Saccharomyces telomeres assume a non-nucleosomal chromatin structure

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The chromatin structures of the telomeric and subtelomeric regions on chromosomal DNA molecules in Saccharomyces cerevisiae were analyzed using micrococcal nuclease and DNase I. The subtelomeric repeats X and Y' were assembled in nucleosomes. However, the terminal tracts of C1.3A repeats were protein protected in a particle larger than a nucleosome herein called a telosome. The proximal boundary of the telosome was a DNase I hypersensitive site. This boundary between the telosome and adjacent nucleosomes was completely accessible to Escherichia coli dam methylase when this enzyme was expressed in yeast, whereas a site 250 bp internal to the telomeric repeats was relatively inaccessible. Telosomes could be cleaved from chromosome ends with nuclease and solubilized as protein-DNA complexes. Immunoprecipitation of chromosomal telosomes with antiserum to the RAP1 protein indicated that RAP1 was one component of isolated telosomes. Thus, the termini of chromosomal DNA molecules in yeast are assembled in a non-nucleosomal structure encompassing the entire terminal C1.3A tract. This structure is separated from adjacent nucleosomes by a region of DNA that is highly accessible to enzymes.

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terminal regions of extrachromosomal ribosomal DNA (rDNA) molecules from *Tetrahymena* (Chiu and Blackburn 1981; Budarf and Blackburn 1986) and slime molds (Edwards and Firtel 1984; Lucchini et al. 1987) have been analyzed, on the basis of nucleosome protection patterns and salt solubility, these telomeric repeats were also found to be assembled in non-nucleosomal chromatin structures.

Specific terminus-binding proteins that copurify with chromosome ends have been isolated from the ciliates *Oxytricha* (Gottschling and Zakian 1986) and *Euplotes* (Price 1990). These proteins bind tenaciously, yet non-covalently, to the native DNA ends, remaining bound even in 2 M salt (Gottschling and Zakian 1986; Price and Cech 1989). Their presence on DNA termini account for only the distal portion of the non-nucleosomal terminal structure defined by chromatin mapping studies, suggesting the presence of additional proteins in the terminal complex (Price and Cech 1987).

Proteins that bind specifically to yeast telomeric repeats in vitro have been identified in *S. cerevisiae* (Berman et al. 1986; Buchman et al. 1988; Liu and Tye 1991). *RAPI*, the gene encoding one of the yeast proteins, has been cloned and shown to be essential for growth (Shore et al. 1986; Buchman et al. 1988; Liu and Tye 1991). *RAPI*, the gene encoding one of the yeast proteins, has been cloned and shown to be essential for growth (Shore et al. 1986; Buchman et al. 1988; Liu and Tye 1991).

**Results**

*X and Y' elements, but not C1,3A repeats, are assembled in nucleosomes*

Micrococcal nuclease (MNase) is a relatively non-sequence-specific endonuclease that is diagnostic for nucleosomes in that it cleaves preferentially in the linker region between nucleosomal core particles (for review, see van Holde 1988). Compared with other organisms, the internucleosomal linker length in *S. cerevisiae* is short, giving an MNase repeat length of ~160 bp (Thomass and Furber 1976; Lohr and Ide 1979).

MNase was used to examine the protection pattern of telomeric sequences [Fig. 1]. Yeast nuclei were isolated from spheroplasts and subjected to MNase digestion. DNA was purified from digested nuclei at different time points and analyzed by Southern blotting. To control for any sequence preference of MNase, a similar time course was performed on naked DNA. The blots of digested chromatin and DNA were hybridized to a C1,3A probe and to probes from conserved regions of the X and Y' elements.

When the MNase chromatin digestions were hybridized with the X or Y' probes, ladders of partial digestion products characteristic of nucleosomes were detected [Fig. 1]. These protection patterns were indistinguishable from profiles obtained by ethidium bromide staining of the gel prior to blotting or by probing with structural genes such as *LEU2* or *TRPI* (data not shown). Thus, the X and Y' sequences, like the bulk of the genome, are assembled in nucleosomes. A different protection pattern was seen when the same blot was probed with the C1,3A probe [Fig. 1]. Instead of nucleosomal-size particles, C1,3A DNA was protected in DNA fragments of a larger and more heterogeneous size than a nucleosome. These data suggest that nucleosomes do not continue out to the very ends of chromosomes in yeast.

**Telomeres possess a DNase I hypersensitive site near the C1,3A-unique DNA junction**

At low concentration, DNase I can be used to identify sequences free of nucleosomes or bound by other non-nucleosomal proteins (for review, see Gross and Garrard 1988). These regions appear as chromatin-specific, DNase I hypersensitive sites. A digestion series on nuclei and naked DNA was performed with DNase I and probed with telomeric sequences [Fig. 2]. In DNase I-digested chromatin, the C1,3A probe hybridized to a heterogeneous smear, corresponding to a population of small DNA fragments averaging 400 bp in size. This smear appeared at very early time points in the digestion when the majority of the DNA was still virtually undigested [Fig. 2]. The fact that the smear persisted for a number of time points without diminishing in size suggested that these DNA sequences were protected in a protein-dependent structure. The early appearance of these small terminal DNA fragments indicated that the proximal boundary of this structure was a hypersensitive site for DNase I. Comparison of the chromatin and naked DNA digestion patterns revealed that the hypersensitive site was chromatin specific [Fig. 2]. When the DNase I-digested chromatin was reprobed with Y' [Fig. 2], X, or single-copy DNA sequences (data not shown), the digestion pattern was similar to that seen for bulk DNA as deduced from the ethidium bromide staining pattern and was thus representative of nucleosomal DNA. Additional hypersensitive sites giving bands at
Figure 1. MNase digestion time courses of nuclei and naked DNA. [Top panels] Sequential probings with C1.3A, X, and Y' of a blot of a 1% agarose gel of an MNase digestion time course of nuclei. DNA was purified from the digested nuclei throughout the time course and loaded in successive lanes on the gel. From left to right, the time points in seconds were 0, 15, 30, 45, 60, 75, 90, 105, 120, 150, 180, 210, 240, 270, 300, 600, 900, and 1200. [Bottom panels] Sequential probings with C1.3A, X, and Y' of a blot of a 1% agarose gel of an MNase digestion time course on purified genomic DNA. The time points in seconds from left to right were 0, 15, 30, 45, 60, 75, 90, 105, 120, 180, 240, 300, 600, 1200, 1200 (2× MNase concentration), 1200 (4×), 1200 (8×), and 1200 (10×). The positions of fragments from the BRL 1-kb ladder are indicated at left.

