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Identification of Human Rap1: Implications for Telomere Evolution

Bibo Li, Stephanie Oestreich,† and Titia de Lange* The Rockefeller University

no recognized orthologs in budding yeast. Here, we hTERT protein (Lendvay et al., 1996; Lingner et al., 1997; three conserved sequence motifs in common with (Lundblad and Szostak, 1989) and Est3p (Lendvay et scRap1, is located at telomeres, and affects telomere al., 1996), have not been found so far. length. However, while scRap1 binds telomeric DNA The lack of sequence similarity between the vertetending the comparison of telomeric proteins to fission scRap1p (Shore and Nasmyth, 1987) is particularly strikyeast, we identify *S. pombe* **Taz1 as a TRF ortholog, ing given that both are duplex telomeric DNA binding indicating that TRFs are conserved at eukaryotic telo- factors and both act as negative regulators of telomere those of vertebrates, contained a TRF-like protein as Steensel and de Lange, 1997; Smogorzewska et al., well as Rap1. We propose that budding yeast pre- 2000). Rap1p is the major telomeric DNA binding activity served Rap1 at telomeres but lost the TRF component, in yeast (Buchman et al., 1988; Klein et al., 1992) and possibly concomitant with a change in the telomeric** every chromosome end in yeast contains \sim 20 high-

zling that there is no ortholog of TRF1 and TRF2 at ascribed a function.

telomeres in budding yeast. Similarly, TIN2 and tankyrase, proteins associated with mammalian telomeres through their interaction with TRF1, have no recognized New York, New York 10021 orthologs in yeast (Smith et al., 1998; Kim et al., 1999). Inversely, orthologs of a number of budding yeast telomeric proteins (for instance, Sir3p, Sir4p, Rif1p, Rif2p, Cdc13p, Stn1p, and Tel2p) have not been identified in Summary mammals, although complete genome information will be required to settle this issue. The telomerase reverse It has been puzzling that mammalian telomeric pro- transcriptase, Est2p, represents a striking exception to teins, including TRF1, TRF2, tankyrase, and TIN2 have this theme, being obviously homologous to the human
no recognized orthologs in budding yeast. Here, we hTFRT protein (Lendyay et al., 1996; Lingner et al., 1997; **describe a human protein, hRap1, that is an ortholog Nakamura et al., 1997). However, human orthologs of** other yeast telomerase components, such as Est1p.

directly, hRap1 is recruited to telomeres by TRF2. Ex- brate TRF proteins and the yeast telomeric protein meres. The data suggest that ancestral telomeres, like length (Kyrion et al., 1992; Marcand et al., 1997; van repeat sequence. affinity Rap1p binding sites within its irregular TG₁₋₃ telomeric repeat tract (Wang et al., 1990; Gilson et al., 1993). Additional Rap1p binding sites occur upstream Introduction of genes for many ribosomal proteins and glycolytic The protection and maintenance of human chromosome

enzymes (for review, see Buchman et al., 1988; Capieaux

ends requires the function of two related factors, TRF1

the duplex telometric TTAGGC

these genes through a tran et al., 1999; Smogorzewska et al., 2000); overexpression in themselves can result in resetting of telomere length
of full-length TRF1 and TRF2 results in gradual telomeric (Marcand et al., 1997). A second putative protein -a et al., 2000).
Although certain parallels between mammalian and **the accompact of scraptite on terminus** of scRap1p based on se-**Although certain parallels between mammalian and quence comparison (Bork et al., 1997; Callebaut and** Mornon, 1997), but this part of scRap1p has yet to be

It was previously noted that scRap1p, like TRF1 and *To whom correspondence should be addressed (e-mail: delange FRE2, binds telomeric DNA with two Myb-type helix-

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Thesent address: Department of Molecular Biology, Massachu. Bianchi et al., 1997, t present address: Department of Molecular Biology, Massachu-
setts General Hospital, Harvard Medical School, Boston, Massachu-
However, this shared DNA binding mode is achieved setts 02114. using substantially different strategies. In scRap1p, two

setts General Hospital, Harvard Medical School, Boston, Massachu-

DNA binding modules are positioned in tandem in the Rap1p, including a region unique to Rap1p, suggesting middle of the polypeptide, while TRF1 and TRF2 each that it is a human ortholog of Rap1p. We therefore refer contain a single C-terminal Myb domain and use homo- to this protein as hRap1. dimerization to place two Myb domains on DNA (Bianchi The BRCT domain is an z**95 aa presumed protein– et al., 1997, 1999; Broccoli et al., 1997). As a con- protein interaction motif first found in BRCA1, p53BP1, sequence of their different architecture, TRF1 and and Rad9p (Koonin et al., 1996) and now recognized in scRap1p recognize telomeric DNA in different ways. The a large number of other proteins linked to DNA repair two Myb domains of scRap1p recognize two tandem (Bork et al., 1997; Callebaut and Mornon, 1997). X-ray GGTGT sites at 8 bp distance (Konig et al., 1996), a crystallography of the XRCC1 second BRCT domains sequence arrangement frequently represented in the revealed an autonomously folded domain composed of yeast telomeric TG₁₋₃ repeat tracts. The two Myb do-** a four-stranded parallel β sheet surrounded by three α

mains in the TRF1 dimer, by contrast, interact with their helices (Zhang et al., 1998). Although the BRCT do **mains in the TRF1 dimer, by contrast, interact with their helices (Zhang et al., 1998). Although the BRCT domains YTAGGGTTR half sites regardless of their relative orien- of the three Rap1p orthologs are highly diverged, most tation and distance, as if the Myb domains are tethered of the conserved patches of hydrophobic residues can** by unusually flexible linkers (Bianchi et al., 1999). A Myb-
type DNA binding domain is also present in the *S. pombe* ARCC1 (Figure 1D). **XRCC1 (Figure 1D). type DNA binding domain is also present in the** *S. pombe* protein Taz1, another telomeric protein that functions **as a negative regulator of telomere length, but no other as revealed by structural analysis (Konig et al., 1996). overt sequence similarity to the TRFs or Rap1p was The first of these domains (called R1) can be recognized** recognized (Cooper et al., 1997; Vassetzky et al., 1999;

a novel human telomeric protein with a role in telomere ScRap1p R1 has some notable peculiarities, including length regulation that has significant sequence identity a long linker between helix 1 and helix 2, and the pres-
(24%–25%) with scRap1p in three different domains ence of phenylalanine and tyrosine residues instead of **ence of phenylalanine and tyrosine residues instead of (24%–25%) with scRap1p in three different domains. Since this human protein has both structural and func- the three highly conserved tryptophan residues typical** tional similarity to yeast Rap1p, we refer to it as hRap1. If Myb domains (Konig et al., 1996; Konig and Rhodes,
In the context of the identification of human Rap1, we
reexamined the structural similarities of the telomer **b** and it carries an insert of \sim 10 aa between helix 1 and 2, **protein complex in** *S. cerevisiae*, *S. pombe*, and verte-
 CONSISTENT CONSISTENT a close relationship between the human brates, revealing previously unappreciated relationships and suggesting a model for the evolution of eukaryotic and yeast Rap1 genes (Figure 1E). Furthermore, the Myb
telomeres. domains of human and yeast Rap1 both lack th

tectable as an \sim 2.5 kb species with a ubiquitous expres-
sion pattern (Figure 1B). The open reading frame in this
cDNA predicts a 399 aa protein of \sim 47 kDa and a Profile
scan motif search revealed that the ORF con **1997), central Myb domains (Konig et al., 1996), and an into a transactivation domain of about 65 aa and a more acidic C terminus, we examined the sequence similarity terminal domain (sometimes referred to as silencing dotail. Alignments revealed an additional region of se- regulation (Hardy et al., 1992a; Kyrion et al., 1992, 1993; quence similarity in the C termini of the human and yeast Liu et al., 1994; Buck and Shore, 1995; Liu and Lustig, proteins that coincides with the main protein–protein 1996). This part of Rap1p binds Rif1p and Rif2p and interaction domain of scRap1p (Hardy et al., 1992b; Mor- interacts with Sir3p and Sir4p (Hardy et al., 1992b; Moretti et al., 1994; Cockell et al., 1995; Liu and Lustig, 1996; etti et al., 1994; Cockell et al., 1995; Liu and Lustig, 1996; Wotton and Shore, 1997) (Figure 1F). Thus, the TRF2- Wotton and Shore, 1997). Alignment of hRap1 with the interacting protein has three domains in common with C terminus of scRap1p reveals significant sequence**

