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The *RPC31* Gene of *Saccharomyces cerevisiae* Encodes a Subunit of RNA Polymerase C (III) with an Acidic Tail

CHRISTINE MOSRIN,¹ MICHEL RIVA,^{1,2} MONICA BELTRAME,³ EVELYNE CASSAR,¹ ANDRÉ SENTENAC,^{1,2} and PIERRE THURIAUX^{1,2*}

Service de Biochimie^{1*} and Laboratoire d'Ingénièrie des Protéines,² Commissariat à l'Energie Atomique, Centre d'Etudes Nucléaires de Saclay, F91191 Gif sur Yvette, France, and European Molecular Biology Laboratory, D-6900 Heidelberg, Federal Republic of Germany³

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The *RPC31* gene encoding the C31 subunit of *Saccharomyces cerevisiae* RNA polymerase C (III) has been isolated, starting from a C-terminal fragment cloned on a λ gt11 library. It is unique on the yeast genome and lies on the left arm of chromosome XIV, very close to a *Not*I site. Its coding sequence perfectly matches the amino acid sequence of two oligopeptides prepared from purified C31. It is also identical to the *ACP2* gene previously described as encoding an HMG1-like protein (W. Haggren and D. Kolodrubetz, Mol. Cell. Biol. 8:1282–1289, 1988). Thus, *ACP2* and *RPC31* are allelic and encode a subunit of RNA polymerase C. The c31 protein has a highly acidic C-terminal tail also found in several other chromatin-interacting proteins, including animal HMG1. Outside this domain, however, there is no appreciable homology to any known protein. The growth phenotypes of a gene deletion, of insertions, and of nonsense mutations indicate that the C31 protein is strictly required for cell growth and that most of the acidic domain is essential for its function. Random mutagenesis failed to yield temperature-sensitive mutants, but a slowly growing mutant was constructed by partial suppression of a UAA nonsense allele of *RPC31*. Its reduced rate of tRNA synthesis in vivo relative to 5.8S rRNA supports the hypothesis that the C31 protein is a functional subunit of RNA polymerase C.

The transcription of the eucaryotic genome is catalyzed by three distinct RNA polymerases, A, B, and C, acting in conjunction with their transcription factors. Enzyme A (or I) synthesizes the large nucleolar precursor of rRNA, enzyme B (or II) synthesizes the mRNAs, and enzyme C (or III) synthesizes 5S rRNA, tRNAs, and a few other small, untranslated RNA species. Unlike the bacterial enzyme, eucaryotic RNA polymerases have no template specificity but select the appropriate class of promoters by recognizing specific preinitiation complexes formed between the promoter and cognate transcription factors (TFIIIA and TFIIIC in the case of enzyme C). They have a much more complex subunit structure than does the bacterial RNA polymerase, since the yeast enzymes dissociate into 12 distinct polypeptidic components under denaturing conditions. This astonishing structural complexity is still poorly understood in functional terms (11, 43, 44).

Biochemical and genetic studies of yeast polymerases have suggested a minimal subunit structure (38) formed of the two large subunits strongly related to the bacterial β' and β subunits (1, 28, 46), three small subunits (ABC27, ABC23, and ABC14) common to all three enzymes (18), and at least one subunit (AC40) (25) identical in enzymes A and C and structurally equivalent to the B44 subunit of enzyme B (23). Another subunit shared by enzymes A and C (AC19) probably has no counterpart in enzyme B (18). In addition, each RNA polymerase contains several specific subunits. There are at least four of them (C80, C53, C34, and C31) in enzyme C (18) and probably also four in RNA polymerase A (38). In the latter case, the specific subunits are polymorphic even between closely related yeast species such as Saccharomyces cerevisiae and S. douglasii (38). The functional role of these small and specific subunits in the cognate RNA polymerase remains to be understood and was genetically approached in the present work by identifying and characterizing the gene encoding the C31 subunit of yeast RNA polymerase C.