~6.5 kb and ~3.5 kb (C1.3A) or ~3.2 kb (Y') were apparent in both the C1.3A and Y' hybridizations to the DNase I chromatin time course [Fig. 2]. Bands at 3.2 or 3.5 kb (or both) were also seen with the X and LEU2 probes [data not shown]. In contrast, the ~6.5-kb band was observed only with the C1.3A and Y' probes.

Figure 2. DNase I digestion time courses of nuclei and naked DNA. [Top panels] Sequential probings with C1.3A and Y' of a blot of a 1% agarase gel of a DNase I digestion time course on nuclei. DNA was purified from nuclei throughout the time course and loaded in successive lanes on the gel. From left to right, the time points in seconds were 0, 15, 30, 60, 90, 120, 150, 180, 210, 240, 270, 300, 330, 360, 390, 420, and 480. [Bottom panels] Sequential probings with C1.3A and Y' of a blot of a 1% agarose gel of a DNase I digestion time course on purified genomic DNA. The time points in seconds from left to right were 0, 15, 30, 45, 60, 90, 120, 150, 180, 210, 240, 270, 300, 330, 360, 390, 420, and 480. The positions of fragments from the BRL 1-kb ladder are indicated at left.
Examination of the naked DNA time courses revealed that C1.3A DNA was digested with both nucleases at a rate similar to other sequences. This result indicated that the C1.3A DNA itself was neither hypo- nor hyper-sensitive to these two nucleases (Figs. 1 and 2).

\textbf{Nuclease mapping of a single, modified yeast telomere defined the size of the telosome}

In the yeast strain used here, ~20% of the C1.3A hybridization was to interstitial, rather than terminal, C1.3A tracts. Therefore, the precise protection pattern of terminal C1.3A tracts could not be deduced using C1.3A as a probe. To map the chromatin structure of telomeric C1.3A DNA in more detail, the telomere region of chromosome VII was used to create a uniquely marked telomere. The ~20 kb distal to ADH4 was deleted and replaced with the URA3 gene on chromosome VII (Walton et al. 1986). Integration of this DNA at ADH4 deleted the most distal ~20 kb of chromosome VII-L. The C1.3A sequence on the transforming fragment was elongated in vivo to full length. This figure is not drawn to scale. Positions of probes used in Fig. 4 are indicated.

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Isolated nuclei from a strain carrying the modified VII-L chromosome were treated for varying times with MNase [Fig. 4]. The locations of the nuclease cut sites were mapped by indirect end labeling [Nedospasov and Georgiev 1980, Wu 1980]. Aliquots of DNA purified from different MNase time points were digested with PstI and hybridized to the 100-bp PstI-Rsal fragment of URA3 [Fig. 4C; panels 2 and 3]. The top band in the gels represents the full-length terminal PstI fragment. Bands below this full-length fragment are MNase partial digestion products from the PstI site toward the terminus. The mapped positions of these partial digestion products are presented schematically in Figure 4B. No partial digestion products were detected where cleavage had occurred within the C1.3A repeats. Therefore, either the telomeric repeats were protected but separated from adjacent nucleosomes by a strong site for MNase or the C1.3A repeats were quickly degraded.

To distinguish between these possibilities, the complementary experiment was done. In this case, the telomere was used in place of a restriction site and the

\textbf{Figure 3. Schematic representation of modifications made to chromosome VII-L. The 3' end of the ADH4 gene was used to target the URA3 gene and a short tract of C1.3A sequence to chromosome VII-L. Integration of this DNA at ADH4 deleted the most distal ~20 kb of chromosome VII-L. The C1.3A sequence on the transforming fragment was elongated in vivo to full length. This figure is not drawn to scale. Positions of probes used in Fig. 4 are indicated.}