Spink et al., 2000).
A search for TRE2-interacting proteins has revealed son et al., 1994; Lipsick, 1996; Konig and Rhodes, 1997). **A search for TRF2-interacting proteins has revealed son et al., 1994; Lipsick, 1996; Konig and Rhodes, 1997). served glycine just beyond helix 1 and they do not have a patch of aliphatic residues beyond helix 2. Finally, Results both human and yeast Rap1 have a histidine residue at** A TRF2-Interacting Protein with Sequence

Similarity to Yeast Rap1p

A standard yeast two-hybrid screen of HeLa cDNAs with

A standard yeast two-hybrid screen of HeLa cDNAs with

part of the human TRF2 protein as bait yie

main, SD) that confers telomeric silencing and length

Figure 1. Sequence and Expression of the Human Ortholog of Yeast Rap1p

(A) Domain structure of hRap1. BRCT, BRCA1 C terminus; Myb, Myb-related HTH motif; Coil, predicted coiled domain; RCT, homology to the Rap1p C terminus; NLS, nuclear localization signal. The original two-hybrid clone and peptides used for raising antibodies are indicated. (B) Ubiquitous expression of hRap1 mRNA. Northern blot (Clontech) probed with the original hRap1 two-hybrid clone or b**-actin. Lane 1, spleen; lane 2, thymus; lane 3, prostate; lane 4, testis; lane 5, ovary; lane 6, small intestine; lane 7, colon (mucosal lining); lane 8, peripheral blood leukocytes.**

(C) Amino acid sequence of hRap1. The BRCT, Myb, Coil, and RCT domains are colored as in (A). The putative NLS is shown in bold.

(D) Alignment of BRCT domains in hRap1 and several other proteins based on the XRCC1 BRCT structure (Bork et al., 1997; Zhang et al., 1998). Four b **sheets (red arrows) and three** a **helices (blue bars) are marked. Bold residues are highly conserved (Bork et al., 1997), and other sequence similarities (Kyte-Doolittle) are shaded.**

(E) Alignment of the Myb domains of Rap1 with human c-Myb R2, TRF1, spTaz1, and scTbf1p using ClustalW. The position of three a **helices in c-Myb (Ogata et al., 1994) are indicated. Omitted amino acids between the** a **helices are indicated in parentheses. The three most conserved tryptophans are in bold. Shaded residues highlight sequence similarities (Kyte-Doolittle).**

(F) Alignment of the C-terminal region of three Rap1 proteins using ClustalW. Green shading highlights sequence similarities in the RCT domain (represented by the C-terminal 108 aa of hRap1). Identities in all three Rap1s are in bold. The most conserved 52 aa are underlined. The boxed region indicates the C-terminal tail domain of scRap1p (Liu and Lustig, 1996).

(G) Overview of the sequence similarities in three Rap1 proteins listing percentage identities (black) and similarities (gray) in each pair-wise comparison.

similarity in the final 108 aa of these proteins (Figure human and yeast Rap1 proteins is given in Figure 1G. human and scRap1p and 49% identity between the Rap1p is not represented as such in hRap1p. This re-**Rap1p proteins from** *K. lactis* **and** *S. cerevisiae* **(Figure gion, positioned between the Myb domain and the RCT 1F). Although its structure is not yet known, this part of domain, is not conserved in sequence and hRap1 has the Rap1 proteins is likely to have an identical fold. a predicted coiled domain that is not present in the**

1F), which we refer to as the *R***ap1** *C T***erminus (RCT) We have not found significant similarity between human domain. The highest level of conservation is found in and yeast Rap1p outside the three domains mentioned the final 52 aa where there is 30% identity between above. Notably, the** *trans***-activation domain of yeast An overall summary of the similarities between the yeast Rap1ps. Furthermore, we were unable to detect**

hRap1. (D) hRap1 can interact with itself through the RCT domain. of telomeric TTAGGG repeats (Chong et al., 1995), the (E) Summary of the TRF2-hRap1 interactions. Constructs on the left in (A, B, and D) and constructions on the right in (C) show LexA is predominantly located at telomeres. Similarly, hRap1 fusions. No transactivation activity was detected for all LexA-hRap1 colocalized with TRF2 in all cells examined (Figures 3D– fusion proteins. Constructs on the right in (A, B, and D) and construc-
tions on the left in (C) represent GAD fusions. β-galactosidase levels
(right column) were measured as described in the Experimental
Procedures. The **transformants. Values** ,**0.05 are indicated as 0. Ab 666 (Figure 3J). We surmise from these data that, as**

The yeast two-hybrid system was used to delineate the were transiently transfected with the dominant-negative tion. Deletion mapping indicated that hRap1 associated that forms inactive heterodimers with the endogenous with a central domain of TRF2 located between positions protein and effectively removes TRF2 from telomeres 123 and 366 (Figure 2A). Although TRF2 and TRF1 show (van Steensel et al., 1998). HeLa cells transfected with significant sequence similarity in the N-terminal half of this construct were examined by IF for the presence of this region (z**29% in the region from aa 45 to 245 in TRF2 on telomeres after gentle extraction with Triton TRF2 [Broccoli et al., 1997]), hRap1 does not interact X-100 to remove the nucleoplasmic proteins. A subpopwith full-length TRF1 in a two-hybrid setting (Figure 2B). ulation of the cells showed a severe loss of the punctate**

The TRF2-interacting domain in hRap1 was mapped to the C terminus (Figure 2C). The RCT domain in scRap1p is required for interaction with Rif1p, Rif2p, Sir3p, and Sir4p (Hardy et al., 1992b; Moretti et al., 1994; Cockell et al., 1995; Liu and Lustig, 1996; Wotton and Shore, 1997). Similarly, deletion mapping indicated that a region between aa 267 and 372 in the C terminus of hRap1 is responsible for the binding to TRF2 (Figure 2C). Yeast two-hybrid experiments also suggested homotypic interactions in the RCT domain. The area required for homotypic interactions in hRap1 is more extensive than that needed for binding to TRF2 and requires sequences from aa 267 to the very C terminus of the protein, including the last 27 aa (Figure 2D). The final 8 aa of the scRap1p RCT domain have been shown to be required for the interaction with Sir3p (Liu and Lustig, 1996). Consistent with the detection of homotypic interactions in the yeast two-hybrid assay, recombinant hRap1 behaves as a dimer in gel-filtration chromatography (D. Rhodes, personal communication). A summary of all the interaction data is given in Figure 2E.

