using the Biopolymer module (http://www.tiros.com). The DNA single strand was docked into the TEN domain model using the HEX program ([42] and http://www.csd.abdn.ac.uk/hex/). A shape and electrostatics correlation was used in the 3D Fastlite mode with a grid dimension of 0.75 Å, receptor and ligand range 180°, twist range 360°, distance range 40 Å and a scan step of 0.75. Two thousand solutions were requested in the initial dock with 20 clusters of solutions found after post-minimization. The complementary RNA base pairs (5′-UAA-3′) were added to the lowest energy dock and visually analyzed.

**RESULTS**

A mutant of the hTERT TEN domain with reduced activity and processivity, and dramatically increased primer $K_m$

To quantitate the contribution of human TEN to telomerase activity and DNA affinity, we performed *in vitro* assays with a panel of hTERT mutants with six amino acid substitutions of the sequence NAAIRS (32,43). This substitution is predicted to have minimal effect on protein structure (44). Telomerase was reconstituted in rabbit reticulocyte lysates (RRL) using hTERT substituted with NAAIRS at one of fifteen positions throughout the protein (Figure 1a), and activity measured with a direct (i.e. non-PCR) assay using oligonucleotide 18GGG [(TTA GGG)]₃ as a substrate (Figure 1b; see Table 1 for oligonucleotide sequences). Relative activity levels were quantitated based on total product intensity, and repeat addition processivity (RAP) was determined from the rate of decrease of product intensity over the first three to five repeats (Figure 1c). As a first step towards examining DNA affinity, the $K_m$ for primer 18GGG was measured by quantitating product intensity at different concentrations of this primer (Figure 1c).

Activity levels of most NAAIRS mutants reflected those measured using *in vivo* reconstituted telomerase (32,43). One exception is the mutation at amino acids 8–13 (here-after called +8 in line with the previous nomenclature), which was substantially more active when reconstituted *in vitro* than *in vivo*. This may be explained by the observation that the extreme N-terminus of hTERT is involved in its nucleolar localization *in vivo* (45).
A subregion of the TEN domain has been shown to be necessary for telomere extension in vivo but not telomerase activity, and has been called the dissociates activities of telomerase (DAT) domain (32). Mutations in the DAT domain (Figure 1, purple) had modest effects on activity and no reduction in RAP. Regions on either side of DAT (regions 1A and 1B) were much more important for activity. The 1B mutant (+170) had substantially reduced RAP, while the 1A mutants (+8, +26) had wt RAP. A mutation in the non-conserved linker region (+200) had no effect on activity or RAP, while the conserved motifs QFP and E [involved in RNA and DNA binding respectively (18,29)] were essential for activity. We confirmed that mutations in the hTERT C-terminus reduced both activity and RAP, as previously shown (43,46).

Analysis of the \( K_m \) for primer 18GGG gave unexpected results. Mutants +26, +74 and +92 showed very modest (~2-fold) increases in \( K_m \), while the \( K_m \) of most other mutants did not differ from wt. The one exception was mutant +170 in region 1B, which had a dramatic (~14-fold) increase in \( K_m \) (Figure 1c, Table 2).

Table 2. \( K_m \) of various DNA primers with wt recombinant hTERT and mutant +170, and the fold difference between these numbers

| Primer  | Sequence                  | \( K_m \) (nM) | Fold difference wt |
|---------|---------------------------|----------------|-------------------|
|         | wt                        | +170           |                   |
| 18GGG   | (TTAGGG)\(_n\)           | 49 ± 29        | 710 ± 140         | 7.8 ± 0.3 |
| 18TTA   | (GGGTTA)\(_n\)           | 65 ± 4         | 1530 ± 100        | 24       | 9.7 ± 5.6 |
| 9GGG    | GGGTTAGGG\(_n\)          | 56 ± 16        | 300 ± 20          | 5        | n.d.     |

The right column lists the \( K_m \) of some of the same primers for endogenous wt human telomerase. Results are the mean ± standard deviation of two to six experiments.