\textbf{Figure 4. Nuclease mapping of the modified chromosome VII-L telomere. [A] The deduced chromatin structure of the terminal 1.2 kb of chromosome VII-L is shown with nucleosomes represented as spheres and the telosome as an ellipse. [B] A diagram of the mapped region of chromosome VII-L is drawn to scale. The locations of the probes are shown as solid and hatched boxes. Symbols marking the location of nuclease cut sites are [W] MNase chromatin cut site, (vertical lines) MNase naked DNA cut sites, and [O] DNase I chromatin cut sites. DNase I naked DNA cut sites are not shown schematically (see D, panel 3). The numbered lines below the map show the location of partial digestion products observed with MNase digestion of chromatin. The thickness of the lines corresponds to the relative abundance of each partial digestion product: the thick lines were detected as major bands and the thin lines as minor bands. The location of PstI [P] and BamHI [B] sites are indicated. [C] Blots of the MNase and DNase I digestions in nuclei from which locations of the partial digestion products mapped in B were deduced. The numbers 1–8 indicate the partial digestion products that correspond to numbered lines in B. The probe used to hybridize to each blot is indicated above each panel. [Panel 1] MNase-digested nuclei probed with the distal probe. The time points in seconds from left to right were 0, 15, 30, 45, 60, 75, 90, 105, 120. [Panels 2 and 3] MNase digested nuclei probed with the URA3 probe, two different gels which were run to differing extents are shown. In panel 2, partial digestion products 6 and 7 were run off the gel. Time points in panel 2 are identical to those in panel 1. In panel 3, the time points in seconds from left to right are 0, 60, 180, 300, and 600. [Panel 4] DNase I-digested nuclei probed with the URA3 probe. Time points in seconds from left to right are 120, 180, and 300. The DNA samples on all blots probed with URA3 were digested with PstI prior to running the gel. [D] Blots of purified DNA samples digested with MNase and DNase I are shown. Hybridization probes are indicated above each panel; the nuclease used is indicated below the panels. The time points in seconds from left to right are 90, 105, 120, 180, 240, and 300 [panel 1]; 60, 120, 180, and 300 [panel 2]; 120, 150, 180, and 210 [panel 3].}
Figure 4. (See facing page for legend.)
unique sequence directly adjacent to the C1.3A repeats was used to probe for the terminal partial digestion products [Fig. 4C, panel 1]. When the terminal partial digestion products were positioned on the map based on the assumption that the C1.3A tract remained intact, they gave the same image of those products seen with the more internal URA3 probe [Fig. 4B]. The mapped positions shown in Figure 4B were the only interpretation that agreed with the data obtained with the URA3 probe in terms of the relative size and abundance of each pair of digestion products arising from the same cleavage event. Thus, by probing directly for the terminal partial digestion products, it was possible to demonstrate that the terminal C1.3A tract persisted during the digestion time course. The naked DNA digests probed at the distal probe [Fig. 4D, panel 1] showed a smear in the region of the gel where distinct partial digestion products were seen with this DNA in chromatin. This result indicated that the protection of distinct terminal partial digestion products was protein dependent.

The chromatin structure deduced from these data is presented in Figure 4A. Highly phased nucleosomes were detected across the URA3 gene. A structure two and one-half times the size of a nucleosome encompassed the most distal 420 ± 75 bp, including the entire tract of C1.3A DNA and 130 bp of adjacent unique DNA. This non-nucleosomal structure is defined here as a telosome. A similar analysis of the chromosome VII-L modified telomere was done with DNase I [Fig. 4C, panel 4]. When the DNase I partial digestion products were examined with the URA3 probe, the observed cut sites were similar to those mapped with MNase, except the most distal site for DNase I was ~100 bp wide and stronger than the corresponding MNase-cut site [Fig. 4B, open circles above the map]. Experiments using the unique DNA next to the C1.3A repeats to probe DNase I digestions were not informative because this probe fell within the ~100-bp-wide DNase I hypersensitive site (data not shown). The results with DNase I redefined the size of the telosome on chromosome VII-L to 320 ± 75 bp. Presumably, the larger size deduced for the telosome from the analysis of MNase-digested DNA was the result of the sequence preference of MNase within the DNase I hypersensitive region on this modified telomere.

Nucleosomes across the URA3 gene at chromosome VII-L were highly phased with respect to underlying sequence. However, the positions of nucleosomes on the telomeric URA3 gene were identical to the positions of the nucleosomes when URA3 was at its normal location on chromosome V [data not shown] or when URA3 was present on a circular plasmid (Thoma 1986). Thus, the positioning of nucleosomes on URA3 appeared to be unaffected by proximity to the telomere as seen with these nucleases. Cells used for chromatin mapping were grown under conditions that repress expression of telomere-adjacent genes (Gottschling et al. 1990). The fact that the placement of nucleosomes on the URA3 gene was the same when URA3 transcription was repressed by proximity to the telomere [Fig. 4A] as it is when the gene is transcribed (Thoma 1986) suggests that changes in nucleosome positioning are not responsible for telomeric position effect repression.

Chromatin structure of a natural, non-α' telomere revealed that proximity to a telomere has little effect on adjacent chromatin structure

The chromatin structure of a single unmodified telomere was also determined. Although the telomeric regions of most yeast chromosomes contain many kilobases of repetitive DNA, on chromosome I, the 3' end of the PHO11 gene is only ~3.6 kb from the right end of the telomere [Venter and Horz 1989]. PHO12, an acid phosphatase gene virtually identical to PHO11, is located subtelomERICally, ~8.5 bp from one end of chromosome VIII [Steensma et al. 1989]. Venter and Horz 1989]. These PHO genes are part of a large duplication in which the ~20 kb of sequence centromere proximal to PHO11 on chromosome I is also present on chromosome VIII [Steensma et al. 1989]. The restriction maps of DNA distal to the PHO genes on chromosome I and chromosome VIII were similar, indicating that the duplicated region extended to within ~500 bp of the terminus of chromosome I-R [data not shown]. This duplicated region does not contain Y' but does contain X DNA (see Materials and methods).

Cloned fragments containing the PHO11 gene were used to construct two isogenic strains in which a 5-kb region containing either PHO11 or PHO12 was deleted. In both strains, mapping was carried out with a probe from within the deleted region, such that each strain contained a single copy of the probe, located either 1.6 kb from the right end of chromosome I [PHO11] or 6.5 kb from one end of chromosome VIII [PHO12]. MNase and DNase I were used to determine the structure of regions distal to PHO11 and PHO12 in chromatin and in naked DNA (Fig. 5). Chromatin mapping was carried out on cells grown in rich medium, conditions known to repress transcription of these PHO genes [Bostian et al. 1983]. Because of the profound sequence specificity of MNase in these regions [Fig. 5A], it was not possible to assess whether the telomere adjacent DNA was in nucleosomes. However, with both nucleases the chromatin structure of these two regions was very similar, except that with DNase I, the hypersensitive site that defined the boundary of the telosome on chromosome I-R was a weaker site on chromosome VIII (Fig. 5B). Although the chromatin cleavage patterns for the two sequences were very similar, they were offset from each other by ~30 bp [Fig. 5A]. Because this ~30 bp difference was also observed on naked DNA controls and with several restriction enzymes [Fig. 5B and data not shown], it is most easily explained by the presence of a 30-bp segment on chromosome VIII that is missing on chromosome I. Thus, proximity to the telomere created a DNase I hypersensitive site near the junction between the C1.3A repeats and unique-sequence DNA but had no other major effects on the chromatin structure of the adjacent sequences that were revealed by these nucleases [Fig. 5B].
Figure 5. Nuclease mapping of a sequence located both at the chromosome I-R telomere and subtelomERICally on chromosome VIII.