TRF2 Recruits hRap1 to Telomeres

The subcellular localization of hRap1 was examined by indirect immunofluorescence (IF) of cultured human cells (Figure 3). A polyclonal rabbit antibody (Ab 765) was raised against [His]₆-tagged hRap1 expressed in a **baculovirus system and subjected to affinity purification. This reagent revealed a prominent punctate nuclear pattern in several different human cell lines (HeLa cervical carcinoma, HT1080 fibrosarcoma cells, and IMR90 primary fibroblasts) (Figures 3A and 3D and data not shown). The same telomeric pattern was seen in interphase nuclei stained with an hRap1-specific, affinitypurified, anti-peptide antibody (Ab 666 directed against aa 211–235, see Figure 1A) (Figure 3G), and exogenously expressed FLAG-tagged hRap1 also localized to telo-**Figure 2. Two-Hybrid Mapping of hRap1-TRF2 and hRap1-hRap1
Interacting Domains
(A) Mapping the hRap1-interacting domain in TRF2. (B) hRap1 does
INCOLOGIZED TRE1 in interphase nuclei (Figures 3A-3C). Since it was
not intera **in yeast, human Rap1 is concentrated at telomeric loci**

throughout the cell cycle.

transactivation with this or other parts of hRap1 in a

LexA fusion context (see below, Figure 2D).

at telomeres depended on its interaction with TRF2, as **suggested by the two-hybrid interaction of these pro-The C Terminus of hRap1 Is a Protein**
 teins and their coimmunoprecipitation (X.-D. Zhu et al.,
 Interaction Domain
 united). To this end, we examined Hel a cells which **Interaction Domain submitted). To this end, we examined HeLa cells which** allele of TRF2 (TRF2^{ABAM}), a truncated version of TRF2

Figure 3. hRap1 Locates to Telomeres through Interaction with TRF2

(A and D) Localization of endogenous hRap1 using indirect immunofluorescence with antibody 765 (green) in interphase nuclei of HeLaI.2.11 cells. (B) Localization of TRF1 in the same cells as in (A) using mouse antibody against TRF1 (red). (C) is a superimposition of (A) and (B). (E) Localization of TRF2 in the same cells as in (D) using mouse antibody against TRF2 (red). This antibody shows some staining at nontelomeric sites. (F) is a superimposition of (D) and (E). (G) Localization of endogenous hRap1 using antibody 666 (red) in interphase nuclei of HeLaI.2.11 $cells$ transiently transfected with TRF2^{ABAM}. **The cells were extracted with Triton X-100 before fixation. (H) Localization of TRF2 in the same cells as in (G) using mouse antibody against TRF2 (green). (I) is a superimposition of (G) and (H). (J) Localization of endogenous hRap1 using antibody 666 (green) on metaphase chromosomes in HeLaI.2.11 cells. DNA was stained with DAPI (blue). Bars in (C), (F),** (I) , and (J) represent $5 \mu m$.

TRF2 pattern, as expected from previous results with hRap1 (Figure 4A). Although a large number of paramethe TRF2^{ABAM} allele (van Steensel et al., 1998) (Figure ters were varied in additional experiments (for instance, **3H). In dual IF experiments for both TRF2 and hRap1, gel conditions, reaction buffer, and protein preparawe found a close correlation between the accumulation tions), we did not find evidence for the binding of teloof the two proteins in a punctate pattern (Figures 3G–3I). meric repeat DNA by hRap1. We also failed to detect a Specifically, all cells that had retained TRF2 at telomeres complex of hRap1 with single-stranded telomeric DNA** had lost the punctate TRF2 pattern (presumably due to TG_{1-3} repeats (data not shown). **expression of the TRF2**^{ABAM} allele) also lacked hRap1 When hRap1 was incubated with TRF2 and a
dots. Based on these data, interaction with TRF2 ap-
dsITTAGGGI₁₂ probe. a ternary complex was formed dots. Based on these data, interaction with TRF2 ap-

pears to be the predominant manner by which hRap1 (Figure 4R) Titration of TRF2 on DNA resulted in two **pears to be the predominant manner by which hRap1 (Figure 4B). Titration of TRF2 on DNA resulted in two accumulates on telomeres. In agreement, telomeric ac- complexes with different mobility (Figures 4A and 4B).**

Since hRap1 carries a Myb motif, it was pertinent to **examine its interaction with DNA. Using a double- reactions with hRap1 and TRF1 (data not shown). A allows binding of TRF2 to this probe, we failed to detect antibody directed to aa 283 to 307 of hRap1 (Ab 664, a protein–DNA complex with baculovirus-expressed see Figure 1A) demonstrated that the hRap1-dependent**

or with duplex probes containing the yeast telomeric

cumulation was observed with a C-terminal fragment of
hRap1 (aa 184 to the C terminus) which retains the TRF2
interacting domain, but lacks the BRCT and Myb do-
mains (data not shown).
TRF2 molecules. When hRap1 was added **tion, we observed an additional complex that migrated TRF2 and hRap1 Form a Complex above the two TRF2 complexes (Figure 4B). The appearon Telomeric DNA ance of this new band was strictly dependent on the**
Since hRap1 carries a Myb motif, it was pertinent to addition of hRap1 and was not observed in bandshift **stranded [TTAGGG]12 probe and a bandshift assay that supershift assay with an affinity-purified, anti-peptide**

Figure 4. hRap1 Binds to TRF2-Telomeric DNA Complexes

(A) Lack of telomeric DNA binding activity for hRap1. Bandshift assays with hRap1 and a ds(TTAGGG)12 probe. The highest concentration of hRap1 is 20 pM and the dilution series represents 3-fold steps. Lane 1 represents reaction with 1 pM TRF2. (B) hRap1 and hTRF2 form a ternary complex. Bandshift as-

says with increasing amounts of TRF2 (lanes 2 to 6, 3-fold steps up to 20 pM) or TRF2 and hRap1 protein (lanes 1 and 7, 2 pM of each protein). (C) The TRF2/hRap1 specific complex can be super-shifted by antibodies against hRap1. Bandshift assays with \sim 2 pM of TRF2 and **hRap1 and antibodies to hRap1 (Ab 664) and tankyrase (Ab 465) as indicated above the lanes. Lanes 4, 7, 8, and 9 contain 250 ng of antibody. Lane 5 contains 50 ng of antibody. Small arrows, TRF2 complex. Large arrows, hRap1-TRF2 complex.**

bandshift complex indeed contained hRap1 protein (Fig- that long telomeres recruit a greater number of Rap1p ure 4C). An antibody to the TRF1-interacting protein, molecules, resulting in inhibition of telomere extension tankyrase (Ab 465), served as a negative control in this in *cis***. How the C terminus of Rap1p regulates telomere experiment (Figure 4C). These data are consistent with elongation has not been clarified, but, the relevant part the observation that hRap1 binds to telomeres in a of Rap1p interacts with two proteins, Rif1 and Rif2, each**

Although hRap1 does not appear to bind DNA by itself, (Hardy et al., 1992b; Wotton and Shore, 1997). we considered the possibility that hRap1 might interact In order to explore a function for hRap1 in telomere with DNA in the context of the TRF2 complex. Such length regulation, we used a previously developed Tet**additional protein–DNA contacts often result in coopera- racyclin (Doxycyclin)-inducible system in the human fitive binding. An example is the interaction of two DNA brosarcoma cell line HTC75 (an HT1080 derivative) (van binding proteins, Mcm1 and** a**2, which bind to each Steensel and de Lange, 1997). HTC75 cells express high other on certain yeast operators, and display strong levels of telomerase, yet maintain their telomeres at a cooperative effects (**.**10-fold enhancement) in band- stable length setting over long periods of growth (300 shift assays with the appropriate target DNAs (Keleher population doublings [PD]). The stable length setting et al., 1989; Smith and Johnson, 1992). However, we of HTC75 telomeres depends on TRF1 and TRF2 (van failed to detect significant cooperativity for TRF2 and Steensel and de Lange, 1997; Smogorzewska et al., hRap1 under the conditions that allow detection of the 2000). Overexpression of TRF1 or TRF2 results in gradternary complex. No enhanced binding was detectable ual telomeric decline although telomerase levels are unwhen hRap1 was added to TRF2 band-shift reactions affected. (for instance, compare lanes 4 and 7 in Figure 4B and We derived five independent clonal HTC75 cell lines lanes 2 and 3 in Figure 4C). In addition, we failed to (F9, F11, F13, F14, and F17) overexpressing full-length, detect an effect of hRap1 on the off-rate of the com- FLAG-tagged hRap1 in an inducible manner. In each plexes (data not shown). We note that in some cases, cell line, Western analysis demonstrated inducible ex**additional protein DNA contacts are known to have a pression of an hRap1 protein that appeared slightly **relatively minor cooperative effect. For instance, the larger than endogenous hRap1, as expected from the binding of the second Myb domain in TRF1 homodimers presence of the FLAG tag (Figure 5A). Induction of hRap1 only increases the affinity by a factor of 10 (Bianchi in these cell lines did not affect the expression of endoget al., 1999). Footprinting analysis of the TRF2-hRap1 enous hRap1 (Figure 5A) or other telomeric proteins complex might provide further insight into this issue. (including TRF1, TRF2, and tankyrase) or telomerase**

