Association of the RENT complex with nontranscribed and coding regions of rDNA and a regional requirement for the replication fork block protein Fob1 in rDNA silencing

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Silencing within the yeast rDNA repeats inhibits hyperrecombination, represses transcription from foreign promoters, and extends replicative life span. rDNA silencing is mediated by a Sir2-containing complex called RENT (regulator of nucleolar silencing and telophase exit). We show that the Net1 (also called Cfi1) and Sir2 subunits of RENT localize primarily to two distinct regions within rDNA: in one of the nontranscribed spacers (NTS1) and around the Pol I promoter, extending into the 35S rRNA coding region. Binding to NTS1 overlaps the recombination hotspot and replication fork barrier elements, which have been shown previously to require the Fob1 protein for their activities. In cells lacking Fob1, silencing and the association of RENT subunits are abolished specifically at NTS1, while silencing and association at the Pol I promoter region are unaffected or increased. We find that Net1 and Sir2 are physically associated with Fob1 and subunits of RNA polymerase I. Together with the localization data, these results suggest the existence of two distinct modes for the recruitment of the RENT complex to rDNA and reveal a role for Fob1 in rDNA silencing and in the recruitment of the RENT complex. Furthermore, the Fob1-dependent associations of Net1 and Sir2 with the recombination hotspot region strongly suggest that Sir2 acts directly at this region to carry out its inhibitory effect on rDNA recombination and accelerated aging.

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results in the expansion or contraction of the array [Kobayashi and Horiuchi 1996; Kobayashi et al. 1998; Johzuka and Horiuchi 2002]. Furthermore, FOB1 is required for hotspot (HOT1) activity, a phenomenon in which specific rDNA sequences can stimulate homologous recombination when placed outside of the array [Lin and Keil 1991]. Significantly, the cis-element sequences required for establishing the RFB and stimulating recombination are overlapping and are found within a region originally identified as an enhancer of RNA polymerase I [Pol I] transcription outside the rDNA array [Elion and Warner 1984; Wai et al. 2001]. How Fob1 functions to generate an RFB or stimulate recombination is not known.

Several lines of evidence suggest that recombination in the yeast rDNA repeats is also negatively regulated through a mechanism that resembles heterochromatic gene silencing and is referred to as rDNA silencing. First, recombination levels in S. cerevisiae are down-regulated by Sir2 [Gotlib and Esposito 1989], an NAD-dependent deacetylase originally identified for its role in chromatin silencing at the mating-type loci and telomeres [for review, see Moazed 2001; Rusche et al. 2003]. Second, Sir2 is required for silencing of Ty1 transposition and transcription of Pol II-dependent reporter genes that are inserted within rDNA [Bryk et al. 1997; Fritze et al. 1997; Smith and Boeke 1997]. Finally, the altered sensitivity of rDNA to micrococcal nuclease and dam methyltransferase in sirΔ cells, as well as loss of rDNA silencing in strains with histone mutations, supports the idea that a Sir2-based silencing mechanism inhibits rDNA recombination by altering chromatin structure [Fritze et al. 1997; Bryk et al. 2002; Hoppe et al. 2002; Park et al. 2002].

Regulation of recombination at rDNA by Fob1 and Sir2 is also a major determinant of budding yeast life span. The accumulation of extrachromosomal rDNA circles (ERCs) excised from the rDNA array can lead to premature cellular senescence in S. cerevisiae [Sinclair and Guarente 1997]. Accordingly, loss of silencing in sirΔ cells results in an increased rate of Fob1-dependent recombination and ERC accumulation and reduces average life span, whereas increasing the dosage of Sir2 suppresses recombination and prolongs average life span [Kaeberlein et al. 1999]. In contrast, FOB1 deletion cells display the opposite aging phenotype. The absence of FOB1 reduces recombination and the formation of ERCs, extends average life span, and as expected, suppresses premature aging in cells lacking Sir2 [Defossez et al. 1999].

In yeast, silencing is best understood at the silent mating-type loci and telomeric regions [Rine and Herskowitz 1987; Gottschling et al. 1990]. Initiation of silencing at these regions involves the association of DNA-binding proteins with cis-acting silencer elements. The silencer-binding factors then recruit a second class of proteins to DNA to form the SIR complex, consisting of Sir2, Sir3, and Sir4 proteins [for review, see Rusche et al. 2003]. As mentioned above, Sir2 is an NAD-dependent deacetylase [Imai et al. 2000; Landry et al. 2000; Smith et al. 2000], and its activity is necessary for the spreading of silencing complexes along chromat via interactions with the N termini of histones H3 and H4 [Braunstein et al. 1993; Hecht et al. 1996; Strahl-Bolsinger et al. 1997; Hoppe et al. 2002; Luo et al. 2002; Rusche et al. 2002]. Sir2 is also the only SIR protein required for rDNA silencing [Bryk et al. 1997; Fritze et al. 1997; Smith and Boeke 1997], and it is part of an rDNA silencing complex called RENT [regulator of nucleolar silencing and telophase exit; Shou et al. 1999; Straight et al. 1999; Visintin et al. 1999]. In addition to Sir2, RENT contains Net1 and Cdc14 [Shou et al. 1999; Visintin et al. 1999]. Net1 is required for rDNA silencing and localization of Sir2 to rDNA [Straight et al. 1999]. Net1 can also associate with Pol I [Shou et al. 2001], but it is unknown if the entire RENT complex, including Sir2, associates with Pol I. Cdc14 is a phosphatase that regulates exit from mitosis [Shou et al. 1999; Visintin et al. 1999], but whether it plays a role in silencing is not known. Moreover, specific silencer elements or DNA-binding proteins that recruit silencing complexes to rDNA have not been described.