(A) [left] DNA digested with MNase [M] or DNase I [D] in nuclei, as well as HindIII-digested naked DNA [H]. Nuclei were prepared from the yeast strain YS11ΔULΔ to determine the structure on chromosome VIII (lanes 8) and from strain YS12ΔULΔ to determine the structure on chromosome I-R (lanes 1). The time points in seconds were D-8, 300; D-1, 300; M-8, 120; and M-1, 180. [Right] The naked DNA controls for these experiments. Time points in seconds were D-1, 60; D-8, 90; M-1, 120; and M-8, 120. All samples were digested with HindIII prior to loading the gels. Marker lanes are X HaeIII and BRL 1-kb ladder [left], and BRL 1-kb ladder [right]. In both panels, all lanes were from the same gel but were rearranged for easier comparison. (B) Schematic drawings of the regions mapped on chromosome I-R [top] and chromosome VIII [bottom] are presented. Cleavage sites are MNase chromatin cut sites, DNase I chromatin cleavage sites, and MNase naked DNA cut sites. A larger triangle or circle represents a stronger cut site. The location of the DNA used as a hybridization probe in A is shown as a solid rectangle. The speckled region represents the terminal tract of C1.3A DNA. HindIII [H] and KpnI [K] sites are indicated. This map is drawn to scale. The arrows in A and B denote the location of the telomere specific DNase I hypersensitive site.

The boundary between the telosome and adjacent nucleosomes is accessible to Escherichia coli dam methylase in vivo

Constitutive expression of the Escherichia coli dam DNA methylase gene in S. cerevisiae results in the in vivo modification of GATC sequences along the yeast genome. Other than minor effects on mitotic recombinational rates, yeast cells expressing the methylase grow normally [Hoekstra and Malone 1985]. The extent of adenine methylation that occurs in vivo can be measured by comparing the relative cleavage of isolated genomic DNA with restriction enzymes that cleave at the sequence modified by the dam methylase: Cleavage at GATC by Sau3A is insensitive to methylation, whereas cleavage at GATC by DpnI occurs only if the site is methylated on both strands. The persistence of large (>10 kb) DNA fragments in DpnI digestions of genomic DNA from cells expressing the dam methylase suggests that some regions of the yeast genome are inaccessible to the methylase activity [Hoekstra and Malone 1985].
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The accessibility of two GATC sites near the telomere of chromosome VII-L was determined in yeast cells expressing the dam methylase gene. This experiment was carried out in a strain bearing a modification of chromosome VII-L very similar to that depicted in Figure 3 in which the URA3 gene is adjacent to the telomere (Gottschling et al. 1990). In this strain there are two GATC sites, 200 bp apart, near the chromosome VII-L telomere: Site 1 is 50 bp and site 2 is 250 bp internal to the start of the terminal C1.3A tract (Fig. 6A). These experiments were performed on cells growing in rich medium, in which case transcription of URA3 is repressed by its proximity to the telomere (Gottschling et al. 1990).

The GATC site closest to the C1.3A tract (site 1) was cleaved by DpnI in ~100% of the DNA molecules isolated from cells expressing the dam methylase. Thus, site 1 was methylated at virtually all chromosome VII-L telomeres in vivo. In contrast, site 2 was cleaved by DpnI in only ~30% of the DNA molecules, whereas this same site was cleaved by Sau3A in all molecules (Fig. 6B). Thus, site 2, located within the URA3 gene, was methylated in 30% of the chromosome VII-L telomeres. When the URA3 gene was at its normal locus on chromosome V (Fig. 6), or when URA3 was located near the chromosome VII-L telomere in mutant strains that relieve transcriptional repression of telomeric genes (Aparicio et al. 1991), the GATC sequence within the URA3 gene (site 2) was 100% methylated (Gottschling 1992). In contrast, the GATC sequence at site 1 located within the DNase I hypersensitive region that links the telosome to adjacent nucleosomal chromatin was completely accessible to the methylase in vivo under all conditions and in all strains tested (Fig. 6 and data not shown). These data showed that the DNA at the proximal boundary of the telosome is highly accessible to enzymes not only in isolated nuclei but also in vivo.

Solubilized telosomes retain bound protein

To investigate the nature of the protein–DNA interactions at telomeres, methods were developed for solubilizing telosomes. Telosomes were released from chromosomes by digestion with MNase or DNase I, or from the modified version of chromosome VII-L by digestion with a restriction enzyme with a recognition site in the nuclease-accessible region. Nuclei were then disrupted, and insoluble material was removed by centrifugation to yield a chromatin supernatant. The chromatin supernatant contained ~80% of the DNA that hybridized to C1.3A. The fraction of nucleosomal DNA in the chromatin supernatant ranged from 10% to 80%, depending on which nuclease was used, and was inversely proportional to the size of the digestion products.

Chromatin supernatants prepared from MNase-digested nuclei were analyzed by agarose gel electrophoresis and Southern blotting, with and without prior digestion by proteinase K (Fig. 7A). Probing of these blots with C1.3A revealed that the heterogeneously sized telomeric fragments were shifted quantitatively in mobility in a protein-dependent manner. This result demonstrated that the solubilized telosomes were protein bound. In different experiments, the telosome migrated to the approximate position of either the 1.6- or the 1.0-kb DNA molecular weight markers.