meres. First, scRap1p recruits Sir3p and Sir4p to telo- during prolonged culturing of the cells. A control cell meres, resulting in the establishment of a heterochro- line (B31) transfected with the empty vector displayed matic state that represses the expression of nearby no changes in telomere length in the presence or abgenes (Moretti et al., 1994; Cockell et al., 1995; Liu and sence of doxycyclin (Figure 5B). Of five hRap1 express-**Lustig, 1996). Because we and others have failed to ing lines, one (F11) showed erratic telomere length demonstrate telomeric silencing in human cells (Broc- changes under uninduced conditions and was not studcoli and Cooke, 1994; H. Tommerup and T. d. L., unpub- ied further. The other cell lines showed consistent lished data), exploration of this potential role for hRap1 changes in telomere dynamics, in each case demonstrais currently not possible. The second known function ble as a gradual elongation in the mean length of the of scRap1p at chromosome ends is the regulation of telomeric restriction fragments (Figures 5C–5E and data telomere length (Conrad et al., 1990; Lustig et al., 1990; not shown). The growth of the telomeres was largely** Kyrion et al., 1992; Liu et al., 1994; Marcand et al., 1997). dependent on the induction of hRap1 expression, al-**Tethering experiments have demonstrated that cells though leaky expression of hRap1 under repressed conevaluate telomere length based on the number of ditions may have led to gradual telomere extension in scRap1p C-terminal domains present at chromosome the lines F9 and F13. The maximum telomere elongation ends (Marcand et al., 1997). The current "protein-count- rate in cells overexpressing hRap1 was 40–50 bp/PD ing" model for telomere length homeostasis dictates (based on the early stages of the F9 and F17 growth**

TRF2-dependent manner. of which contributes to telomere length homeostasis

activity (data not shown).

hRap1 Affects Telomere Length The effect of hRap1 on telomere length homeostasis ScRap1p has two well-defined functions at yeast telo- was determined by measuring telomere length changes

Figure 5. Overexpression of Full-Length hRap1 Results in Telomere Elongation

(A) Western analysis of the inducible expression of the FLAG-tagged hRap1 in three F cell lines. Extracts were prepared from cells grown in parallel in the presence (uninduced) or absence (induced) of doxycyclin for four population doublings and probed with the hRap1 antibody 765 that detects both the endogenous hRap1 and the induced, FLAG-tagged hRap1.

(B) Telomere length is maintained stably in clonal HTC75 line B31 containing empty expressing vector pUHD10–3. Chart showing the median telomere lengths versus PD in cells grown under induced (1**) or uninduced (**2**) conditions.**

(C–E) Time course of changes in telomere length in clonal HTC75 lines expressing FLAG-tagged hRap1 (F9 in C, F17 in D, and F13 in E) grown in the absence and presence of doxycyclin (+ and - induction, respectively). Top, genomic blotting analysis of telomeric restriction fragments **in HinfI/RsaI digested DNA, probed with a TTAGGG repeat probe. Bottom, chart showing the changes of median telomere lengths in each corresponding cell line.**

PD in F14 (data not shown). However, much more mod- and TRF2 (Bianchi et al., 1997; Broccoli et al., 1997; Kim which showed extension at a rate of 17 bp/PD (Figure 29% sequence identity in this region (Broccoli et al., 5E). Similar clonal variations in response were previously 1997; Smith and de Lange, 1997). Alignment of Taz1 observed with HTC75 clones expressing TRF1 (van with this part of TRF1 and TRF2 revealed significant Steensel and de Lange, 1997). **Subsete and Steen and Steensel and Steensel and definition** Steensel and S

Prompted by the identification of the ortholog of yeast 180 aa (Figures 6A and 6B). Rap1p at human telomeres, we examined the possibility

of other such relationships, Taz1 was identified in S. TRF1 and TRF2: a Myb domain that is substantially more **of other such relationships. Taz1 was identified in** *S.* **TRF1 and TRF2: a Myb domain that is substantially more** *pombe* **by a one-hybrid screen using telomeric DNA and similar to that of the TRFs than to scRap1p R1 (Figure also by a screen for loss of telomeric silencing (Cooper 1E) and the TRFH domain (Figure 6A). Furthermore, the et al., 1997; Nimmo et al., 1998). The ORF of Taz1 showed overall architecture of Taz1 seems to resemble that of a Myb domain near the C terminus but no other se- the TRFs since each of these proteins bind to DNA as quence similarity to TRFs or Rap1p had been recog- a homodimer (Bianchi et al., 1997; Broccoli et al., 1997;** $nized.$ The identification of the key sequences con**served in scRap1 and hRap1 showed that Taz1, which Taz1, like the TRFs, acts as a negative regulator of telolacks the BRCT and the RCT domains, is not a member mere length (Cooper et al., 1997). Based on these strucof this family. tural and functional similarities, we suggest that Taz1 is**

TRF1 and TRF2 have been identified in mice and hu- a member of the TRF family of telomeric proteins.

curves). Telomere growth occurred at a rate of 30 bp/ with protein–protein interaction domains in both TRF1 est rates of telomere elongation were found in F13, et al., 1999). The mammalian TRFs have approximately dard ClustalW settings, Taz1 can be aligned to TRF1 Taz1 Is a Member of the TRF Protein Family yielding 24% sequence identity and 47% similarity over

mans and sequence alignments have shown that in addi- It is not clear whether Taz1 is more closely related to tion to their C-terminal Myb motif, the TRFs have a defin- TRF1 or to TRF2. Although the TRFH and Myb domains ing sequence motif near their N termini. This domain, of Taz1 are more similar to those of TRF1, the region called TRF homology (TRFH) domain here, is a region between the TRFH and the Myb domain bears more of z**200 aa with unknown function, although it overlaps similarity to TRF2. The TRF1 and TRF2 proteins are** A

B

cal residues are in bold; sequence similarities are shaded (Kyte-**Doolittle). that cells measure telomere length through counting**

contrast, the N terminus of Taz1 has no overriding acidic of overexpression of human Rap1 is very similar. In-

Proteins that bind to the duplex part of the telomeric required for stable telomere length maintenance. It will repeat array have been studied in vertebrates and in the be necessary to establish the effect of diminished hRap1 yeasts *S. cerevisiae* **and** *S. pombe***. The identification activity to settle this issue. of the human ortholog of the budding yeast telomeric Although the hRap1-interacting partner TRF2 is reprotein Rap1p now suggests an evolutionary relation- quired for the protection of chromosome ends (van ship between the telomeric complexes in these three Steensel et al., 1998; Karlseder et al., 1999), it has yet systems. We propose that telomeres in vertebrates and to be established whether hRap1 contributes to this** in *S. pombe* represent an ancestral situation in which a function. A possible role for yeast Rap1p in maintaining **TRF-like protein and a Rap1-like factor both functioned telomere integrity is suggested by the instability of teloat telomeres, whereas the arrangement found in** *S. cere-* **meres in rap1–17 cells (Kyrion et al., 1992). In addition,** *visiae* **is apparently different, representing telomeres overexpression of yeast Rap1p induces a mild chromo**without the TRF module. We propose that the telomeric some instability phenotype (Conrad et al., 1990) and

complex in *S. cerevisiae* **evolved through a combination of gene loss and nonorthologous gene displacement and that these changes were accompanied by an alteration in the telomeric DNA sequence. Our findings may allow a more meaningful comparison of telomere biology in yeast, vertebrates, and other eukaryotes.**

Structural and Functional Similarities of Human and Yeast Rap1

Human Rap1 has three domains in common with yeast Rap1p. It contains a central Myb motif with specific sequence features also present in the yeast Rap1p R1 Myb domain. In addition, human Rap1, like yeast Rap1p, carries a BRCT domain at the N terminus and the Rap1 proteins share a conserved motif (the RCT domain) in their C termini. Functionally, hRap1 also displays strong similarities to scRap1p. Like the yeast protein, hRap1 is an integral component of the telomeric complex and the majority of the protein is detectable at chromosome ends by indirect immunofluorescence. Furthermore, in both yeast and human Rap1, the C terminus of the protein functions to bind other telomeric proteins.