To investigate how silencing is initiated at rDNA, we performed a high-resolution mapping of Net1 and Sir2, two subunits of the RENT silencing complex, along the entire 9.1-kb length of an rDNA repeat by chromatin immunoprecipitation [ChIP]. Each rDNA unit yields a 35S precursor rRNA and a 5S rRNA, separated by two nontranscribed spacers, NTS1 and NTS2 (see Fig. 1A). Our data show that both silencing proteins are associated primarily with two regions: one region within NTS1 and a second that overlaps the Pol I promoter and part of the 35S rRNA gene. The NTS1 region associated with silencing proteins includes sequences necessary for FOB1-dependent replication fork block and recombination activities. Surprisingly, we find that FOB1 is required for rDNA silencing at NTS1, and we show that Fob1 is primarily associated with this region of rDNA. In contrast, lower levels of Fob1 are localized to the Pol I promoter/35S region, and deletion of FOB1 has no effect on silencing at this location. Consistent with these observations, in fob1Δ cells, we detect dramatically reduced associations of Net1 and Sir2 with NTS1 but unaffected or increased associations with the Pol I promoter/35S region. Both Fob1 and Pol I physically interact with the RENT complex, suggesting two distinct pathways recruit the RENT complex to rDNA.

Results

Net1 and Sir2 are preferentially associated with two regions within rDNA

The structure of a 9.1-kb rDNA repeat unit and important functional elements are shown in Figure 1A. The association of Net1 or Sir2 with rDNA by ChIP has been studied previously using a limited number of sites [ranging from 1 to 7; Gotta et al. 1997; Straight et al. 1999; Armstrong et al. 2002; Bryk et al. 2002; Buck et al. 2002; Hoppe et al. 2002]. Because Sir2 associates preferentially with a few sites in the nontranscribed spacer (NTS) as compared with a few sites within the 35S coding region
(Gotta et al. 1997; Hoppe et al. 2002), it might be expected that silencing complexes would be found mainly at the NTS. To obtain a more comprehensive picture of where Net1 and Sir2 are associated with rDNA, we designed a panel of 68 primers to amplify fragments of ~0.25 kb in length that spanned an rDNA repeat [Fig. 1A].

We constructed yeast strains in which the endogenous copy of the NET1 or SIR2 gene was modified to encode a protein with the TAP tag at its C terminus [Net1–TAP or Sir2–TAP]. The TAP tag is a dual epitope tag consisting of a calmodulin-binding peptide separated from two Protein A repeats by a TEV-protease cleavage site. Both NET1–TAP and SIR2–TAP strains exhibited the same levels of rDNA silencing as the parental untagged strain, suggesting that the modified proteins were fully functional [Supplementary Fig. 1]. Cells were cross-linked with formaldehyde, and Net1–TAP or Sir2–TAP was immunoprecipitated from extracts containing sheared chromatin using an IgG resin. Whole-cell extract chromatin [WCE] or immunoprecipitated chromatin [IP] from untagged or TAP-tagged strains was used as template DNA for quantitative PCR, using the panel of primers shown in Figure 1A.

We generated a graphical representation of Net1 and Sir2 association across an rDNA repeat [Fig. 1B] by using the ratio of IP material to WCE [input] material to calculate relative fold enrichment values for each DNA fragment. Background binding was defined by immunoprecipitation of rDNA fragments from an untagged strain and immunoprecipitation of DNA fragments from CUP1, a repetitive, nonsilenced locus. Unexpectedly, we found that Net1 and Sir2 were not found in a single peak spanning the NTS region. Instead, the data showed two major peaks that overlapped most of NTS1 but only part of NTS2 [Fig. 1B]. Notably, the NTS1 peak coincided with the replication fork block region [RFB; Fig. 1B]. This region contains a polar replication fork block and a number of cis-elements required for FOB1-dependent rDNA recombination [Lin and Keil 1991; Kobayashi and Horiuchi 1996; Kobayashi et al. 1998; Jozzuka and Horiuchi 2002]. The NTS2 peak overlapped the Pol I transcription initiation region [TIR; Fig. 1B], but the majority of the peak was in fact spread toward the 18S region of the transcribed 35S rRNA gene. We also observed a much smaller peak at the 5S rRNA gene. The relative associations of Net1 and Sir2 with rDNA fragments closely mirrored each other, but more material was consistently immunoprecipitated by Net1–TAP than Sir2–TAP. Because levels of Net1–TAP were higher than those of Sir2–TAP [Fig. 1G, lanes 2,3], and Sir2 is also required for silencing at other loci, this observation may reflect higher levels of Net1 association with rDNA. Additionally, whereas Net1 is associated with rDNA throughout the cell cycle, Sir2 is partially released toward the end of mitosis [Straight et al. 1999]. Finally, differences in cross-linking efficiency can also explain the difference in association of rDNA fragments with Net1 and Sir2 in our experiments.

Examples of the ChIP data used to obtain the graph in Figure 1B are shown in Figure 1C–E. The enrichment of DNA regions corresponding to most of the 25S rRNA by Net1 or Sir2 was near the background level of binding [Fig. 1C]. Similarly, DNA from CUP1 was not significantly enriched in Net1–TAP and Sir2–TAP immunoprecipitations [Fig. 1C]. In contrast, high levels of DNA fragments from NTS1 and NTS2/18S regions were immunoprecipitated by Net1–TAP and Sir2–TAP [Fig. 1D,E]. As another way of quantifying relative binding, we performed multiplex PCR with primers for both rDNA and CUP1, using CUP1 as an internal control for calculating relative fold enrichment. Examples of the data and the calculated relative fold enrichment are shown in Figure 1F and are consistent with the two peaks of Net1/Sir2 association in Figure 1B.