Proteins in yeast extracts can bind to exogenously added telomeric DNA in vitro to yield a specific gel shift (Berman et al. 1986). Competition experiments were carried out to determine whether the proteins responsible

Figure 6. In vivo methylation of the region near the telomere of the modified chromosome VII-L. [A] The diagram represents the structure near the telomere of chromosome VII-L after modification by insertion of the URA3 gene (see Fig. 3). The thick shaded region represents the terminal C1.3A tract, the solid rectangle indicates the sequence used for hybridization in B, the open rectangle represents the URA3 gene, and 1 and 2 mark the locations of GATC sites. Site 1 is also a BamHI site. The lower portion of the diagram depicts the expected fragments, A through E, detected by the hybridization probe if DpnI cleavage does or does not occur at sites 1 or 2. B DNA was isolated from a strain in which the E. coli dam methylase gene is expressed in vivo and which contains a mutation in the URA3 gene (strain UCC1023, marked ura3-52) or from the same strain after modification of chromosome VII-L (strain UCC1021, marked URA3-TEL + ura3-52). DNA was digested with HindIII alone (marked 0) or with HindIII plus either BamHI, DpnI, or Sau3A and run on an agarose gel. The DNA was hybridized to the probe shown in A. The open arrowhead indicates the HindIII band from the ura3-52 locus, the closed arrowhead indicates the band from the same locus after digestion with Sau3A. Fragments A–E are derived from the telomere region of chromosome VII-L as shown schematically in A. The marker lane is the 1-kb BRL ladder.
petitor suggested that this complex was formed prior to disruption of the nuclei, providing additional evidence that telomeric sequences are protein bound in vivo. The increased mobility of the shifted complex that was observed in the presence of competitor from ~1.6 kb to ~1.0 kb suggested that some protein or proteins were competed away. In many experiments done in the absence of competitor (including the one shown in Fig. 7A), telosomes migrated at ~1.0 kb. Thus, the component of the soluble telosomes that was competed for in this experiment did not appear to copurify with telosomes consistently, suggesting that it was not an integral part of the protein–DNA complex.

The RAPI protein interacts with yeast telomeres in vitro (Buchman et al. 1988) and in vivo (Conrad et al. 1990, Lustig et al. 1990; Susel and Shore 1991). To determine whether RAPI protein is present in telosomes, RAPI antiserum was added to a chromatin supernatant obtained from MNase-digested nuclei. In these experiments, 30–50% of the telosomes were specifically precipitated by the RAPI antibody (Fig. 8A, left panel). In contrast, 2-μm DNA, which is nucleosomal (Livingston and Hahne 1979; Nelson and Fangman 1979), was not precipitated by the antiserum (Fig. 8A, right panel). No precipitation of telomeric fragments was observed with the preimmune sera (data not shown). This experiment was repeated in the presence of 10- and 100-fold molar excess of C1.3A DNA. In the presence of 100-fold molar excess of C1.3A DNA, the immunoprecipitation of the chromosome VII-L telomeric fragment was reduced about twofold (Fig. 8B). The fact that competition with the specific competitor was not complete suggested that RAPI protein was associated with the telosome in nuclei prior to the addition of the specific competitor, indicating that the RAPI protein is a component of telosomes in vivo. The partial competition for the RAPI binding to the telosome could account for the change in mobility of the protein bound telosomes seen with addition of competitor (Fig. 7B).

Discussion

Most yeast DNA sequences, including the subtelomeric repeats X and Y' (Fig. 1), are assembled in nucleosomes. In contrast, the C1.3A repeats at the very ends of yeast chromosomal DNA molecules are assembled in a non-nucleosomal chromatin structure, herein called a telosome. The conclusion that the terminal C1.3A repeats are in telosomes comes from studies using two nucleases, MNase and DNase I, and from examination of the protection patterns of both total C1.3A DNA (Figs. 1 and 2) and two individual telomeres, a modified form of the chromosome VII-L telomere (Fig. 4) and the natural end of chromosome I-R (Fig. 5). This conclusion is also supported by gel-retardation studies indicating that C1.3A repeats were protein associated prior to disruption of isolated nuclei (Figs. 7 and 8).

Telosomes were found to differ from nucleosomes in several ways. First, telosomes contained about twice as

Figure 7. Gel mobility-shift assays on isolated telomeric chromatin. (A) Aliquots of chromatin supernatants prepared from MNase-treated nuclei either were loaded directly onto agarose gels containing ethidium bromide [−] or treated with SDS and proteinase K prior to loading [+] (+). Two samples each without [−] and with [+] proteinase K treatment are shown. Molecular weight markers are BRL 1-kb ladder. This blot is hybridized with the C1.3A probe. (B) Gel mobility-shift assays of chromatin supernatants from MNase-treated nuclei with 0, 10×, and 100× molar excess of C1.3A competitor DNA added to the nuclei prior to lysis. Duplicate lanes with [+] and without [−] treatment with SDS and proteinase K are shown. The amount of competitor added to each lane is indicated below the panel. This blot is probed with the unique sequence DNA next to the chromosome VII-L telomere (solid box, Fig. 4B).

for the gel mobility-shift displayed by telosomes had associated with C1.3A DNA in vivo or in vitro. Gel mobility-shift experiments were performed in the presence of 10- or 100-fold molar excess of cloned C1.3A DNA that was added to nuclei prior to their disruption. Because the C1.3A competitor fragments were similar in length to endogenous telomeres, a C1.3A probe could not be used to analyze these gels. Instead, the competition blots were probed with the unique DNA adjacent to the C1.3A repeats on the modified chromosome VII-L telomere (Fig. 4). Although addition of specific competitor increased the relative mobility of the protein-bound telosomes, 100% of the detectable chromosome VII-L telosomes still displayed a protein-dependent gel shift in the presence of a 100-fold molar excess of C1.3A DNA (Fig. 7B). When this same blot was probed with C1.3A, only a small fraction of the competitor C1.3A fragments were shifted in mobility by in vitro protein binding (data not shown), indicating that this experiment was done under conditions where competitor DNA was in excess over free protein. The fact that some of the telosome-associated proteins were not competed with the specific complemen-