Yeast and human Rap1 may also be similar with regards to their effect on telomere maintenance. Yeast Rap1p acts as a negative regulator of telomere length. A number of rap1 mutants in yeast display telomere elongation phenotypes (Kyrion et al., 1992; Liu et al., 1994) and mutation of the Rap1p binding sites in yeast telomeres can result in run-away telomere elongation (McEachern and Blackburn, 1995; Krauskopf and Blackburn, 1996). Furthermore, the scRap1p-interacting factors, Rif1p and Rif2p, display characteristics consistent with a negative regulatory pathway that limits telomere elongation (Hardy et al., 1992b; Wotton and Shore, Figure 6. S. pombe Taz1 Is a Member of the TRF Family

(A) Alignment of the TRFH domain of mouse and human TRF1,

mouse and human TRF2, and S. pombe Taz1 using ClustalW. Identi-

cal residues are in bold: sequence similari **(B) Overview of the sequence similarities in TRF and Taz1 proteins, the number of Rap1p C-terminal domains at individual**

Moderate overexpression of scRap1p results in slight paralogs with strikingly different N termini (Chong et al., the cause the increased abundance of scRap1p titrates

1995; Broccoli et al., 1997). The first 45 aa of TRF2 are

1995; Broccoli et al., 1997). The first 45 aa of extension (e.g., one of the Rif proteins). The phenotype **or basic nature. creased hRap1 level in the telomerase positive cell line HTC75 resulted in a moderate telomere elongation. It is Discussion possible that the mild telomere extension phenotype is due to titration of an hRap1-interacting protein that is**

listing percentage identity (black) and similarity (gray). chromosome ends.

several rap1^{ts} mutants have effects consistent with **scRap1p acting to protect chromosome ends (Lustig et al., 1990). Like inhibition of TRF2, complete loss of Rap1p in yeast is lethal but this phenotype may be primarily due to nontelomeric functions.**

A BLAST search with full-length hRap1 against the budding yeast genome did not yield scRap1p or klRap1p. Similarly, a search with Taz1 yielded several Myb motif proteins but not the TRFH domains of any of the four mammalian TRFs represented in the nonredundant database. These findings illustrate current difficulties in finding highly diverged but biologically informative orthologs. Precise definition of the most conserved motifs in the TRF and Rap1 families may facilitate the identification of telomeric proteins in other organisms.

S. pombe **Taz1 Is a Member of the TRF Family**

We propose that Taz1 is a member of the TRF family of duplex telomeric DNA binding proteins. This family is defined by the presence of two regions of sequence similarity: a Myb motif that is closely related to c-Myb R1/R2 and a TRF homology domain. TRF family proteins undergo homotypic interactions and bind telomeric DNA as dimers or higher order oligomers (Bianchi et al., 1997; Broccoli et al., 1997; Spink et al., 2000). In addition, each of these proteins have a negative effect on telomere Figure 7. Model for the Evolution of Eukaryotic Telomeres length maintenance in telomerase expressing cells (A) A model for the evolution of duplex telomeric DNA binding pro- (Cooper et al., 1997; van Steensel and de Lange, 1997; teins. See Discussion for details. TRF/Taz1 and Rap1 proteins are

nance, possibly by interfering with recombination at *pombe* **ortholog of Rap1 has not been identified and its presence telomeres (Nakamura et al., 1998). This function is remi- at** *S.pombe* **telomeres is speculative. Loss of the TRF module is niscent of the protective activity of TRF2, which appears proposed to have occurred in budding yeasts**. The proposed struc-
 proposed to a suppropriate DNA ropein at mammalian ture of the telomeric complex in the last to suppress inappropriate DNA repair at mammalian ture of the telomeric complex in the last common ancestor of tungi
telomeres (van Steensel et al., 1998). Secondly, Taz1
plays a role in meiosis. Taz1 mutants display a def in the horse-tail stage of meiosis I in *S. pombe* (Cooper TRF tether.

et al., 1998; Nimmo et al., 1998), in which telomeres (B) Prepone **associate with the spindle pole body and facilitate the eukaryotes with the exception of budding yeasts. For references on** rapid migration of the meiotic nucleus through the cell

(Chikashige et al., 1994, 1997; Kohli, 1994). This telo-

mate lineage of eukaryotes is based on the phylogeny listed in http://

mate lineage of eukaryotes is based **taz1**² **cells showing loss of clustering of telomeres at the spindle pole body and a marked drop in meiotic**

TRF family suggests that a TRF-like protein originally motif and has no known telomeric DNA binding activity. was engaged on telomeres in the last common ancestor Thus, either scRap1p has gained a Myb domain or of fungi and vertebrates. According to this view, *S. cere-* **hRap1 has lost one. Determination of the telomeric DNA Hemi- and Eu-ascomycetes (represented in Figure 7B into this issue. It is of obvious interest to establish the by budding yeasts and** *Neurospora***, respectively). function of the single Myb motif in hRap1.**

hRap1 to telomeres, this may have been one of the a TRF-like Myb domain, the TTAGGG repeat binding

Smogorzewska et al., 2000).
Taz1 has two additional functions. First, it suppresses
a telomerase-independent pathway of telomere mainte-
DNA contacts of hRap1 in vivo have not been analyzed. The S.

(B) Preponderance of TTAGGG-like repeats at telomeres of most eukaryotes with the exception of budding yeasts. For references on

recombination and spore viability. It will be of interest
to determine whether TRF1 or TRF2 has a role in mam-
malian meiosis, which features an analogous chromo-
some reorganization in the bouquet stage of meiosis l
(see **or two Myb domains. ScRap1p has two Myb domains Evolution of Eukaryotic Telomeres and binds to telomeres through a direct interaction with The recognition of** *S. pombe* **Taz1 as a member of the the telomeric DNA, while human Rap1 has a single Myb** *visiae* **most likely lost the TRF-like telomere binding binding activity and the number of Myb domains in Rap1 component at some point during the divergence of the in** *S. pombe* **or another fungus could provide insight**

Given that human TRF2 facilitates the binding of The genome of *S. cerevisiae* **encodes one protein with**

protein Tbf1p (Brigati et al., 1993; Bilaud et al., 1996). Rap1 sites appear to be the only common motif in the Tbf1p has not been ascribed a function at telomeres so highly diverged telomeric sequences of budding yeasts far. It is unlikely that Tbf1p is a remnant of the budding (McEachern and Hicks, 1993; Krauskopf and Blackburn, yeast TRF gene since Tbf1p lacks a TRFH domain and 1996, 1998). The DNA binding activity of scRap1p is an *S. pombe* **Tbf1p ortholog is reported in the database. also important for its role as a transcriptional regulator.**

proteins? One possibility is that during the evolution of described here has a similar function, it is not excluded the budding yeasts, including *S. cerevisiae, K. lactis***, that mammals have a second ortholog of Rap1 with a and several** *Candida* **species, the telomeric DNA se- role in gene expression. quence may have been altered such that the TRF com**ponent could no longer bind to the telomeric repeat **Experimental Procedures array. A mutation in the telomerase RNA template could have caused this alteration, and binding of Rap1 to the Isolation of hRap1 new version of the telomeric DNA might have rescued A two-hybrid screen was carried out with the yeast reporter strain** the potential lethality of this change (McEachern and
Blackburn, 1995; Krauskopf and Blackburn, 1996). An HeLa S3 matchmaker cDNA library (Clontech), and a LexA-fused
alternative scenario is that loss of the TRF gene resul **alternative scenario is that loss of the TRF gene resulted frame shift mutation leading to a TRF2 protein with 444 aa that is tions in their telomerase RNA template gene that allowed tions 200 to 399, see Figure 1A) resulting from this screen was used**