Previous ChIP experiments have shown that in sir2Δ cells, the acetylation levels of histones H3 and H4 associated with the mating-type and telomeric silent chromatin as well as with a few sites in rDNA are increased [Braunstein et al. 1993; Suka et al. 2001; Armstrong et al. 2002; Bryk et al. 2002; Hoppe et al. 2002]. We used an antibody that recognizes histone H3 diacetylated at Lys 9 and 14 to examine H3 acetylation levels across the rDNA repeat in Sir2+ versus sir2Δ cells. Figure 2 shows...
Region-specific requirements for rDNA silencing

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the effect of deleting SIR2 on H3 acetylation levels for rDNA regions corresponding to the 25S rRNA, NTS1, and NTS2/18S. We observed an increase in H3 acetylation at each of the above rDNA regions in the range of threefold to fivefold in sir2Δ compared with SIR2+ cells but no effect on H3 acetylation at the control CUP1 locus (Fig. 2A,B, left panels). Furthermore, in four independent experiments, deletion of SIR2 resulted in an increase in H3 acetylation in the range of twofold to sixfold for all rDNA fragments tested (data not shown). As controls, we tested changes in acetylation of H3 at silent chromatin regions at the HMR silent mating-type locus and a telomeric DNA region. As expected (Suka et al. 2001; Bryk et al. 2002; Hoppe et al. 2002), deletion of SIR2 resulted in a large increase in H3 acetylation at both loci but had no effect on H3 acetylation at the control ACT1 locus (Fig. 2A,B, right panels). As an additional control, we performed ChIP with an anti-Sir2 antibody and with the same cross-linked chromatin used above for immunoprecipitation of acetylated H3 to directly compare H3 acetylation with Sir2 occupancy. As expected (Strahl-Bolsinger et al. 1997), the anti-Sir2 antibody efficiently immunoprecipitated HMR and telomeric DNA fragments but not the unsilenced ACT1 locus (Fig. 2A,C, right panels). Consistent with the results presented in Figure 1, we observed the highest levels of Sir2 binding at the NTS1 and NTS2/18S regions and less binding at regions corresponding to the 25S rRNA coding sequences (Fig. 2A,C; data not shown). Therefore, although the highest levels of Sir2 binding are observed at the NTS1 and NTS2/18S regions, deletion of SIR2 causes an increase in H3 acetylation throughout rDNA.

Silencing at NTS1 requires Fob1

The association of silencing proteins with the RFB region of NTS1 was especially intriguing; this section of rDNA is required for FOB1-dependent recombination and contains a replication fork block. Furthermore, the FOB1 and SIR2 genes oppositely affect the formation of extrachromosomal rDNA circles and the rates of cellular senescence (Kaeberlein et al. 1999). These observations...
suggested that Fob1 might block or negatively regulate silencing, but instead we found that silencing at NTS1 was abolished in the absence of FOB1 (Fig. 3A,B). To assess silencing, we deleted FOB1 in strains that contained a Ty1 transposon element bearing a Pol II-transcribed reporter gene [Ty1–mURA3] inserted either outside the rDNA array or within NTS1 [Fig. 1A, R1 reporter, Smith and Boeke 1997]. Cells were 10-fold serially diluted and spotted on complete medium as a plating control and on medium without uracil to monitor silencing of mURA3. Consistent with previous observations, the reporter gene was not silenced outside the array but was efficiently silenced within NTS1, as indicated by poor growth on URA− medium [Fig. 3A, cf. rows 1 and 2; Smith and Boeke 1997]. However, in fob1Δ cells, the reporter gene at NTS1 was completely derepressed [Fig. 3A, cf. rows 2 and 4]. The extent of derepression in fob1Δ cells was comparable to what has been observed in sir2Δ cells [Fig. 3A, row 6]. Western analysis of whole-cell extracts indicated that the levels of Net1–TAP and Sir2–TAP proteins were not altered in fob1Δ cells [Fig. 3D], ruling out the possibility that deletion of FOB1 disrupts silencing at NTS1 by reducing the cellular levels of these silencing proteins. The loss of silencing was specific to FOB1, as addition of a single-copy plasmid containing the FOB1 gene under the control of its own promoter restored silencing to fob1Δ cells [data not shown].

We next tested if FOB1 influences silencing in regions located farther from the NTS1 enhancer. We deleted the FOB1 gene in a strain in which the Ty1–mURA3 reporter gene is inserted in NTS2, adjacent to the 5S gene