MATERIALS AND METHODS

Strains and media. Strains are listed in Table 1. Yeast genetic techniques and media were described by Sherman et al. (45). Except for tryptophan, the amino acid requirements were also met by adding 0.1% Casamino Acids to the standard minimal medium. Reduction in the wild-type growth rate was obtained by adding 2% sterile filtered glucosamine to the Casamino Acids medium containing 0.5% glucose as the carbon source (27). The 5-fluoro-orotic acid medium was described by Boeke et al. (4). For experiments with *Escherichia coli*, strain DH5 α F'I⁹, obtained from Bethesda Research Laboratories, Inc., was used. Bacterial media were as described by Davis et al. (8).

Plasmid constructions. The RPC31 plasmids pJC31, pC326, pC329, and pSE359-RPC31 were constructed by inserting the 2.5-kilobase (kb) EcoRI fragment spanning *RPC31* (isolated from plasmid YCp50-C31; see Fig. 1) into the pJDB110 (33), pEMBLYCp32, pFL39, and pSE359 vectors. Plasmid pEMBLYCp32 (constructed by C. Mann) is pEMBLYi32 (2, 14) with a 1.4-kb ARS1-CENIV cassette inserted at the AatII site. Plasmid pFL39 is a centromeric vector obtained from F. Lacroute. It harbors a 0.83-kb EcoRI-PstI TRP1 fragment (without ARS1) and a 0.8-kb TagI fragment containing CENVI and an uncharacterized ARS fragment. Plasmid pSE359 (obtained from C. Mann) is a 5.3-kb YRp vector derived from TRP1 YRp7 (33). It has an EcoRI, SacI, BamHI, SalI, and SphI multiple cloning site and bears the SUPM11-o ochre (UAA) supressor allele corresponding to a tRNA^{Tyr} mutated at the anticodon. Plasmid pC50 was constructed by inserting the 2.1-kb Spel-EcoRI RPC31 fragment into the centromeric vector pUN50

^{*} Corresponding author.

TABLE 1. Yeast strai

Strains	Genotype"	Source or reference ^b
YNN281	MATa ade2-101 ura3-52 lys2-801 his3 $\Delta 200$ trp1- Δ	YGSC
YNN282	MATα ade2-101 ura3-52 lys2-801 his3Δ200 trp1-Δ	YGSC
D27-7c	MATa ade2-101 ura3-52 lys2-801 his3 Δ 200 trp1-901 leu2-3,112 rpc160-41 pep4::HIS3	14
D34-3a	MATα ade2-101 ura3-52 lys2-801 his3* trp1-Δ800 leu2-3,112	This work
D34-3b	MATa ade2-101 ura3-52 lys2-801 his3* trp1∆800 leu2-3,112	This work
CDM62	MATα ade2-101 ura3-52 lys2-801 his3* trp1-Δ800 leu2-3,112 rpc31Δ::LEU2 pep4::HIS3 (pC326)	This work
DM824	MATα ade2-101 ura3-52 lys2-801 his3* trp1Δ800 leu2-3,112 rpc31Δ::LEU2 pep4::HIS3 (pSE359-Tyr4)	This work
Le788	MATa ura3 met2 met4 lys10 arg8 ade1 his6 pha2	F. Hilger
KJ382-23A	MATa spoll ura3 cyh2 ade2 his7 home3 tyrl canl RPC3::pJC31	This work
DM32-1	MATa ura3-52 trp1 RPC31::Y1pC327	This work
DM33-1a	MATa ura3* met2 pha2 his6 ade1	This work
DM34-7d	MATa trp1 RPC31::Y1pC327 ura3*	This work
D55-7b	MATa ura3* met4 petx ade2	This work

^a his3* is either his3Δ200 or his3-11,15; ura3* is an unidentified ura3 allele (possibly ura3-52).

^b YGSC, Yeast Genetic Stock Center, Berkeley, Calif.

(9). To construct pBSC31-5, we digested plasmid pC329 with *Not*I, subjected it to a brief digestion with exonuclease III, and treated it with mung bean nuclease. The plasmid was further cut with *Eco*RI, and the small 1.2-kb fragment was ligated to the pBluescript KS (Stratagene) plasmid. The resulting plasmid (pBSC31-5) bearing no ATG before the first ATG of the C31 coding region was used for in vivo transcription and translation experiments as recommended by Stratagene (transcription) and by Promega Biotech (translation).