Perusal of the large collection of available telomeric
DNA sequences indicates that the TTAGGG repeat of
present day vertebrate telomeres is the most pervasive
discussion (GenBank accession number: AF262988) encoded a **telomeric sequence in eukaryotes, also occurring in ki- predicted ORF of 399 aa. netoplastida, slime molds, and most branches of fungi (e.g., Ustilago, Pneumocystis, Neurospora, and Asper- Two-Hybrid Assays gillus) (reviewed in Henderson, 1995) (Figure 7B). The The LexA-TRF2 full-length fusion was generated by inserting a PCR majority of other eukaryotes have telomeric sequences** fragment encoding TRF2 aa 2–389 into pBTM116 (Bartel et al., 1993)
 that are closely related to the TTACCC repeats for at the EcoRI site, followed by replacement of that are closely related to the TTAGGG repeats, for
example TAGGG in Giardia, TT[T/A]GGG in tomato,
TTAGGC in C. elegans, and TTGGGG in the ciliate Tetra-
of sequences encoding the indicated amino acids from the LexA**hymena. Even the heterogenous telomeric repeat of** *S.* **TRF2 full-length clone followed by insertion into the polylinker region** *pombe* **was recently recognized as being composed of of pBTM116. GAD-TRF2 hybrid was generated by cloning the LexAa TTACAGG repeat motif (Hiraoka et al., 1998). Thus, TRF2 StuI/BamHI fragment into pACT2 (Clontech). The LexA-hRap1** the last common ancestor of all eukaryotes most likely
carried TTAGGG repeats or a closely related sequence
at its telomeres. Budding yeast telomeric DNA repre-
sents a striking departure from the TTAGGG motif, con-
sents **taining telomeric repeats that are very different in length tion between TRF1 and hRap1. Expression of fusion proteins was and sequence (e.g., ACGGATTTGATTAGGTATGTGG verified by Western blotting using anti-LexA or anti-GAD monoclonal TGT in** *K. lactis* **[McEachern and Blackburn, 1994]). Pos-

sibly the precursor of the Hemiascomycetes suffered a Two-hybrid experiments were performed in the yeast strain L40.**

We propose a model for the evolution of the telomeric values from individual transformants differed by <30% from the complex in which the ancestral eukaryotic telomeres average. **were composed of TTAGGG repeats and bound a TRFlike protein as well as a Rap1-like factor. During the Recombinant hRap1 and Generation of hRap1 Antibodies evolution of** *S. cerevisiae* **and other budding yeasts, a

N**-terminally [His]₆-tagged full-length hRap1 protein was expressed
 telomerase mutation may have caused a shift in the in insect cells using the Bac-To-Bac **telomerase mutation may have caused a shift in the in insect cells using the Bac-To-Bac Expression System (Monsato** telomeric sequence, resulting in loss of the tethering
function of the TRF module (or vice versa). In agreement,
TRF1 does not bind telomeric DNA from S. cerevisiae
rem, Ab 765, was affinity purified using hRap1 coupled to (Zhong et al., 1992). The optimal binding site for current $\overline{YPDLPEC\text{-}COOH}$ starting at aa 211 and hRap1-283 (NH₂-DPP
day human TRF1 contains two (overlapping) copies of TPEEDSETOPDEEEEEEEKVSC-COOH) starting at aa 283 o **the sequence YTAGGGTTR (Bianchi et al., 1999), a pre- were synthesized (BioSynthesis, Lewisville, TX), conjugated to KLH cise representation of the TTAGGG repeat array, and (Pierce, Rockford, IL), and used to immunize NZW rabbits (Covance).** minor changes in this sequence strongly inhibit TRF1 The resulting immune sera, Ab 666 (against hRap1-211) and Ab
binding (Zhong of al. 1993) Hapish of al. 1994), The 664 (against hRap1-283), were affinity purified again binding (Zhong et al., 1992; Hanish et al., 1994). The price bound peptides (hRap1-211and hRap1-283 respectively)
DNA binding activity of scRap1 may have rescued cells using standard procedures (Harlow and Lane, 1988). **bearing telomeres containing Rap1 recognition sites. Once telomere function became independent of TRFs, Immunofluorescence and Metaphase Chromosome Spread divergence of the telomeric sequence could occur, as Preparation of metaphase chromosome spreads and indirect fluolong as the Rap1 binding sites were preserved. Indeed, rescence techniques are described in Chong et al., 1995. In brief,**

Why does *S. cerevisiae* **lack the TRF class of telomeric Although there is no indication that the ortholog of Rap1**

in selection of yeasts with specific compensatory muta- mutated from aa 389. The hRap1 two-hybrid cDNA clone (aa posi-Rap1 to bind to telomeric DNA independently. as probe on a cDNA library made from a human breast cancer cell
Perusal of the large collection of available telomeric line (MDA) (Kratzschmar et al., 1996), yielding 26 pos

Sibly the precursor of the Hemiascomycetes suffered a
major change in the telomerase RNA template region
(see also discussion in Brun et al., 1997).
We propose a model for the evolution of the telomeric
We propose a model

day human TRF1 contains two (overlapping) copies of TPEEDSETQPDEEEEEEEEKVSC-COOH) starting at aa 283 of hRap1

phate-buffered saline (PBS), fixed in 2% formaldehyde in PBS for al., 1992) and filled in with [α -³²P]dCTP and Klenow enzyme. **10 min, and permeabilized with 0.5% Nonidet P-40 in PBS for 10 min. To prepare metaphase chromosome spread, HeLa I.2.11 cells Tetracyclin Inducible Expression of hRap1 in HTC75 Cells** were treated with colcemid (0.1 µg/ml, 90 min), harvested by trypsin-

ization, washed with serum and PBS, hypotonically swollen in 10 rying an N-terminal FLAG epitope tag was created by PCR cloning **ization, washed with serum and PBS, hypotonically swollen in 10 rying an N-terminal FLAG epitope tag was created by PCR cloning** mM Tris (pH 7.4), 10 mM NaCl, and 5 mM MgCl₂ (37°C, 15 min), and
sedimented onto cover slips for 15 s at 3000 rpm in a Sorvall was generated by inserting paired oligos encoding a FLAG tag into **sedimented onto cover slips for 15 s at 3000 rpm in a Sorvall was generated by inserting paired oligos encoding a FLAG tag into RT6000B tabletop centrifuge. Chromosome spreads were immedi- BamHI site of pUHD10–3. A fragment representing the complete ately fixed and permeabilized as described above. To examine the coding region of hRap1 was generated by PCR using Pfu-polymerhRap1 localization in interphase HeLa1.2.11 cells transfected with** ase and the primer pair 5'-GCGCAGATCTGCGGAGGCGTTGGAT
hTRF2^{4BAM}, cells were washed with PBS, extracted with Triton X-100 TTG-3' and 5'-GCGCGGATCCTTATT **buffer (0.5% Triton X-100, 20 mM HEPES [pH 7.9], 50 mM NaCl, 3 and cloned into the BamHI site. The resulting plasmid was introtwice, fixed with 3% formaldehyde, 2% sucrose in PBS for 10 min, and independent clonal cell lines (F lines) were isolated using cloning permeabilized with Triton X-100 buffer for 10 min, washed with PBS cylinders. Growth conditions of the cells and methods to measure twice followed by antibody staining. Most nucleoplasmic proteins telomere lengths were described previously (van Steensel and de are extracted in this method. Residual signals represent predomi- Lange, 1997). nantly DNA bound proteins.**

Endogenous hRap1 was detected with polyclonal antibodies 765 Acknowledgments and 666 (see above). TRF1 was detected with polyclonal mouse anti-TRF1 antibody (Ab 3, S. Smith and T. d. L., unpublished data),

We thank Daniela Rhodes for help in reviewing the hRap1 Myb

and BRCT domains and for communication of unpublished susults.