Figure 3. Fob1 is required for rDNA silencing at NTS1 but not at NTS2/18S. Silencing was assessed by monitoring the growth of 10-fold serial dilutions of cells on −URA medium. Complete medium was used as a plating control. [A] Both FOB1 and SIR2 are required for Ty1–mURA3 silencing at NTS1. Silencing was assayed using strains containing a Ty1–mURA3 insertion either outside rDNA or at NTS1 [Smith and Boeke 1997]. The approximate location of this reporter (R1) within rDNA is shown in Figure 1A. [B,C] In another reporter gene system, FOB1 and SIR2 are both required for silencing at NTS1 [B, R3 reporter], but only SIR2 is required for silencing at NTS2, near the 35S coding region [C, R4 reporter]. See Figure 1A for the locations of R3 and R4 reporter genes. [D] The levels of Net1–TAP and Sir2–TAP proteins do not change in the absence of Fob1 as shown by Western blotting of whole-cell extracts. Actin is shown as a loading control.
The R2 reporter is typically poorly silenced (Smith and Boeke 1997), and no loss of silencing was detected in *fob1Δ* cells (data not shown). To further test the role of *FOB1* in silencing at the NTS2/18S region, we integrated a plasmid containing the *mURA3* reporter gene without Ty1 elements into three sites: outside the rDNA at the *LEU2* gene and inside rDNA at two locations corresponding to peaks of Net1 and Sir2 association [Fig. 1A, R3 and R4 reporters]. As expected, the *mURA3* gene inserted at either NTS1 or NTS2/18S was strongly silenced [Fig. 3B,C, cf. rows 1 and 2], and this silencing was *SIR2*-dependent [Fig. 3B,C, rows 5 and 6]. The greater concentration of silencing proteins at NTS2/18S compared with NTS1 [Fig. 1B] was consistent with the approximately twofold to fivefold stronger silencing for the R4 reporter in NTS2 compared with the R3 reporter in NTS1 [Fig. 3B,C, cf. row 2]. However, although silencing at NTS1 was abolished in *fob1Δ* cells, deletion of *FOB1* had no effect on silencing of an identical reporter gene at NTS2/18S [Fig. 3B,C, cf. row 4]. Thus, we have identified a novel role for the fork-blocking protein Fob1 and determined that the requirements for silencing of Pol II promoters across an rDNA repeat are not uniform.

**Fob1 colocalizes with Net1 and Sir2 to the RFB region of NTS1**

Fob1–GFP has been shown to localize to the nucleolus (Defossez et al. 1999), but it is not known if Fob1 actually associates with rDNA. To determine if Fob1 is associated with the NTS1 region, we mapped the association of Fob1 with rDNA by ChIP, using a strain in which the endogenous *FOB1* gene was modified to produce a protein with a C-terminal TAP tag [Fob1–TAP; Fig. 1G, lane 4]. Cells expressing Fob1–TAP displayed wild-type levels of silencing, suggesting that the modified protein was functional [Supplementary Fig. 1]. A graph representing the enrichment of rDNA regions in Fob1–TAP immunoprecipitations is shown in Figure 4A. We observed a major peak of Fob1 association, which precisely overlapped the NTS1 RFB region and the peak of Net1–TAP association with this rDNA region [Fig. 4A,C]. Two smaller peaks of binding occurred at the 5S gene and near the NTS2/18S region [Fig. 4B]. In contrast, Fob1 did not associate with the 25S region or the *CUP1* locus [Fig. 4B].

In summary, Fob1, Net1, and Sir2 displayed similar association profiles with respect to the NTS1 region.

**Fob1 is required for localization of Net1 and Sir2 to NTS1**

To further investigate the nature of the silencing defect in cells lacking Fob1, we mapped the associations of Net1–TAP and Sir2–TAP along an rDNA repeat in *fob1Δ* cells by ChIP. The enrichment of NTS1 DNA by Net1–TAP or Sir2–TAP was greatly reduced in *fob1Δ* cells [Fig. 5A,B]. However, similar levels of Net1–TAP and Sir2–TAP were associated with the NTS2/18S region in...
FOB1 and fob1Δ cells [Fig. 5C]. In addition, there was a reproducible increase in relative association toward the interior of 35S [cf. Figs. 5A and 1B]. These findings are consistent with the observed loss of silencing specifically at NTS1 but not NTS2/18S in fob1Δ cells [Fig. 3], and show that Fob1 is required for the proper localization of Net1 and Sir2 to rDNA.

**Fob1 specifically associates with the RENT complex**

The precise colocalization of Fob1 with Net1 and Sir2 at NTS1, as well as its requirement for Net1 and Sir2 to associate with this region, suggested that Fob1 might physically associate with the RENT complex. To test this possibility, we performed coimmunoprecipitation experiments from extracts prepared from yeast strains in which the endogenous copies of Net1 and Fob1 were modified to produce Net1–HA3 and Fob1–Myc13. Cells expressing Fob1–Myc13 displayed wild-type levels of silencing, suggesting that the modified protein was functional [Supplementary Fig. 1]. Immunoprecipitation of Net1–HA3 or Sir2 resulted in coprecipitation of Fob1–Myc13 [Fig. 6A, lane 5, 6B, lane 4], and immunoprecipitation of Fob1–Myc13 resulted in coprecipitation of both Net1–HA3 and Sir2 [Fig. 6A, lane 7]. Deletion of SIR2 had no effect on the amount of Net1–HA3 and Fob1–Myc13 that coprecipitated together, indicating that Net1 and Fob1 can associate independently of Sir2 [Fig. 6C, cf. lanes 3 and 4, 7 and 8]. However, we consistently coprecipitated more Sir2 than Fob1 with Net1 [Fig. 6A–C], suggesting that the Fob1–Net1 interaction was weaker than the Sir2–Net1 interaction. We also tested the interaction of Fob1 with RENT using GST pull-down assays. We purified bacterially expressed GST–Fob1 and a GST–control protein [Fig. 6D, lanes 1, 2], incubated them with whole-cell yeast extracts, and analyzed the bound fractions by Western blotting. We found that GST–Fob1 associated with all three subunits of RENT (Net1–3HA, Sir2, and Cdc14–Myc9), whereas the GST–control protein did not [Fig. 6D, cf. lanes 4 and 5]. Furthermore, GST–Fob1 did not bind Sir3 or Sir4 (silencing proteins not required for rDNA silencing), Tup1 (a general transcriptional corepressor), or Act1 (an abundant cytoskeletal protein; Fig. 6D, cf. lanes 4 and 5), suggesting that the association between Fob1 and the RENT complex was specific.