Figure 7.
Figure 8  Immunoprecipitations of chromatin supernatants with anti-RAP1 antiserum. (A) Chromatin supernatants were prepared from nuclei digested with MNase for 2 or 4 min and immunoprecipitated with a RAP1 antiserum. DNAs were purified from the immunosupernatant [S] and the immunopellet [P], subjected to electrophoresis in a 1% agarose gel, and blotted. The blot was probed sequentially with C1.3A [left] and 2 μm of DNA [right]. The molecular weight markers are BRL 1-kb ladder. (B) Chromatin supernatants were prepared from BamHI-digested nuclei with 0, 10×, and 100× molar excess C1.3A competitor DNA. Chromatin supernatants were immunoprecipitated, and DNAs were purified and prepared as in A. The amount of C1.3A competitor DNA is indicated below the panel. The blot was probed with the unique DNA adjacent to the C1.3A tract on the modified chromosome VII-L (solid box, Fig. 4B). Molecular weight markers are BRL 1-kb ladder. This blot was quantified by storage phosphor imaging (Johnston et al. 1990).

much DNA as nucleosomes (Fig. 4). Second, unlike nucleosomes, telosomes contained a variable amount of DNA (Figs. 1, 2, 4, and 5). Telosomes protected the most distal 245–395 bp of DNA, starting from a point ~50 bp internal to the start of the simple repeats and encompassing the entire terminal tract of C1.3A repeats (Fig. 4A). Because the lengths of these terminal tracts vary from telomere to telomere, telosomes from different chromosomes can differ in size by as much as 150 bp of DNA (Figs. 1, 2, and 4). Additional experiments with soluble telosomes indicated that at least some telosomal proteins were not displaced by intercalation of ethidium bromide (J.H.Wright and V.A. Zakian, unpubl.), in contrast, nucleosomes are dissociated by ethidium bromide (McMurray and van Holde 1986). Although these data do not preclude the possibility that histones participate in forming telosomes, they do demonstrate that conventional nucleosomes do not continue out to the very ends of chromosomal DNA molecules.

In contrast to internucleosomal linker regions, the region that links the telosome to adjacent nucleosomal DNA contains a strong hypersensitive site for DNase I (Figs. 2, 4, and 5). The DNase I hypersensitivity of these sites is protein dependent in that the C1.3A unique DNA boundary was not a preferred site for DNase I in naked DNA (Figs. 2, 4, and 5). Proximity to the telomere is the cause of this hypersensitivity. This conclusion was reached by comparing the digestion patterns of very similar DNA sequences located telomerically and subtelomerically on two different chromosomes (Fig. 5). The only major difference in the deduced chromatin structure of a 1.3-kb region present on both chromosomes was a strong DNase I hypersensitive site that was observed only on chromosome I where this sequence was adjacent to a telosome (marked by an arrow in Fig. 5A). Another hypersensitive site was observed with both the C1.3A and Y' probes, generating a band at ~6.5 kb [Fig. 2]. The size of this DNA fragment was approximately the same as the unit length of the Y' element. This result is consistent with the possibility that Y' junctions are also hypersensitive to DNase I. In yeast, DNase I hypersensitive sites are found flanking centromeres (Bloom and Carbon 1982), at ARSs (Thoma et al. 1984; Brown et al. 1991), and at other sites bound by nonhistone proteins such as at the GAL1.10 upstream activating sequence [UAS] (Lohr 1984). Thus, DNase I hypersensitivity at telomeres and at other genomic sites may mark transition points between different chromatin structures.

A GATC site within the region containing the DNase I hypersensitive site on the modified form of chromosome VII-L was 100% accessible in vivo to methylation by E. coli dam methylase when the enzyme was expressed in yeast. In contrast, a GATC site 200 bp internal to this accessible site, located within the URA3 gene, was relatively inaccessible to the methylase (Fig. 6). In fact, this site was less accessible to dam methylase at the telomere than when it was located at internal sites on the chromosome (Fig. 6). When the nucleosome–telosome linker region contained a recognition site for BamHI, the site was BamHI accessible in nuclei at >80% of telomeres (J.H. Wright and V.A. Zakian, unpubl.). The accessibility of this linker region to in vivo methylation on both DNA strands and to digestion by DNase I, MNase, and BamHI in nuclei suggests that this region is not protein associated or that it protrudes from the protein–DNA complex.

Because the telosome–nucleosome linker region is nucleosome-accessible, telosomes can be released from chromosomes as protein–DNA complexes (Figs. 7 and 8). The RAP1 protein was identified as one component of solubilized telosomes (Fig. 8), corroborating our previous results demonstrating that RAP1 is associated with isolated termini from a short, multicopy linear plasmid (Conrad et al. 1990). Telomeres in yeast and other organisms have been shown to associate with the synaptonemal complex in meiosis (Moens and Pearlman 1990) and with the nuclear envelope in interphase nuclei (e.g., Agard and Sadat 1983; Rawlins et al. 1991). In addition to
its binding to telomeres, the RAP1 protein fractionates with the nuclear scaffold and can mediate loop formation between noncontiguous RAP1-binding sites in vitro (Hofmann et al. 1989; Cardenas et al. 1990). Moreover, RAP1 protein has been shown by immunofluorescence with the termini of metiotic chromosomes [F. Klein and S. Gasser, pers. comm.]. Thus, RAP1 is a good candidate for a protein that mediates telomere associations both with other telomeres and with the nuclear scaffold. Although 30–50% of the telomeres had bound RAP1 as shown by specific immunoprecipitation with a RAP1 antiserum [Fig. 8], only a small fraction of the RAP1 protein copurified with telomeric chromatin (J.H. Wright and V.A. Zakian, unpubl.). Thus, most of the RAP1 protein is either free in the cell, associated with some of the many other RAP1-binding sites in yeast DNA (Buchman et al. 1988), or sequestered in structures such as the nuclear scaffold (Cardenas et al. 1990).