(Ab 2, X.-D. Zhu and T. d. L **Hammamatsu C4742–95 camera using Open Lab program (for other Received January 20, 2000; revised April 25, 2000. IF experiments). IP lab images were noise filtered, corrected for background, and merged using Adobe PhotoShop. References**

Cells grown to semiconfluence on 15 cm dishes were trypsinized,
Cells grown to semiconfluence on 15 cm dishes were trypsinized,
washed with media, harvested by centrifugation, washed with ice-
two-hybrid system to detect p **Bianchi, A., Smith, S., Chong, L., Elias, P., and de Lange, T. (1997). tein content of extracts was measured using the Bradford assay TRF1 is a dimer and bends telomeric DNA. EMBO J.** *16***, 1785–1794. (BioRad, Hercules, CA) using bovine serum albumin as a standard.**

and transferred to nitrocellulose membrane by electrophoresis. Pon- Lange, T. (1999). TRF1 binds a bipartite telomeric site with extreme ceau S-staining confirmed equal loading of the samples. Blots were spatial flexibility. EMBO J. *18***, 5735–5744. preincubated in 10% nonfat milk in PBS containing 0.5% Tween 20 Bilaud, T., Brun, C., Ancelin, K., Koering, C.E., Laroche, T., and dry milk powder and 0.1% Tween 20 in PBS. Blots were incubated telobox protein. Nat. Genet.** *17***, 236–239.** For 12–16 hr at 4°C with anti-hRap1 antibody 765, followed by three

10 min washing steps. Subsequently, blots were incubated for 30

min with horseradish peroxidase-conjugated donkey-anti-rabbit an-

tibody (Amersham) and

France bacanomias-expressed, phase and principle and in 2 pro-

terms were used for bandshift assays. Binding reactions (in $\frac{1}{3}$ and Shore, D. (1993).

erol, 0.5 μ g E. coli DNA, 20 mM GJycine-KOH [pH 9.0], 10 mM D **Broccoli, D., and Cooke, H.J. (1994). Effect of telomeres on the run on a 0.6% agarose gel in 0.1**3 **TBE at 4**8**C. Gels were dried onto** DE81 paper and analyzed by autoradiography or by exposure on a **interphase location of adjacent regions**
Phosphorimager, DNA probes were prepared as described pre- some. Exp. Cell Res. 212, 308–313. **Phosphorimager. DNA probes were prepared as described pre-**

HeLaI.2.11 cells (van Steensel et al., 1998) were washed in phos- containing (TTAGGG)12 repeats was purified from pTH12 (Zhong et

hTRF2^D**B**^D**M, cells were washed with PBS, extracted with Triton X-100 TTG-3**9 **and 5**9**-GCGCGGATCCTTATTTCTTTCGAAATTCAATCCT-3**9 duced into HTC75 as described previously (van Steensel et al., 1998)

Proteins (65 m**g) were separated on 8% SDS-polyacrylamide gels Bianchi, A., Stansel, R.M., Fairall, L., Griffith, J.D., Rhodes, D., and de**

Gilson, E. (1997). Telomeric localization of TRF2, a novel human

damage-responsive cell cycle checkpoint proteins. FASEB J. *¹¹***, Bandshift Assays 68–76.**

viously (Zhong et al., 1992). In brief, the HindIII/Asp718 fragment Broccoli, D., Smogorzewska, A., Chong, L., and de Lange, T. (1997).

to double-stranded regions of telomeric DNA. Trends Cell Biol. *7***, (1995). Identification fo a new family of tissue-specific basic helix-**

Buchman, A.R., Kimmerly, W.J., Rine, J., and Kornberg, R.D. (1988). 3813–3822. Two DNA-binding factors recognize specific sequences at silencers, Karlseder, J., Broccoli, D., Dai, Y., Hardy, S., and de Lange, T. (1999). upstream activating sequences, autonomously replicating sequences, p53- and ATM-dependent apoptosis induced by telomeres lacking and telomeres in *Saccharomyces cerevisiae*. Mol. Cell. Biol. 8,

between transcriptional activators, silencers, and telomeres as re- Mol. Cell. Biol. *9***, 5228–5230.**

Callebaut, I., and Mornon, J.-P. (1997). From BRCA1 to RAP1: a 935–948.

Chikashige, T., Ding, D.-Q., Fundbiki, H., Haraguchi, T., Mashiko, Konig, P., and Rhodes, D. (1997). Recognition of telomeric DNA.
Chromosome movement in fitsion years. Science 264, 270–273.
Chikashiga V. Ding. D.Q. Imai V

Chikashige, Y., Ding, D.Q., Imai, Y., Yamamoto, M., Haraguchi, T.,
and Hiraoka, Y. (1997). Meiotic nuclear reorganization: switching
the position of centromeres and telomeres in the fission yeast Schiz-
mour suppressors. N

Chong, L., van Steensel, B., Broccoli, D., Erdjument-Bromage, H., under-anchored metanoprotease-desintegrin protein with
Hanish, J., Tempst, P., and de Lange, T. (1995). A human telomeric integrin binding sequence. J. Biol

Cockell, M., Palladino, F., Laroche, T., Kurion, G., Liu, C., Lustig,
A.J., and Gasser, S.M. (1995). The carboxy termini of Sir4 and Rap1 peats. Nature 383, 354-357.
affect Sir3 localization: evidence for a multicomponent **telomere turnover in yeast. Proc. Natl. Acad. Sci. USA** *95***, 12486– required for yeast telomeric silencing. J. Cell Biol.** *129***, 909–924.**

12491. Conrad, M.N., Wright, J.H., Wolf, A.J., and Zakian, V.A. (1990). RAP1 protein interacts with yeast telomeres in vivo: overproduction alters Kyrion, G., Boakye, K.A., and Lustig, A.J. (1992). C-terminal trunca-

5173. Cooper, J.P., Nimmo, E.R., Allshire, R.C., and Cech, T.R. (1997). Regulation of telomere length and function by a Myb-domain protein

*visiae***. Genes Dev.** *⁷***, 1146–1159. Cooper, J.P., Watanabe, Y., and Nurse, P. (1998). Fission yeast Taz1** protein is required for meiotic telomere clustering and recombina-

Gilson, E., Roberge, M., Giraldo, R., Rhodes, D., and Gasser, S.M.

(1993). Distortion of the DNA double helix by RAP1 at silencers and

multiple telomeric binding sites. J. Mol. Biol. 231, 293-310.

Goscon M. and Buiard H

Gossen, M., and Bujard, H. (1992). Tight control of gene expression
in mammalian cells by tetracyclin-responsive promoters. Proc. Natl.
Acad. Sci. USA 89, 5547-5551.
Hanish I.P. Yanowitz J. and de Lange T. (1994). Strippen

Hanish, J.P., Yanowitz, J., and de Lange, T. (1994). Stringent se**quence requirements for telomere formation in human cells. Proc. Liu, C., and Lustig, A.J. (1996). Genetic Analysis of Rap1p/Sir3p**

Hardy, C.F.J., Balderes, D., and Shore, D. (1992a). Dissection of a
carboxy-terminal region of the yeast regulatory protein RAP1 with Liu, C., Mao, X., and Lustig, A.J. (1994). Mutational analysis defines **carboxy-terminal region of the yeast regulatory protein RAP1 with Liu, C., Mao, X., and Lustig, A.J. (1994). Mutational analysis defines** effects on both transcriptional activation and silencing. Mol. Cell. **Biol.** *12***, 1209–1217. in** *Saccharomyces cerevisiae***. Genetics** *138***, 1025–1040.**

protein involved in transcriptional silencing and telomere length reg- telomere elongation leads to senescence in yeast. Cell *57***, 633–643.**

(Cold Spring Harbor, NY: Cold Spring Harbor Press). length. Science *250***, 549–553.**

Henderson, E. (1995). Telomere DNA structure. In Telomeres, E.H. Marcand, S., Gilson, E., and Shore, D. (1997). A protein-counting Blackburn and C.W. Greider, eds. (Cold Spring Harbor, NY: Cold mechanism for telomere length regulation in yeast. Science *275***, Spring Harbor Press), pp. 11–35. 986–990.**

Human telomeres contain two distinct Myb-related proteins, TRF1 Hiraoka, Y., Henderson, E., and Blackburn, E.H. (1998). Not so pecu-

Brun, C., Marcand, S., and Gilson, E. (1997). Proteins that bind Hollenberg, S.M., Sternglanz, R., CHeng, P.F., and Weintraub, H. 317–323. loop-helix proteins with a two-hybrid system. Mol. Cell. Biol. *15***,**

210–225. Keleher, C.A., Passmore, S. , and Johnson, A.D. (1989) Yeast Repres- $\text{Sor } \alpha$ 2 binds to its operator cooperatively with yeast protein Mcm1.

vealed by functional analysis of a yeast DNA-binding protein. Mol.
Cell. Biol. 8, 5086–5099.
Buck, S.W., and Shore, D. (1995). Action of a RAP1 carboxy-terminal by login E. Laroche, T. Cardonas, M.E. Hofmann, L.E. Y. Schwe

Buck, S.W., and Shore, D. (1995). Action of a RAP1 carboxy-terminal
silencing domain reveals an underlying competition between HMR
and Gasser, S.M. (1992). Localization of RAP1 and Topoisomer-
ase II in nuclei and meiotic

widespread BRCT module closely associated with DNA repair. FEBS Kohli, J. (1994). Telomeres lead chromosome movement. Curr. Biol. Lett. *⁴⁰⁰***, 25–30.** *⁴***, 724–727.**

Capieaux, E., vigilals, M.-L., Seitenac, A., and Golleau, A. (1969).