**RNA polymerase I interacts with Net1 and Sir2**

Our ChIP data indicated that Net1 and Sir2 associated with the NTS2/18S region of rDNA in a Fob1-independent manner. Noticeably, this peak of association with rDNA overlaps the Pol I transcription initiation region. Previous work had demonstrated that the Net1 subunit of RENT is associated with Pol I and is required for optimal Pol I transcription in vitro [Shou et al. 2001]. Our data raise the possibility that this interaction may also...
be important for rDNA silencing and the association of RENT with rDNA. To provide further support for this idea and to specifically address whether other subunits of the RENT complex, in particular Sir2, were also associated with Pol I, we carried out a series of immunoprecipitations. As expected from previous studies (Straight et al. 1999; Shou et al. 2001), immunoprecipitation of Sir2 coprecipitates Net1–HA3 and Fob1–Myc13 (Fig. 7A, lanes 6,8). Furthermore, we found that these interactions were not affected by the presence of ethidium bromide, indicating that they were not DNA-dependent (data not shown). To determine whether Sir2 is also associated with Pol I, the coding regions of RPA135 and RPA190 were modified to express TAP-tagged proteins, and these subunits were immunoprecipitated from whole-cell yeast extracts. Both Pol I subunits associated with Sir2 in the presence of ethidium bromide [Fig. 7B, lanes 5,6]. These results show that both the Net1 and Sir2 subunits of RENT are associated with Pol I and suggest that the localization of Net1 and Sir2 to the NTS2/18S region may result from their physical association with Pol I.

Discussion

We have shown that the Net1 and Sir2 subunits of the rDNA-specific silencing complex RENT are associated primarily with two regions within an rDNA repeat that overlap the NTS1/RFB and transcription initiation regions. The presence of the RENT complex at these functional elements is consistent with a role for silencing in regulation of specific rDNA activities. We show that Fob1 also associates with the NTS1 replication fork block region and is required for silencing of Pol II-transcribed genes at this location. In the absence of Fob1, Net1 and Sir2 are no longer associated with the NTS1 RFB region but are still present at the NTS2/18S region, where silencing occurs independently of Fob1. Both Fob1 and Pol I physically associate with the RENT complex, suggesting two mechanisms for recruitment of silencing proteins to rDNA [Fig. 8] and providing additional evidence that the integrity and function of rDNA is regulated by silencing proteins. Because the overall structural organization of rRNA genes is conserved in organisms ranging from yeast to humans [for review, see Reeder 1999], the intersection of silencing, recombination, and transcriptional pathways described here may also be conserved in other eukaryotes.
Previously identified SIR2 coding region. Notably, these areas overlap with two pre-gion of NTS2 and includes more than 1 kb of the 35S other peak overlaps the PolI transcription initiation sequences that overlap the RFB region, whereas the occupancy (Fig. 1B). The peak within NTS1 is centered out the 35S rRNA region is not uniform. Net1 and Sir2 Binding of the RENT complex to the NTS and through- Distribution of silencing complexes in rDNA

Figure 7. The RENT complex physically associates with RNA polymerase I. [A] Western blots showing that Net1–TAP coimmunoprecipitates with Sir2 [lanes 6, 8] and the largest subunits of RNA polymerase I [Rpa135 and Rpa190, lanes 6, 8] from whole-cell extracts. Act1 serves as a loading control. [−] Un- tagged; [+] tagged; (*) a cross-reacting band. One percent of whole-cell extract [input] and 25% of bound material are shown in A and B. [B] Western blots showing that Rpa135–TAP [lane 5] and Rpa190–TAP [lane 6] coimmunoprecipitate with Sir2.

Binding of the RENT complex to the NTS and throughout the 35S rRNA region is not uniform. Net1 and Sir2 are not exclusively or continuously associated with the nontranscribed region, displaying instead two distinct peaks of association, separated by 1 kb of relatively low occupancy [Fig. 1B]. The peak within NTS1 is centered on sequences that overlap the RFB region, whereas the other peak overlaps the PolI transcription initiation region of NTS2 and includes more than 1 kb of the 35S coding region. Notably, these areas overlap with two previously identified SIR2-responsive regions at NTS1 and 18S, which display altered micrococcal nuclease and dam methyltransferase sensitivities in the absence of SIR2 [Fritze et al. 1997]. In addition, the Net1 and Sir2 association profiles are consistent with qualitative silencing assays, indicating that silencing of Ty1–mURA3 reporters is stronger at a site within NTS1 and near the start of the 18S region as compared with two sites near the 5S gene [Smith and Boeke 1997]. Similarly, the relatively low association of Net1 and Sir2 with certain sections of the 35S RNA coding region is consistent with the relatively weaker silencing of reporter genes inserted at these locations [Smith and Boeke 1997; J. Huang and D. Moazed, unpubl.]. The reduced associations of Net1 and Sir2 with the center of the NTS [Fig. 1B] may be the result of the transcriptional activity of Pol III at the 5S gene or the presence of a barrier element. The left bound- ary of RENT association with the NTS2/18S region coincides with the main binding site of the cohesin complex within rDNA, adjacent to the 5S promoter [Laloraya et al. 2000]. Mutations in some cohesin subunits perturb boundary functions that limit the spreading of silencing at the silent mating-type loci [Donze et al. 1999], suggesting that sites of cohesin association may act as barrier elements. Thus, RENT association may be reduced near the 5S because of the presence of cohesins.