Disruption of the RPC31 gene. The rpc31::HIS3a and rpc31::HIS3b alleles were produced by inserting the HIS3 gene (a 1.7-kb BamHI fragment of plasmid pSZ63 [33]) in both orientations at the BamHI site of RPC31. The last 6 amino acids of the RPC31 coding sequence were replaced by 30 others in rpc31::HIS3a and by 6 unrelated amino acids in rpc31::HIS3b (see Fig. 3). rpc31\Delta::LEU2 was constructed by replacing the 810-base-pair (bp) SalI fragment of RPC31 with the 2.2-kb XhoI-SalI LEU2 marker from YEp13 (33). Heterozygous RPC31/rpc31::ΔLEU2 or RPC31/rpc31::HIS3 diploids were constructed in strain YNN281 \times YNN282 by the allele replacement technique (42). Their genetic structures were confirmed by Southern analysis with the 2.1-kb XbaI-SalI RPC31 fragment (see Fig. 1) as a probe. The rpc31::Tn10-13, -33, and -38 alleles were obtained by transposon mutagenesis (19). Their approximate insertion sites were determined by restriction mapping. Tn10-33 appeared to be inserted about 100 bp upstream from the RPC31 coding sequence itself. Tn10-13 generated an N-terminal fusion of lacZ to RPC31, as shown by the blue color of the corresponding mutant on YPD supplemented with 5-bromo-4chloro-3-indolyl-B-D-galactopyranoside (X-Gal) at 40 mg/ liter.

In vitro mutagenesis. Plasmid pC329 (*TRP1 RPC31*) or pC50 (*TRP1 RPC31*) was mutagenized with hydroxylamine (40) and directly introduced into the $rpc31\Delta$::*LEU2* strain CDM62(pC326) by transformation. Each transformant may receive several distinct plasmid molecules (which would prevent the detection of recessive phenotypes), but the cellular plasmid content of individual cells should become homogeneous after a few rounds of mitotic segregation. Accordingly, the initial transformants were pooled and respread on selective medium at appropriate dilutions to form individual colonies, which were then replica plated on 5-fluoro-orotic acid medium. Defective mutants are unable to grow on 5-FOA medium, which selects for colonies that have lost pC326. This selection was done at different tem-

peratures (16, 25, 30, and 37°C) to identify conditional mutants in addition to fully defective ones. No conditional RPC31 mutant was obtained, but 11 temperature-sensitive TRP1 mutants were isolated, indicating that plasmid-borne conditional mutants were generated and detected under our experimental conditions. A total of 225 fully defective rpc31 mutants were obtained (corresponding to 1.5% of the colonies). The mutagenesis was probably close to saturation because 166 GC \rightarrow AT transitions are possible on the *RPC31* coding sequence, of which 45 are likely to be silent mutations, as they would correspond to substitutions in the acidic C-terminal domain (see Results). The six nonsense alleles were constructed by oligonucleotide-directed mutagenesis of the pC50 plasmid with a kit from Amersham Corp. Sequencing of the entire *RPC31* coding sequence showed the absence of spurious additional mutations in the corresponding clones.

In vivo labeling of RNAs. Exponential cultures (10 ml) on Casamino Acids medium supplemented with 0.02% uracil and tryptophan were transferred by filtration to a medium containing 5 \times 10⁵ Bq of tritiated uracil at a final concentration of 0.5 mg/liter. After 15 min of growth, the culture was chilled and harvested by centrifugation at 4°C. The pellet was suspended in 2.5 ml of cold TE buffer (10 mM Tris hydrochloride [pH 7.4], 10 mM EDTA) with 0.5% sodium dodecyl sulfate, and the nucleic acids were extracted in the same volume of phenol (saturated in TE buffer) under vigorous agitation at 65°C for 1 h. The aqueous phase was precipitated with sodium acetate (final concentration, 0.3 M) plus 2.5 volumes of ethanol. The pellet was ethanol washed, dried, and suspended in 200 µl of water. The yield was about 200 µg of RNA, as measured by UV A_{260} and A_{280} . RNAs were separated by electrophoresis on a 6% polyacrylamide gel as previously described (14).