A subregion of the TEN domain has been shown to be necessary for telomere extension in vivo but not telomerase activity, and has been called the dissociates activities of telomerase (DAT) domain (32). Mutations in the DAT domain (Figure 1, purple) had modest effects on activity and no reduction in RAP. Regions on either side of DAT (regions 1A and 1B) were much more important for activity. The 1B mutant (+170) had substantially reduced RAP, while the 1A mutants (+8, +26) had wt RAP. A mutation in the non-conserved linker region (+200) had no effect on activity or RAP, while the conserved motifs QFP and E [involved in RNA and DNA binding respectively (18,29)] were essential for activity. We confirmed that mutations in the hTERT C-terminus reduced both activity and RAP, as previously shown (43,46).

Analysis of the \( K_m \) for primer 18GGG gave unexpected results. Mutants +26, +74 and +92 showed very modest (~2-fold) increases in \( K_m \), while the \( K_m \) of most other mutants did not differ from wt. The one exception was mutant +170 in region 1B, which had a dramatic (~14-fold) increase in \( K_m \) (Figure 1c, Table 2).

Primers with different permutations of the 3’-end have up to 100-fold differences in dissociation rate from endogenous human telomerase (47), accounting for the distinctive 6 nt pausing pattern of telomerase extension. Since the pausing pattern of recombinant human telomerase is
very similar to that of endogenous telomerase purified from 293 cell extracts (Supplementary Figure S1), it is highly likely that recombinant telomerase shows the same pattern of dissociation at different template positions. We therefore also measured the Kₘ of the primer 18TTA ([GGGTTA]₃; see Figure 6a for an illustration of the alignment of this primer with the RNA template relative to that of primer 18GGG. Mutant +170 showed an even greater increase in Kₘ with primer 18TTA (Table 2 and Supplementary Figure S1). For wt telomerase the Kₘ of primers 18GGG and 18TTA were not substantially different (Table 2); since these primers are predicted to have a ~100-fold difference in dissociation rate, this implies that Kₘ may not be a good reflection of primer affinity (Kₐ) for telomerase. This is also the case for endogenous human telomerase from 293 cells; the Kₘ of 18GGG and 18TTA were lower for endogenous telomerase than recombinant, but were similar to each other (Table 2). We therefore embarked on an exploration of the kinetic parameters of mutant +170 telomerase in order to identify the defect(s) responsible for its dramatic Kₘ increase.

During the course of this study, we detected two additional amino acid mutations in the panel of NAAIRS mutants used (32) relative to a wt hTERT sequence in Genbank (33). We corrected these amino acids and confirmed that they have no effect on telomerase activity or on any of the kinetic parameters measured in this study (Supplementary Figure S2 and Table S1).

**Table 3. Kₐ of various DNA primers with wt recombinant and overexpressed hTERT and mutant +170, and the fold difference between these numbers**

| Primer       | Recombinant | Overexpressed in 293T |
|--------------|-------------|-----------------------|
|              | Kₐ (nM)     | Kₐ (nM)               | Fold difference |
|              | wt +170     | wt +170               |                |
| Bio-18GGG    | 164 ± 63    | 361 ± 142             | 2.2            |
| Bio-18TAG    | n.d.        | n.d.                  |                |
| Bio-12GGG    | n.d.        | n.d.                  |                |
| Bio-18TAG    | 0.5 ± 0.3   | 600 ± 70              | 1200           |
| Bio-12GGG    | 0.9 ± 0.4   | 510 ± 90              | 567            |
| Bio-18GGG    | 0.5 ± 0.1   | 1590 ± 770            | 3180           |

Results are the mean ± standard deviation of three to four experiments. n.d., not determined.

**Mutant +170 has a defect in DNA binding affinity**

Since the Kₘ for a primer-enzyme interaction (measured by activity) is not necessarily equal to the affinity constant (Kₐ) for the same interaction, we next measured primer affinity directly using a pulldown assay that we had developed for *Tetrahymena* telomerase (11). Biotinylated 18-nt telomeric primers were incubated with purified recombinant enzyme and recovered with Neutravidin beads, and the yield of telomerase measured by dot blot against hTR (Supplementary Figure S3). The Kₐ of primer binding to mutant +170 was only 2-fold greater than that of wt telomerase (Table 3), confirming that Kₘ is not a good measure of Kₐ for recombinant human telomerase.