The deacetylase activity of the Sir2 subunit of RENT is required for rDNA silencing [Imai et al. 2000]. In addition, deletion of SIR2 results in an increase in levels of acetylated histones H3 and H4 at the rDNA 5S region [Armstrong et al. 2002], a threefold increase in the levels of diacetylated H3 [Lys 9 and 14] at one site in the NTS [Bryk et al. 2002], a 1.6- to 2.4-fold increase in acetylated H4 at two sites in NTS2 near the PolI promoter region [Hoppe et al. 2002], and increases in both H3 [K9/K14] and H4 acetylation at a site within the NTS1 and transcriptional initiation regions [Buck et al. 2002]. Our ChIP analysis indicates that although Net1 and Sir2 are localized primarily to two rDNA regions, Sir2 is required for global hypoacetylation of histone H3 associated with rDNA chromatin [Fig. 2; J. Huang, unpubl.].

Recruitment of RENT to the replication fork block region of NTS1 by Fob1

Our data show that Fob1 is required for the association of the RENT complex with the rDNA NTS1 region and for silencing at this location. Our finding that Fob1 is present at the replication fork block region of NTS1 is consistent with Fob1’s requirement for the replication fork block and recombination activities of cis-elements within this region. Additionally, we find that Fob1 is physically associated with RENT in vivo, suggesting that Fob1 is a recruitment factor for silencing complexes. Interestingly, Fob1 possesses homology to two highly conserved domains of retroviral integrases, which are known to mediate DNA cleavage and strand-transfer reactions [Dlakic 2002], raising the possibility that Fob1 binds to DNA at or near the RFB directly.

Our finding that Fob1 is required for rDNA silencing was surprising, given its role in promoting recombina- tion. One possible explanation for our results is that si- lencing complexes are recruited to the RFB region by Fob1 to counteract the recombination potential of rep- lication fork barriers that are also generated by Fob1. In prokaryotic systems, blocked replication forks can lead to double-strand breaks that are substrates for homolo- gous recombination (for review, see Rothstein et al. 2000), and similar processes may occur at the rDNA of yeast. Alternatively, association of RENT with the Fob1 protein may inhibit the replication fork-blocking activity of Fob1 and thereby Fob1-stimulated recombination. Our data are also consistent with a model in which the RENT complex acts directly on Fob1 to inhibit its fork- blocking activity, for example, by deacetylating it. Fur- thermore, our findings are consistent with the opposing roles of Fob1 and Sir2 in regulation of rDNA recombin-
tion and yeast lifespan (Gottlieb and Esposito 1989; Kobayashi and Horiuchi 1996; Defossez et al. 1999; Kaebelerlein et al. 1999; Johzuka and Horiuchi 2002). Fob1, a positive regulator of rDNA recombination, recruits its own inhibitor, Sir2, to rDNA. Therefore, in the absence of the inhibitory function of Sir2, the enhancement of recombination by Fob1 is unopposed, and increased recombination results in increased accumulation of ERCs and reduced life span in $sir2/H9004$ cells. Because rDNA recombination requires Fob1, in $fob1/H9004$ cells recombination levels are dramatically reduced, and because Sir2 is no longer recruited to the NTS1 region, silencing at this region is also abolished.

It is notable that a positive regulator of recombination is located in a region of the genome that, because of its repetitive organization, is inherently recombinogenic. In wild-type cells, the levels of recombination at rDNA are significantly lower than predicted for a large and repetitive locus, despite the stimulatory presence of Fob1. Moreover, in the absence of Fob1 or both Fob1 and Sir2, rDNA recombination levels remain low (Defossez et al. 1999, Kaebelerlein et al. 1999), suggesting that an additional level of regulation is present that suppresses recombination. Some level of recombination is still necessary, which may explain the positive role of Fob1. For example, unequal sister-chromatid exchange during recombination results in either the gain or loss of repeats, allowing the maintenance of a favorable number of repeats as growth conditions dictate. Contraction of repeats also facilitates the removal of dominant-negative mutations, maintaining the integrity of the array. The ability of Fob1 to promote recombination and to recruit the RENT complex may provide a mechanism for fine-tuning rDNA recombination levels. In addition, our findings are consistent with a model that in wild-type yeast cells, rDNA recombination and replicative life span are regulated by the dual silencing and recombination activities of Fob1.

Association of RENT with the NTS2/35S region

Net1 and Sir2 are also associated with an rDNA region of ~1.5–2 kb spread over part of NTS2 and the 35S rRNA coding region [Fig. 1B]. Because association with this region is not Fob1-dependent [Fig. 5A], an as-yet-unidentified protein recruits silencing complexes specifically to this region. We suggest that the most likely candidate is RNA polymerase I and/or its associated transcription factors. Net1 binds purified Pol I complexes and is required for optimal Pol I-dependent transcription in vitro (Shou et al. 2001), and rDNA silencing is impaired in cells that lack a functional Pol I (Buck et al. 2002). Our findings that both the Net1 and Sir2 subunits of RENT associate with Pol I (Fig. 7A,B) support the hypothesis that the polymerase itself recruits the RENT complex to the NTS2/18S region of rDNA (Fig. 8).

It remains unclear why silencing complexes are recruited to an area of heavy transcription initiation. This region, like the NTS1 replication fork block region, may be involved in stimulating recombination. For example, optimal $HOT1$ activity requires a cis-element located within the Pol I transcription initiation region in addition to the sequences located at NTS1 [Keil and Roeder 1984; Voelkel-Meiman et al. 1987]. Thus, the $HOT1$ element of NTS2 may contribute to hotspot activity within rDNA. Furthermore, there is evidence supporting a link between recombination and Pol I activity, because it has been shown that $HOT1$ sequences fail to stimulate recombination at an ectopic location in the absence of Pol I [Huang and Keil 1995]. This observation suggests that the presence of Pol I or its transcriptional activity may be required for rDNA recombination. Analogous to the situation with Fob1 at NTS1, the RENT complex might associate with Pol I to suppress Pol I-stimulated recombination. Alternatively, Pol I-dependent silencing in the NTS2 region may influence recombination rates by regulating the activity of DNA replication origins within rDNA. Recently, it has been shown that in rDNA, functional replication origins are clustered and separated by large regions where initiation firing is suppressed in a $SIR2$-dependent manner (Pasero et al. 2002). In principle, clustering of origin-firing would reduce the number of active replication fork blocks within NTS1 regions, thereby minimizing the likelihood of recombination (Pasero et al. 2002).