Some of the telosomal proteins detected in the gel mobility-shift assay were not competed away from the endogenous telomeric DNA by 100-fold molar excess of cloned C1.3A DNA added prior to nuclear lysis [Fig. 7B]. This result is consistent with the interpretation that the complex was formed before the competitor DNA was added, that is, in vivo. The salt-stable terminus-binding proteins also do not readily exchange with added telomeric DNA ends (Gottschling and Zakian 1986; Raghuraman and Cech 1989). The low exchange rate exhibited by some of the yeast telosomal proteins suggests that the yeast telomere may contain analogs of the Oxystricha terminus-binding proteins. This possibility is strengthened by preliminary data indicating that the solubilized telosomes from yeast are more salt stable than nucleosomes [J.H. Wright and V.A. Zakian, unpubl.].

In summary, we analyzed the chromatin structures of the X and Y' elements found in subtelomeric regions of yeast chromosomes and of the simple repeats at the very ends of chromosomal DNA molecules. X and Y' are organized in nucleosomes, as is the URA3 gene when it is moved next to a yeast telomere. Proximity to the telomere has very little effect on the chromatin structure of adjacent DNA sequences as seen with these nucleases: URA3 has the same chromatin structure at the telomere as at other sites in the genome, and a region 1.6 kb from the end of chromosome I has the same chromatin structure as a very similar region 6.5 kb from the end of chromosome VIII. The C1.3A repeats at the very ends of yeast chromosome are organized into telosomes that can be released as protein-DNA complexes by the actions of several nucleases. RAP1 protein is one component of soluble telosomes. Experiments similar to those described here for RAP1 may be carried out to determine whether other proteins that bind to yeast telomeric repeats in vitro [Liu and Tye 1991] are present in the telosome. It will also be of interest to examine the structure of telosomes in mutant strains [e.g., see Lustig and Petes 1986; Lundblad and Szostak 1989, Sussel and Shore 1990] and under conditions [Runge and Zakian 1989, Conrad et al. 1990] where telomere structure or function is altered.

Materials and methods

Yeasts strains

Most experiments were carried out in strain JW2 [MATa lys5 ura3-51 hisG leu2-3,112 trpl1::hisG ade2 ade5 can1 cyh2 prb1::hisG]. To make JW2, the protease B gene PRB1 [Moehle et al. 1987] was deleted and replaced with the bacterial hisG sequence in the yeast strain DM30 [Runge et al. 1991] by use of previously published methods [Alani et al. 1987]; URA3 was then inserted on the left telomere of chromosome VII [Gottschling et al. 1990] as diagramed in Figure 3. The plasmid used to modify chromosome VII-L was similar to the VII-L URA3-TEL plasmid described in Gottschling et al. [1990], with an additional 60 bp of sequence between the URA3 gene and the C1.3A DNA as described in Conrad et al. [1990]. This 60 bp contains a copy of the lac operator, as well as a concatamer of BamHI sites. All yeast transformations were done according to Ito et al. [1983].

The strains used for analysis of the telomeric regions of chromosomes I and VIII were, respectively, YS12AUL and YS12AUL. These strains were derived from YS18-2 [MATa his3-11 his3-15 leu2-3 leu2-112 ura3-432 ura3-601 can1 pho10:URA3] obtained from W. Horz; YS18-2 was modified by deleting a 5-kb EcoRI fragment containing PHO12, to produce YS12AUL, or containing PHO11, to produce YS11AUL. The PHO11 and PHO12 deletions were obtained by transforming YS18-2 with a DNA fragment containing the HIS3 gene flanked by short stretches of DNA found at both the PHO11 and PHO12 loci. The fragment used for the transformation was isolated from pPH011, a plasmid with an insert containing a 5-kb EcoRI fragment of the PHO11 gene [obtained from W. Horz, Venter and Horz 1989]. A 4.6-kb Dral–KpnI fragment within the 5-kb EcoRI insert in pPH011 was replaced with a 1.8-kb BamHI fragment containing the HIS3 gene. The resulting 2.2-kb EcoRI fragment was used to transform YS18-2. Transformants were screened by Southern blotting for the PHO11::HIS3 deletion/insertion or the PHO12::HIS3 deletion/insertion. These strains also contained a third, unidentified locus that hybridized to the probe used in the chromatin mapping analyses. This cross-hybridizing locus was deleted in both strains by transformation with a 2.7-kb fragment containing a 2.2-kb Xhol–SalI fragment of the LEU2 gene flanked by 230 bp BclI–HindIII and 200 bp KpnI–EcoRI fragments subcloned from pPH011. The result of these manipulations was the construction of two strains in which the 230-bp HindIII–KpnI fragment of pPH011 hybridized uniquely to either the PHO11 or PHO12 locus. The distance between PHO11 and the end of the chromosome in YS12AUL is not large enough to contain a Y' element. The lack of Y' on chromosome I-R was confirmed directly in YS18-2 [Venter and Horz 1989]. Two X probes hybridized to the terminal 10 kb of chromosome I-R in a strain called AB972 [de Steenstra et al. 1989, H.Y. Steensma, pers. comm.]. In both YS12AUL and YS11AUL, the right end of chromosome I had the same restriction map as in AB972. Thus, the right end of chromosome I and the PHO12 end of chromosome VIII in JW2 both contain at least some sequences from the X element.