Since endogenous human telomerase exhibits a lower Kₘ than recombinant enzyme (Table 2), we also measured direct primer binding affinities for endogenously-assembled human telomerase. We reconstituted wt and mutant +170 telomerase *in vitro* in 293T cells, purified it by immunopurification with an hTERT antibody (40) and employed the same direct primer binding assay. Under our transfection conditions in 293T cells, overexpressed telomerase activity exceeds the endogenous levels by ~200-fold, resulting in <1% contribution by the endogenous wt enzyme. To our surprise, the Kₛₐ of primer binding of 18-nt primers to overexpressed cellular mutant +170 were 500–1200-fold greater than those of wt telomerase (Figure 2, Table 3). This indicates that recombinant human telomerase does not exactly recapitulate the DNA binding properties of the endogenously-assembled enzyme, and this discrepancy masks the dramatic defect in DNA binding affinity of mutant +170.
hTERT mutant +170 retains wt global structure

One potential explanation for the large effect of mutation +170 on $K_m$, $K_d$ and processivity is an effect on enzyme structure. To rule this out, we fractioned wt and mutant +170 telomerase (reconstituted in RRL) over parallel 10–40% glycerol gradients. The use of crude translation lysate resulted in products from contaminating polymerases and nucleases, some even more abundant than the extension products from mutant +170 (Figure 3a). Nonetheless, telomerase extension products from both enzymes were detected (Figure 3a, n + 4 bands) and eluted with almost identical profiles (Figure 3a, right panel), indicating a similar overall composition and structure.

Another indicator of wt conformation is the ability to bind to hTR. This was measured by immunoprecipitation of in vitro translated hTERT with an antibody to an N-terminal tag, followed by dot blotting to quantify yield of bound hTR (Figure 3b, top panel). When normalized to the yield of [35S]-labelled hTERT (Figure 3b, bottom panel), mutant +170 bound hTR at least as efficiently as wt hTERT (levels of 130 ± 10%). To measure the ability of mutant +170 hTERT to assemble with hTR within human cells, we also reconstituted wt and mutant telomerase in telomerase-negative WI38 VA13/2RA cells (which lack endogenous expression of hTERT and hTR). The amount of hTR that immunoprecipitated with mutant hTERT was equal to that with wt hTERT (Figure 3c; levels of 95 ± 19%), demonstrating that in vivo assembly of the telomerase complex was intact. Assembly of mutant +170 hTERT with hTR was slightly less efficient in 293T cells, at 63% of wt levels (Figure 3d).

Increased $K_m$ of mutant +170 is not attributable to a defect in translocation rate or interaction with 5'-end of primer

It has been proposed that the effect of the TEN domain on RAP is due to a defect in the telomerase translocation step (31). It is difficult to directly measure translocation rate with a mutant that shows almost no detectable second repeat addition, so we instead examined the effect of mutant +170 on translocation indirectly. In vitro reconstituted telomerase was used to extend primer 18GGG in a reaction with $^{32}$P-dTTP as the only nucleotide. This reaction results in the addition of two
Increased \( K_m \) of mutant +170 is not attributable to a defect in translocation rate or interaction with 5'end of primer. (a) Direct telomerase activity assay with wt or mutant recombinant telomerase using a titration of primer 18GGG, in the presence of \(^32\)P-dTTP and no other nucleotide, in order to measure activity in the absence of translocation. LC: labelled 100 nt oligonucleotide as a recovery and loading control; +1: primer plus 1 nucleotide. (b) Activity assay in the presence or absence of primer 5TTA (5'-GGTAA-3', 50 \( \mu \)M), with \(^32\)P-dGTP alone (right four lanes) or \(^3\)P-dGTP, dATP and dTTP (left four lanes), using wt or mutant +170 (m) recombinant telomerase. (c) Activity assay using a titration of primer 5GGT and recombinant telomerase. LC: labelled 100 nt oligonucleotide as a recovery and loading control; +3: primer plus 3 nt.