DNA sequences. Overlapping restriction subfragments of the 2.5-kb EcoRI fragment (see Fig. 1) were inserted in the pBS⁺ or pBS⁻ plasmid (Stratagene). The sequence was determined on both strands by the dideoxy method with modified T7 DNA polymerase (Sequenase, United States Biochemical Corp.). Open reading frame and acid-base-amino acid mapping was done with the DNA Strider program (26). Searches of data bases, secondary structure predictions, and sequence alignments were performed at computer facilities at CITI2, Paris, France, on a VAX8530 computer (Ministère de la Recherche et de la Technologie) with the FASTA package program of Pearson and Lipman (34).

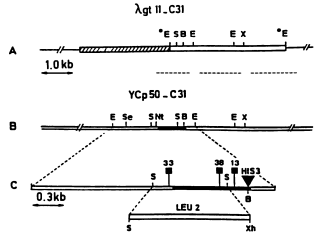


FIG. 1. Restriction map of the RPC31 region. (A) \gt11-C31 insert (3.6 kb), with approximate locations of three transcripts, as deduced from Fig. 2. (B) YCp50-C31 insert (13.5 kb). (C) Expanded view of the 2.5-kb EcoRI fragment spanning RPC31, with approximate locations of deletion and insertion mutations. 33, 38, and 13 indicate three insertions of the Tn10-LUK (URA3⁺) minitransposon. HIS3 indicates the 1.7-kb yeast BamHI HIS3 cassette inserted in either orientation and leading to a recessive-lethal (rpc31::HIS3a) or viable (rpc31::HIS3b) phenotype (see also Fig. 3). LEU2 indicates a Sall-XhoI (2.2-kb) cassette. Solid bars indicate the RPC31 coding sequence. Open boxes indicate yeast genomic DNA outside the RPC31 gene. The hatched box indicates the lacZ DNA on λ gt11. Thin lines indicate vector DNA. Broken lines indicate approximate locations of the three transcripts hybridized by the λ gt11-C31 DNA (see Fig. 2). Restriction sites: B, BamHI; E, EcoRI; Nt, NotI; S, SalI; Se, SpeI; X, XbaI; Xh, XhoI. *E denotes artificial EcoRI sites created by the construction of the λ gt11-C31 clone.

Microsequencing of oligopeptides. RNA polymerase C prepared as described by Huet et al. (18) showed the usual polypeptide pattern except that the 37-kilodalton polypeptide was absent from the preparation. The enzyme (2 mg) was loaded on a 12% polyacrylamide gel, and the subunits were separated by electrophoresis under denaturing conditions. The band containing the C31 subunit was cut from the gel and incubated with tolylsulfonyl phenylalanyl chloromethyl ketone-treated trypsin for 18 h at 37°C in 0.1 M ammonium acetate (pH 8.3) at a protein/enzyme ratio of 1:50 (wt/wt). The peptides were separated by high-pressure liquid chromatography on an Aquapore RP300 reversed-phase column (10 cm by 2.1 mm; Brownlee Laboratories). The column was developed at 40°C with a linear gradient of 25 mM ammonium acetate (pH 6.2) to 60% acetonitrile-50 mM ammonium acetate (pH 6.2) at 200 μ l/min for 45 min. A₂₁₄ peaks were collected and chromatographed in a second system (0.115% trifluoroacetic acid to 60% CH₂CN-0.1% trifluoroacetic acid). Fractions corresponding to A_{214} peaks were spotted on a precycled, Polybrene-coated, trifluoroacetic acid-activated glass fiber filter for sequencing in a 477 Sequenator (Applied Biosystems).

RESULTS

Cloning of gene *RPC31*. Starting from a previously isolated λ gt11-C31 clone (39), the yeast insert (a 3.6-kb fragment delimited by two *Eco*RI sites generated by the cloning procedure) was subcloned, and an internal 2.1-kb *SalI-XbaI* fragment (Fig. 1A) was used as a probe to screen a yeast genomic library harbored by the centromeric plasmid YCp50

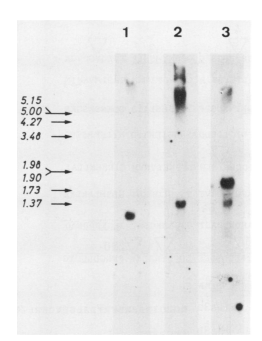


FIG. 2. Northern blot analysis. Yeast RNA (15 µg) from strain YNN281 was electrophoresed in a 1% agarose gel after denaturation with glyoxal, transferred to a nitrocellulose membrane, and hybridized with the three consecutive 0.6-, 1.2-, and 1.8-kb *Eco*RI fragments (lanes 1, 2, and 3, respectively) of the λ gt11-C31 clone (Fig. 1A) under high-stringency conditions. Denaturated λ DNA restriction fragments were used as molecular weight markers. Numbers at left are in kilobases.