The association of Net1 and Sir2 with the Pol I transcription initiation region is strikingly polar, with the highest area of occupancy located toward the 35S coding region and decreasing association toward the middle of the 35S gene [Figs. 1B, 5A]. This association pattern is consistent with the role of Net1 as a Pol I transcription factor (Shou et al. 2001) and with the spreading of silenc-
ing to sequences flanking the rDNA array in the direction of Pol I transcription [Buck et al. 2002]. However, because Net1 remains localized to the nucleolus throughout the cell cycle, and Sir2 is required for hypo-acetylation of H3 throughout the rDNA, it is unlikely that low levels of RENT association with rDNA regions corresponding to the middle and 3′-end of 35 rRNAs reflect dissociation from Pol I during transcription elongation [Fig. 2, Shou et al. 1999, Straight et al. 1999; Visintin et al. 1999]. We favor the possibility that this polar pattern of localization mirrors the mode of association of Pol I or one of its associated factors with rDNA [Figs. 1, 8]. Finally, we note that the recruitment of RENT to rDNA by either Pol I or Fob1 may provide redundant and/or novel opportunities for regulation of the cell cycle functions of this complex.

Materials and methods

Yeast strains and plasmids

A strain table is included in the Supplemental Material. NET1, SIR2, FOBI, RPA135, and RPA190 genes were modified with the C-terminal TAP tag as described [Rigaut et al. 1999], and the NET1 gene was modified with a C-terminal HA3 tag by integrating the plasmid pDM239 as described [Straight et al. 1999]. The Myc9-tagged Cdc14 strain was a kind gift from R. Deshaies ([California Institute of Technology, Pasadena, CA). The RPA190 and RPA135 genes were modified with C-terminal HA3 tags, the FOBI gene was modified with the C-terminal Myc13 tag, and the FOBI and SIR2 deletion strains were generated by replacing their open reading frames with the promoter (California at Berkeley, Berkeley, CA).

The NET1 gene was modified with a C-terminal HA3 tag by integrating the plasmid pJR531 (the kind gift of J. Rine, University of California at Berkeley, Berkeley, CA). The RDN1–NTS1 mURA3–LEU2 gene from plasmid pJR531 was created similarly to [Smith and Boeke 1997]. The RDN1–NTS2 mURA3–LEU2 gene was modified by replacing their open reading frames with the TRP1 promoter (California Institute of Technology, Pasadena, CA). The mURA3 silencing regions were generated by transformation with pDM316 cut with SmaI PCR product of NTS2 and pDM704 cut with EagI PCR product of NTS1. All transformations were performed with the lithium acetate method [Guthrie and Fink 1991], and proper integration was confirmed by PCR.

pDM316 (LEU2::mURA3) was constructed by ligation of a 1.6-kb BamHI–EagI PCR product containing the mURA3 gene into pRS305. pDM312 (RDN1–NTS2::mURA3–LEU2) was generated by ligation of a 1-kb Xhol–EcoRI PCR promoter of NTS2 and 1.6-kb EcoRI–EagI PCR product containing the mURA3 gene into pRS305 digested with Xhol and EagI. pDM704 (RDN1–NTS1::mURA3–LEU2) was created similarly to pDM312 except using the 0.5-kb Xhol–EcoRI PCR product of NTS1. The mURA3 gene was amplified from pJJ560-2 [Smith and Boeke 1997], and NTS regions were amplified from genomic DNA.

ChIP assays

ChIP assays were carried out as described [Suia et al. 2001], with modifications. Yeast cultures (50 mL) were grown to an OD600 of 1.5–1.8 and cross-linked with 1% formaldehyde at room temperature for 15 min. The reaction was quenched with glycerol at a final concentration of 125 mM for 5 min. Cells were washed twice with cold TBS [20 mM Tris-HCl at pH 7.6 and 150 mM NaCl] and frozen at −80°C. Cell pellets were resuspended in 400 µL of lysis buffer [50 mM HEPES–KOH at pH 7.5, 500 mM NaCl, 1 mM EDTA at pH 8.0, 1% Triton X-100, 0.1% sodium deoxycholate, 0.1% SDS, and protease inhibitors] and bead-beat with glass beads (beads and Mini Beadbeater, Biospec Products) twice for 30 sec. Lysates were sonicated three times for 20 sec at 40% amplitude [Branson Digital Sonifer], and centrifuged at 13,000 rpm for 5 and 15 min. To obtain input DNA, 50 µL of clarified lysate was used, and for each immunoprecipitation reaction.