hTERT mutant +170 is defective in nucleotide polymerization, particularly opposite position 51 of the template

Since \( K_m \) is a measure of total enzyme activity, it is likely that many kinetic parameters contribute to this value. For example, since DNA dissociation rate varies at each template position (47), it is possible that polymerization rate \( k_{\text{pol}} \) also varies across the telomerase template; this has not previously been tested. We therefore propose the telomerase kinetic scheme in Figure 5, illustrated for one round of addition of a TTAGGG repeat. The scheme includes rate constants for association and dissociation of primers of six permutations (e.g. \( k_{\text{off}}(G\hat{G}G) \) and \( k_{\text{off}}(G\hat{G}\hat{G}) \)); polymerization at the six positions (e.g. \( k_{\text{pol}}(G\hat{G}G) \)); and translocation \( k_{\text{trans}} \). The 21 rate constants shown here, plus nucleotide association constants and possible protein conformational changes that are not shown, contribute to the overall \( K_m \) for processive extension of a primer. In non-processive reactions including a single dNTP, such as that in Figure 4a, the \( K_m \) would represent a portion of this scheme, illustrated by the dashed box in Figure 5 for primer 18GGG.
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Figure 5. Proposed kinetic scheme for human telomerase. (a) Alignment of an elongated 18 nt DNA primer (black) with the RNA template (red) before (top) and after (bottom) translocation and addition of two Gs. Numbers 1–6 are assigned to each position of nucleotide addition within a single telomeric repeat. (b) Proposed kinetic scheme for telomerase. E = telomerase, S = DNA substrate, with 3'-permutation of each DNA primer in parenthesis. ES = telomerase-DNA complex. It has been demonstrated that the dissociation rate \( k_{\text{off}} \) of a particular DNA sequence is the same whether that DNA is the substrate or product of extension (47), so only one \( k_{\text{off}} \) has been included for each sequence. Note that the rate constant for translocation \( k_{\text{trans}} \) is potentially composed of at least two other rate constants, for RNA-DNA dissociation and realignment, respectively (50). Protein conformational changes and nucleotide association constants are not shown in this scheme. The dashed box indicates the portion of this scheme measured in a reaction with 18GGG as the DNA primer and dTTP as the only nucleotide. since we cannot distinguish between measurement of \( k_{\text{pol}} \) and rates of protein conformational change (since we do not know which is rate limiting), we have termed the resulting rate constant \( k_{\text{obs}} \) (Table 4). The \( k_{\text{obs}} \) at each position is similar for recombinant telomerase and telomerase overexpressed in 293T cells; note that the units are \( \text{h}^{-1} \), demonstrating that telomerase is a slow enzyme. For wt telomerase, \( k_{\text{obs}} \) varies by a factor of 10–20, with the slowest addition being at position 5 (opposite RNA nucleotide 51, see Figure 6a) and the fastest at position 1 (opposite RNA nucleotide 49). Note that these differences do not correlate with the ~100-fold differences in enzyme dissociation measured for these six primers (47), indicating that primer dissociation is not rate-limiting in this experiment.

Mutant +170 demonstrated a reduced \( k_{\text{obs}} \) at each template position (Figure 6b, Table 4). Most strikingly, there was a complete lack of detectable addition to primer 18TAG when using recombinant telomerase (Figure 6b, top panel). This reflects an inability to copy position 51 of the RNA template and thus an inability to add the second G in the repeat TTAGGG. This was confirmed by the pattern of addition to primer 18TTA; mutant +170 telomerase is able to add a single G to this primer, but not the second G. Telomerase reconstituted in 293T cells presumably retained more contaminating nucleases and polymerases after purification, since it produced more bands representing nuclease activity or non-templated primer elongation (Fig. 6b, lower panel, bands without asterisks), which complicates interpretation of these reactions. In particular, a band of slightly slower mobility is obscuring the extension products of primer 18TAG. Nevertheless, the trend for a reduction in \( k_{\text{obs}} \) for mutant +170 reflects that of recombinant telomerase, and this appears to be most dramatic with primer 18TAG.