The yeast strains used for the dam methylase experiments were UCC1023 [MATa ura3-52 lys2-801:dam + lys2 ade2-101 trpl1::hisG 2 Dam+ URA3-TEL] and UCC1021 [UCC1023 with adh4::URA3-TEL]. In these strains the E. coli dam methylase gene was integrated at the LYS2 gene on yeast chromosome II-R as described [Gottschling 1992], where it is expressed constitutively, presumably from a fortuitous promoter within the bacterial DNA. The dam methylase gene was isolated as a 1.5-kb HindIII–PvuII fragment from pMFH1 [Hoeckstra and Malone 1985].
Preparation of yeast chromatin and DNA

Nuclease digestion of chromatin was carried out in nuclei isolated by modification of methods described in Kenna et al. (1989). Cells were grown to 2 × 10^6–1 × 10^7 cells/ml at 30°C in YEPD (Zakian and Scott 1982). Cells were washed once with water, once with SCE-60 (1 M sorbitol, 0.1 M sodium citrate, 60 mM EDTA at pH 7.5), resuspended in 25 ml of SCE-60 with 14 mM β-mercaptoethanol and 0.5 mg/ml Zymolyase-100T (ICN), and incubated for 30 min at 30°C. Spheroplasts were collected by centrifugation and washed with 40 ml of cold SPC (1 M sorbitol, 20 mM PIPES at pH 6.3, 10 mM CaCl₂, 1 mM PMSF, 1 mM iodoacetic acid). The remaining steps of the procedure were done at 4°C. The spheroplast pellet was resuspended in 2 ml of SPC, added to 40 ml of FPC (9% Ficoll, 20 mM PIPES at pH 6.3, 10 mM CaCl₂, 1 mM PMSF, 1 mM iodoacetic acid), and mixed by inversion. Cell lysis was allowed to occur for 15 min on ice. To pellet nuclei, the lysate was centrifuged at 13,000 rpm in a Sorvall SS-34 rotor for 20 min at 4°C. DNA was isolated for the naked DNA controls as described in Runge et al. (1989).

Nuclease assays

For chromatin digestions, nuclear pellets were washed in ~10 volumes of PC (50 mM PIPES at pH 6.3, 0.1 mM CaCl₂, 75 mM NaCl, 6 mM MgCl₂) and resuspended in 7.5 × 10^6 cells/ml in PC + 6 mM CaCl₂ for MNase assays, and in PC for DNase I assays. Nuclei were aliquoted into 1.5-ml microtubes at 200 xg/ml of nuclei in 1.5-ml microtubes. Tubes were shaken at 4°C for 30 min in an Eppendorf model 5432 tube mixer. The bottoms of the tubes were punctured with a needle, and the liquid spun into new tubes. The disrupted nuclei were centrifuged at 10,000g for 5 min. The supernatant contained the soluble chromatin. This method is an adaptation of procedures developed by M. Saunders and K. Bloom (pers. comm.). For the BamHI-digested chromatin supernatants, no other nuclease digestion was done; BamHI digestion was done at 4°C during the nuclear lysis with glass beads with 500 units of enzyme (NEB). The C1.3A competitor DNA was a 7.5-kb Sall fragment from the plasmid YLPF (R. Wellinger, A. Wolf, and V.A. Zakian, in prep.), which contains two ~300-bp tracts of C1.3A DNA cloned from natural yeast telomeres by the methods described in Wang and Zakian (1990). The competitor was added to the nuclei prior to their disruption. Each ~300-bp tract was considered the molar equivalent of a yeast telomere.

Assay for in vivo methylation

DNA was isolated from UCC1021 or UCC1023 cells that were grown to stationary phase in YEPD. DNA was digested by the appropriate restriction enzymes, separated by electrophoresis in a marine gel apparatus. All Southern blots were done according to methods described in Kenna et al. (1990). The probe used to map chromosome I-R and chromosome VII-L telomeres was a pair of complementary 60-bp oligonucleotides containing the lac operator (Conrad et al. 1990). The probe used to map chromosome 1-R and chromosome VIII was a 230-bp HindIII–KpnI probe subcloned from pPHOU (Jayaram et al. 1983). The URA3 probe was a 100-bp PstI–RsaI fragment from the 5′ end of the URA3 gene (Rose et al. 1985). The probe for the unique DNA immediately adjacent to the C1.3A tract on the modified chromosome VII-L telomere was a pair of complementary 60-bp oligonucleotides containing the lac operator (Conrad et al. 1990). The probe used to map chromosome I-R and chromosome VIII was a 230-bp HindIII–KpnI probe subcloned from pPHOU (Venter and Horz 1989). The 2-µm probe was a 1.3-bp HindIII fragment isolated from recombinant DNA plasmid CV20 (Jayaram et al. 1983).

Gel mobility-shift assays

Chromatin supernatants were subdivided into aliquots in duplicate at 40 µl per tube on ice. Four microliters of 1% SDS and 10 µg of proteinase K were added to one, and then one or both was incubated at 37°C for 20 min. Gel dye and glycerol were added just prior to loading the gel. The gels were run at 10.7 V/cm for 3 hr using the electrophoresis conditions described above.
Immunoprecipitations
Chromatin supernatant (250 μl) was mixed with 16 μl of RAPI antiserum or with preimmune serum (Conrad et al. 1990) and incubated on ice. Protein A-Sepharose (50 μl) was added in two aliquots for a total incubation time of 60 min. The beads were washed four times with binding buffer. Material retained by the beads was eluted with 1% SDS at 65°C. DNAs from both supernatant and pellet fractions were purified for DNA isolated for naked DNA controls, except that 10 μg of tRNA was added as carrier prior to ethanol precipitation. The samples were suspended in TE + 2 μg of RNase A and subjected to standard gel electrophoresis and Southern blotting.

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