Thirty microliters of a 50% slurry of prewashed IgG1–agarose beads [Sigma] was incubated with each lysate at 4°C for 2 h. For Sir2 or diAcH3 ChIP experiments, 150 µL of lysate was incubated at 4°C overnight with 1.5 or 2.0 µg of antibody [polyclonal anti-Sir2, Hoppe et al. 2002; anti-Ack9/Ack14 H3, Upstate Biotechnology], and further incubated with Protein A Sepharose beads at 4°C for 2 h. Beads were washed three times in lysis buffer, once with 10 mM Tris–HCl [pH 8.0], 0.25 M LiCl, 0.5% NP-40, 0.5% sodium deoxycholate, and 1 mM EDTA, and once with TE [pH 8.0] at room temperature. Beads were eluted by incubating with 100 µL of 50 mM Tris–HCl [pH 8.0], 1 mM EDTA [pH 8.0], and 1% SDS at 65°C for 15 min. Eluate was transferred to a fresh tube and pooled with a final bead wash of 150 µL of TE with 0.67% SDS. For input DNA, 200 µL of TE with 1% SDS was added to 50 µL of lysate. All samples were incubated at 65°C overnight, combined with 250 µL of TE, 15 µg of glycerol, and 100 µg of Proteinase K, and incubated at 37°C for 2 h. After addition of 55 µL of 4 M LiCl, samples were extracted once with phenol:chloroform:::soyamyl alcohol and once with chloroform. Precipitated and washed DNA was resuspended in 50 µL of TE with 10 µg of RNase A and incubated at 37°C for 1 h. PCR reactions [12.5 µL contained 2.0 µL of template DNA (1:8 dilution of IP and 1:2,000 dilution of WCE for TAP-tag strains; 1:8 dilution of Sir2 and 1:16 dilution of di-AcH3 IPs, and 1:25,000 dilution for WCE), 1.25 µCi of [α-32P]dCTP. PCR parameters were 1 cycle of 95°C for 2 min, 55°C for 30 sec, and 72°C for 1 min, followed by 21 [multicopy genes] or 28 [single-copy genes] cycles of 95°C for 30 sec, 55°C for 30 sec, and 72°C for 1 min, and a final step of 72°C for 4 min. Primers of 20-nt oligonucleotides were designed with Primer3 software [http://www.basic.nwu.edu/biotools/Primer3.html] to amplify products −250 bp for rDNA or −150 bp for ACT1 or CUP1. Sequences are available upon request. Primers for HMR-E and telomeric sequences (0.6 kb from the end of Chromosome VI-R) have been previously described [Hoppe et al. 2002]. Samples were run on 6% polyacrylamide gels at 100 V for 45 min. PCR products were quanititated using QuantityOne software. Relative fold enrichment was determined by calculating the ratio of rDNA[IP] to rDNA[WCE] and normalizing the data such that the untagged background ratio in Figures 1B, 4A, and 5A is −1. Each set of experiments was performed at least three times and produced similar binding profiles. For multiplex PCR, fold enrichment values for each strain were calculated as follows: [rDNA[IP]/CUP1[IP]]/[rDNA[WCE]/CUP1[WCE]]. The untagged strain value was normalized to 1, yielding the rDNA fold enrichment for the tagged strains. In Figure 2B, the relative fold enrichment is defined as the ratio of SIR2 to sir2Δ for values from the following calculation: [rDNA[IP]/CUP1[IP]]/[rDNA[WCE]/CUP1[WCE]].
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Silencing assays

Cells were grown in YEPD to an OD_{600} of 1.6–1.8 and concentrated to two-thirds the original volume. Three microliters each of 10-fold serial dilutions were spotted on appropriate media. Plates were incubated at 30°C for 2–3 d.

Immunoprecipitation reactions

Reactions were performed essentially as described previously [Straight et al. 1999]. One percent of input whole-cell extract and 25% of bound fractions were run on SDS-PAGE gels, blotted, and probed with the indicated antibodies.

Purification of GST fusion proteins

GST–Fob1 protein was expressed from pDM708, which was constructed by ligation of a 1.7-kb Bcl–XhoI PCR product containing the FOB1 gene [amplified from genomic DNA] with pGEX-6P-1 (Amersham Pharmacia). The GST–control protein was constructed by ligation of a 1.7-kb BclI–Purification of GST fusion proteins

GST pull-downs

Purified GST fusion proteins (5 μg) were bound to glutathione agarose [50 μL of packed resin] at 4°C for 1 h in 200 μL of yeast lysate buffer [50 mM HEPES-NaOH at pH 7.5, 100 mM NaCl, 10% glycerol, 1 mM EDTA, 2 mM DTT, 1 mM PMSF, 0.1% NP-40, and protease inhibitors]. Beads were washed twice with a column and washed with wash buffer [PBS with 50 mM NaCl, 0.5 mM DTT, 0.1% NP-40] and with wash buffer without NP-40. Column was eluted with 50 mM Tris-HCl [pH 8.0], 250 mM NaCl, and 10 mM glutathione. Peak fractions were pooled and dialyzed against 50 mM HEPES-NaOH [pH 7.6], 300 mM NaCl, and 30% glycerol and stored at −80°C.

GST pull-downs

GST pull-downs

Purified GST fusion proteins (5 μg) were bound to glutathione agarose [50 μL of packed resin] at 4°C for 1 h in 200 μL of yeast lysate buffer [50 mM HEPES-NaOH at pH 7.5, 100 mM NaCl, 10% glycerol, 1 mM EDTA, 2 mM DTT, 1 mM PMSF, 0.1% NP-40, and protease inhibitors]. Beads were washed three times with a column and washed with wash buffer [PBS with 50 mM NaCl, 0.5 mM DTT, and 0.1% NP-40] and with wash buffer without NP-40. Column was eluted with 50 mM Tris-HCl [pH 8.0], 250 mM NaCl, and 10 mM glutathione. Peak fractions were pooled and dialyzed against 50 mM HEPES-NaOH [pH 7.6], 300 mM NaCl, and 30% glycerol and stored at −80°C.

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Association of the RENT complex with nontranscribed and coding regions of rDNA and a regional requirement for the replication fork block protein Fob1 in rDNA silencing

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