Since primers 18TTA and 18TAG can each hybridize to two positions on the RNA template, it is theoretically possible that they preferentially hybridize at the template 5’-end. This is unlikely, since primer 18TAG displays a monophasic dissociation rate that reflects binding to the 3’-end of the template (47). If true, however, this would mean that an inability to add the second G could be due to a defect in translocation rather than a defect in addition opposite RNA position 51. To address this, we made use of a primer with 3 nt changed such that the 3’-end of the primer can only align at the 3’-end of the template (primer 18ACT; see Figure 6a, bottom). wt telomerase was able to processively extend this primer almost as well as primer 18TAG (Figure 6c). This indicates that human telomerase does have the ability to utilize the 3’-end of its RNA template, and thus an inability to extend primer 18TAG likely represents a polymerization defect rather than a translocation defect. As predicted, mutant +170 is unable to add nucleotides to primer 18ACT.

The rate constants representing nucleotide association are not shown in Figure 5 for simplicity, but may contribute to final \( K_m \) values. As a start towards measuring these affinities, we titrated dGTP in a non-processive reaction using primer 18TTA (Figure 6d). The \( K_m \) for dGTP of wt recombinant telomerase was 1.3 ± 0.4 \( \mu \text{M} \), while that of mutant +170 was ~3-fold higher at 4.2 ± 1.8 \( \mu \text{M} \).
Table 4. $k_{obs}$ using DNA primers with each of six repeat permutations, with wt recombinant and overexpressed hTERT and mutant +170

| Primer | Sequence | Recombinant $k_{obs}$ (h$^{-1}$) | Overexpressed in 293T $k_{obs}$ (h$^{-1}$) |
|--------|----------|---------------------------------|----------------------------------|
|        |          | wt +170 Fold defect      | wt +170 Fold defect   |
| 18GGG  | (TTAGGG)$_3$ | 2.7 ± 1.4 0.7 ± 0.5 4  | 2.9 ± 1.8 0.8 ± 0.5 4 |
| 18GTT  | (TAGGGT)$_3$ | 1.7 ± 0.7 0.5 ± 0.2 3  | 2.5 ± 0.8 0.4 ± 0.2 6 |
| 18GTT  | (AGGGTT)$_3$ | 1.3 ± 0.7 0.3 ± 0.2 4  | 1.8 ± 0.5 1.9 ± 1.1 1 |
| 18TTA  | (GGGTAT)$_3$ | 0.9 ± 0.2 0.09 ± 0.02 10 | 0.7 ± 0.2 0.2 ± 0.1 3 |
| 18TAG  | (GGTATG)$_3$ | 0.21 ± 0.03 <0.006 >35 | 0.15 ± 0.06 n.q. n.q. |
| 18AGG  | (GGTATG)$_3$ | 0.41 ± 0.06 0.06 ± 0.01 7 | 0.34 ± 0.09 0.15 ± 0.12 2 |
| Overall |          | 0.25 ± 0.09 0.029 ± 0.006 9 | n.d. n.d. n.d. |

The rates in the first six rows were calculated for reactions including a single radiolabelled nucleotide, i.e. non-processive reactions. The rate in the bottom row represents a processive reaction with the primer 18GGG. Results are the mean ± standard deviation of three experiments. n.d., not determined; n.q., not quantifiable due to contaminating products.
Therefore it is possible that this mutation affects the ability of nucleotides to enter the enzyme active site, possibly via an allosteric interaction of the TEN domain with the active site. However, this 3-fold defect in $K_m$ for dGTP is insufficient to explain the >35-fold defect in addition to primer 18TAG; furthermore, the latter reaction was carried out at a dGTP concentration of 10 $\mu M$. Thus the data in Figure 6b and Table 4 point to a defect of mutant +170 in $k_{pol}$ (particularly $k_{pol.5}$) and/or a defect in protein conformational changes during catalysis.