The yeast H⁺-ATPase gene is controled by the promoter binding

factor TUF. J. Biol. Chem. 264, 7437-7446.

Chikashige, Y., Ding, D.-Q., Funabiki, H.,

Kratzschmar, J., Lum, L., and Blobel, C. (1996). Matargidin, a mem- *osaccharomyces pombe***. EMBO J. 16, 193–202.**
 brane-anchored metalloprotease-desintegrin protein with an RGD

protein. Science 270, 1663–1667.
 protein. Science 270, 1663–1667. *proche T. Kurien, G. Liu, C. Lustia 2000 2000 28000 <i>270, 28000 270, 28000 <i>270, 28000 270, 28000 270, 270, 270, 270, 270, 271, 271, 2*

telomere structure and decreases chromosome stability. Cell 63,
139–750. T39–750.
2.5159–750. The All Line D.O. L.O. L. T.D. (4007). 5173.

structure regulate telomere position effects in *Saccharomyces cere-* **in fission yeast. Nature** *385***, 744–747.**

Isolation and functional analysis of a *Kluyveromyces lactis* **RAP1 tion. Nature** *392***, 828–831.**

de Lange, T. (1998). Ending up with the right partner. Nature 392,
153–754. Lendvay, T.S., Morris, D.K., Sah, J., Balasubramanian, B., and Lund-
Cilcon E. Poberge M. Circlde B. Phodes D. and Cesser S.M. blad, V. (1996). Se

Natl. Acad. Sci. USA *91***, 8861–8865. interaction in telomeric and** *HML* **silencing in** *Saccharomyces cere-*

Hardy, C.F.J., Sussel, L., and Shore, D. (1992b). A RAP1-interacting Lundblad, V., and Szostak, J.W. (1989). A mutant with a defect in ulation. Genes Dev. *6***, 801–814. Lustig, A.J., Kurtz, S., and Shore, D. (1990). Involvement of the Harlow, E., and Lane, D. (1988). Antibodies, a Laboratory Manual silencer and UAS binding protein RAP1 in regulation of telomere**

McEachern, M.J., and Blackburn, E.H. (1994). A conserved sequence Note Added in Proof motif within the exceptionally diverse telomeric sequences of budding yeasts. Proc. Natl. Acad. Sci. USA *91***, 3453–3457. The article cited herein as "X.-D. Zhu et al., submitted" is now in**

T. (2000). Cell cycle regulated association of Rad50/Mre11/Nbs1 elongation caused by telomerase RNA gene mutations. Nature *376***, with TRF2 and human telomeres. Nat. Genet., in press. 403–409.**

McEachern, M.J., and Hicks, J.B. (1993). Unusually large telomeric repeats in the yeast *Candida albicans***. Mol. Cell. Biol.** *13***, 551–560. Moretti, P., Freeman, K., Coodly, L., and Shore, D. (1994). Evidence**

that a complex of SIR proteins interacts with the silencer and telomere-binding protein RAP1. Genes Dev. *8***, 2257–2269.**

Nakamura, T.M., Cooper, J.P., and Cech, T.R. (1998). Two modes of survival of fission yeast without telomerase. Science *282***, 493–496.**

Nakamura, T.M., Morin, G.B., Chapman, K.B., Weinrich, S.L., Andrews, W.H., Lingner, J., Harley, C.B., and Cech, T.R. (1997). Telomerase catalytic subunit homologs from fission yeast and human. Science *277***, 955–999.**

Nimmo, E.R., Pidoux, A.L., Perry, P.E., and Allshire, R.C. (1998). Defective meiosis in telomere-silencing mutants of *Schizosaccharomyces pombe***. Nature** *392***, 825–828.**

Ogata, K., Morikawa, S., Nakamura, H., Sekikawa, A., Inoue, T., Kanai, H., Sarai, A., Ishii, S., and Nishimura, Y. (1994). Solution structure of a specific DNA complex of the Myb DNA-binding domain with cooperative recognition helices. Cell *79***, 639–648.**

Shore, D., and Nasmyth, K. (1987). Purification and cloning of a DNA binding protein from yeast that binds to both silencer and activator elements. Cell *51***, 721–732.**

Smith., D.L., and Johnson, A.D. (1992) A molecular mechanism for combinatorial control in yeast: MCM1 protein sets the spacing and orientation of the homeodomains of an a**2 dimer. Cell** *68***, 133–142.**

Smith, S., and de Lange, T. (1997). TRF1, a mammalian telomeric protein. Trends Genet. *13***, 21–26.**

Smith, S., Giriat, I., Schmitt, A., and de Lange, T. (1998). Tankyrase, a poly(ADP-ribose) polymerase at human telomeres. Science *282***, 1484–1487.**

Smogorzewska, A., van Steensel, B., Bianchi, A., Oelmann, S., Schaefer, M.R., Schnapp, G., and de Lange, T. (2000). Control of human telomere length by TRF1 and TRF2. Mol. Cell. Biol. *20***, 1659– 1668.**

Spink, K.G., Evans, R.J., and Chambers, A. (2000). Sequence-specific binding of Taz1p dimers to fission yeast telomeric DNA. Nucleic Acids Res. *28***, 527–533.**

van Steensel, B., and de Lange, T. (1997). Control of telomere length by the human telomeric protein TRF1. Nature *385***, 740–743.**

van Steensel, B., Smogorzewska, A., and de Lange, T. (1998). TRF2 protects human telomeres from end-to-end fusions. Cell *92***, 401–413.**

Vassetzky, N.S., Gaden, F., Brun, C., Gasser, S.M., and Gilson, E. (1999). Taz1p and teb1p, two telobox proteins in *Schizosaccharomyces pombe***, recognize different telomere-related DNA sequences. Nucleic Acids Res.** *27***, 4687–4694.**

Wang, S.-S., and Zakian, V. A. (1990). Sequencing of *Saccharomyces* **telomeres cloned using T4 DNA polymerase reveals two domains. Mol. Cell. Biol.** *10***, 4415–4419.**

Wotten, D., and Shore, D. (1997). A novel Rap1p-interacting factor, Rif2p, cooperates with Rif1p to regulate telomere length in *Sacharomyces cerevisiae***. Genes Dev.** *11***, 748–760.**

Zhang, X., Morera, S., Bates, P.A., Whitehead, P.C., Coffer, A.I., Hainbucher, K., Nash, R.A., Sternberg, M.J., Lindahl, T., and Freemont, P.S. (1998). Structure of an XRCC1 BRCT domain: a new protein-protein interaction module. EMBO J. *17***, 6404–6411.**

Zhong, Z., Shiue, L., Kaplan, S., and de Lange, T. (1992). A mammalian factor that binds telomeric TTAGGG repeats in vitro. Mol. Cell. Biol. *13***, 4834–4843.**

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McEachern, M.J., and Blackburn, E.H. (1995). Runaway telomere press: Zhu, X.-D., Kuster, B., Mann, M., Petrini, J.H.J., and de Lange,