Homology model of the human TEN domain with a docked DNA primer

The crystal structure of the Tetrahymena TEN domain was used to perform homology modelling of the hTERT TEN domain using a threading technique (Figure 7b). Although of low sequence similarity, the essential features of the human TEN domain model and the Tetrahymena structure are very similar; in particular, the predicted DNA binding groove and the region mutated in this study are highly similar (Figure 7b, red and yellow, respectively). This model is also similar to a previously published homology model of the human TEN domain (48). The regions of most difference in all the models are the poorly conserved loops (Supplementary Figure S4).

The 3 nt DNA primer 5'-TTA-3' was docked into the TEN domain model using the HEX programme (42). The top-scoring solution was positioned with its 3'-end in the pocket formed by amino acids 170–175, and showed polar interactions with Val81, Asn95, Lys94, Tyr176 and Gln177 (Supplementary Figure S5 and Table S2). The DNA substrate is also positioned in a way that the further addition of bases to the 5'-end could lead to interactions with residue Pro188.

During extension, the 3'-end of the DNA primer is predicted to form a duplex with the RNA template. The addition of the complementary RNA base pairs to the docked DNA primer resulted in a steric clash between the RNA and a large loop in the TEN domain. However, it is plausible that this loop is sufficiently flexible to move out of the way, allowing interaction with an RNA–DNA duplex (Supplementary Figure S6).

DISCUSSION

Kinetic analyses of telomerase extension using the hTERT TEN domain mutant +170 has demonstrated that a major defect of this mutant is in nucleotide addition opposite position 51 of the RNA template. This mutant also exhibits a defect in affinity for an 18 nt DNA primer, corroborating previous data implicating the TEN domain as a contributor to DNA binding affinity. However, our results demonstrate that the defect in catalysis of mutant +170 is independent of its defect in DNA binding, since catalytic rate measurements were carried out at saturating DNA concentration. These observations expand the numerous functions of this important region of hTERT.

The observed catalytic defect may reflect a difference in the actual polymerization rate of telomerase at this position, or, since these amino acids are not part of the catalytic triad of aspartates, a protein conformational change during this step of catalysis. The latter conclusion is supported by the finding of a possible conformational change in the vicinity of the TEN domain in a mutant of the nearby amino acid Q169 (30), although it should be emphasized that we are proposing a defect in a conformational change during catalysis rather than in the steady-state conformation of the enzyme; the former would be considerably more difficult to measure.

Together with previous data demonstrating that amino acids in this vicinity of Tetrahymena TERT have the ability to crosslink to a DNA primer (19,28), our data suggest that this region of the TEN domain is involved in positioning the 3'-end of the primer in the active site (Figure 7a). Docking of the DNA primer in our human TEN homology model supports this proposition (Supplementary Figure S5). Conformational changes in the enzyme are undoubtedly occurring as the RNA–DNA duplex is realigned with the active site after addition of each dNTP, and we propose that the TEN domain plays a key role in mediating these. Amino acids 170–175 are particularly involved in the transition from –TAG to –TAGG; this transition may be the one requiring the greatest conformational change, or possibly the other five transitions are primarily mediated by other amino acids. The former possibility is supported by the finding that an amino acid in a completely different region of Tetrahymena TERT also leads to a defect in addition of the second G in the telomeric repeat (49). This is also the position of slowest addition (Table 4).

Mutant +170 affects nucleotide addition to some extent at every position of the RNA template. This indicates that these amino acids remain in the vicinity of the telomerase active site throughout the addition cycle. Furthermore, the DNA binding affinity provided by these amino acids is also localized to the 3'-end of the primer. We postulate that the two defects, while functionally separable, are related in that they both result from reduced binding of amino acids 170–175 to the 3'-end of the DNA. This region of the TEN domain is therefore not involved in the classical ‘anchor site’ at the 5'-end of the DNA primer. In Tetrahymena TERT, amino acid W187 remains close to DNA at the 3'-template boundary throughout the addition cycle (28). Combining these two observations results in a model whereby the distance between these amino acids increases over the addition cycle (Figure 7a). Since the predicted length of the template RNA–primer DNA duplex is 27 A˚, the distance between W187 and Q168 in Tetrahymena TERT is 24 A˚, and the corresponding distance in the human TEN domain is also 24 A˚, we propose that the structures in Figure 7b represent the ‘stretched’ state of the TEN domain just prior to translocation. Our model (illustrated in Figure 7a) is a refinement of previous models for the function of the TEN domain in that we propose: (i) the conformational change may be within the TEN domain itself, rather than between the TEN domain and the rest of the protein (28), and (ii) the conformational change is one
The extent of the DNA affinity defect of overexpressed cellular mutant +170 (500–1000-fold for 18-nt primers) is the most significant observed so far for the human telomerase TEN domain (24,27,29,30,50). Interestingly, this DNA binding defect was barely detectable using recombinant mutant +170 telomerase expressed in RRL. This may explain discrepancies in the literature regarding the extent of DNA affinity contributed by this region of the hTERT TEN domain (24,30).

DNA nucleotides 12–24 nt upstream of the primer 3'-end have been shown to crosslink to the *Tetrahymena* TEN domain (11,19) and provide extra binding affinity of DNA to the isolated human TEN domain (24). It is therefore likely that other amino acids in the TEN domain contribute to the classical telomerase 'anchor site' at the 5'-end of the primer (13–15). The intervening nucleotides may 'loop-out' during processive elongation as previously proposed (13,14), with the TEN domain functioning to tether the primer 5'-end to the site of nucleotide addition.

DNA affinity is known to be a determinant of telomerase repeat addition processivity (51). In the case of hTERT mutant +170, decreased RAP is not rescued at high primer concentration (Supplementary Figure S1), suggesting that the decrease in RAP is a consequence of failure to efficiently copy an entire repeat, rather than due to the DNA affinity defect of this mutant. Mutation of hTERT amino acids L13 and L14 also leads to a decrease in RAP without a decrease in DNA affinity (31). It was proposed that the decrease in RAP of the latter mutant is due to altered conformational changes at the translocation step, although a specific defect in extension of primers pairing near the 3'-end of the template was also observed (31).

Previous studies on the biochemical properties of mutations in the TEN domain point to a number of functions for this region. These data, in combination with our new results, lead us to propose that the hTERT TEN domain comprises at least three functional domains (Figure 7b):

(i) A DNA binding groove that is essential for DNA affinity (Figure 7b, yellow, red and green). This groove seems to be conserved between species, since it was first identified in the structure of the *Tetrahymena* TEN domain, and can be crosslinked to DNA primers in this species (19,28). In human TERT, N95 has been shown to directly contribute...
to affinity (29) and modelling supports a direct interaction with the DNA primer (Supplementary Figure S5). Our modelling data indicate that the 3′-end of the DNA is bound by the 170–175 pocket, while the 5′-end of the DNA extends towards P188 (Supplementary Figure S5).

(ii) An overlapping region that is involved specifically in conformational changes during nucleotide addition, the function elucidated in this study (Figure 7b, yellow and blue). It remains possible that part of this region (blue, L13 and L14) instead forms an interaction with a different part of TERT that is important during translocation (31).

(iii) A protein–protein interaction domain, part of the previously defined N-DAT domain (32), that is involved in telomerase recruitment to the telomere (Figure 7b, purple). Mutation of amino acids 122–136 in hTERT impacts little on enzyme activity, processivity, $K_m$, or primer $K_d$ as measured by a direct binding assay (29, this study). Instead, there is evidence that targeting these mutant telomerase molecules to the telomere overcomes their defect in vivo (52,53). The role of these amino acids in interacting with other proteins is supported by their location on the surface of our homology model (Figure 7b, purple).

Other proposed functions of the hTERT TEN domain include low-affinity RNA binding that has not been localized to particular amino acids (18,23,54,55) and a nucleolar localization domain in a region not shared with Tetrahymena TERT (45).

Compared to many other enzymes, telomerase has a complicated mechanism of action that involves many movements of the RNA template and DNA primer relative to the enzyme’s active site. Not surprisingly, $K_m$ is not an accurate prediction of the finer mechanistic details of telomerase. This study represents one of the first attempts to assign quantitative values to multiple kinetic parameters of human telomerase. This process has uncovered the importance of an essential N-terminal domain of the protein in events at the enzyme’s active site.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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