(41). One of the plasmids obtained (YCp50-C31) has a 13.5-kb Sau3A insert which contains a NotI site (8 bp) and includes the whole RPC31 gene (Fig. 1B). Northern (RNA) hybridization showed that the initial 3.6-kb insert hybridized to three transcripts of about 1.2, 1.5, and 1.7 kb (Fig. 2) which could be approximately located as depicted in Fig. 1A. The 1.2-kb transcript corresponded to the RPC31 fragment of λ gt11-C31. Its size was consistent with a gene product of 31 kilodaltons.

We determined the 1.279-bp sequence of the Sall-EcoRI fragment spanning RPC31 (Fig. 1). The coding sequence (Fig. 3) is identical to that of the ACP2 gene independently isolated by Haggren and Kolodrubetz (15) and interpreted by them as encoding an HMG1-like protein (our arguments against this interpretation are given below, in the amino acid sequence analysis section). The corresponding region is present in a single copy on the yeast genome (Fig. 4), showing that RPC31 and ACP2 are the same gene. To firmly establish that this gene encodes the C31 protein, we digested a purified preparation of C31 with trypsin and microsequenced two of the resulting oligopeptides. In both cases, there was a perfect match between the observed and predicted amino acid sequences (Fig. 3). One oligopeptide (GGSNNYMSNDP) included the Met at position 14, allowing us to define the initiator codon as indicated in Fig. 3. The *RPC31* reading frame indicates a theoretical M_r of 27,707, somewhat lower than the M_r of 31,000 indicated by electrophoretic migration in polyacrylamide gels (18). This discrepancy is unlikely to reflect a postranscriptional modification of the gene product, since an apparent M_r of 31,000 was also observed when the RPC31 gene was expressed in vitro from plasmid pBSC32-5 (data not shown). Similar minor discrepancies have been observed for other subunits of RNA polymerase C (1, 25).

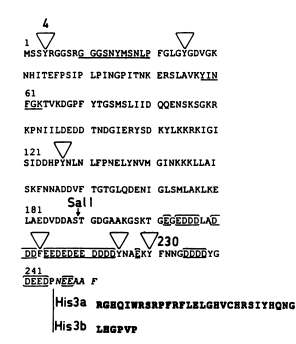


FIG. 3. Amino acid sequence of the *RPC31* gene. The two oligopeptides determined by microsequencing are underlined. The Asp-Glu acidic tail is bracketed. The *Sall* site defining the border of the two DNA probes used in Fig. 4 is indicated by an arrow. The predicted amino acid substitutions generated by the *rpc31::HIS3a* (lethal) and *rpc31::HIS3b* (viable) insertions are shown in boldface type (see the text). Open triangles represent substitutions of UAU (Tyr) codons by a nonsense UAA (*rpc31-Tyr4*) or UAG (*rpc31-Tyr230*) triplet. EMBL/GenBank accession number X51498.

Chromosome mapping. Since our data leave no doubt that RPC31 encodes the C31 subunit of RNA polymerase C, we conserved the gene symbol RPC31 initially proposed (39). This gene was assigned to chromosome XIV by the method of Falco and Botstein (10). Plasmid pJC31 (URA3 RPC31) was integrated into RPC31 in the MATa strain K382-23A by targeted integrative crossover (31). The resulting RPC31:: pJC31 (ura3 RPC31) transformant (strain KJ382-23A; Table 1) was crossed to three MATa ura3⁻ strains, K396-22B, K381-9D, and K393-35C, which together harbor at least one auxotrophy marker on each chromosome (21). The spontaneous ura3 segregants (formed at a frequency of about 2%) were prototrophic for all but one (met4) of the heterozygous markers, thereby indicating that RPC31, like MET4, maps on chromosome XIV. A more precise mapping was obtained by crossing the MATa RPC31::Y1pC327 (ura3 RPC31) strain DM32-1 or DM34-7d to the ura3 mutant strains DM33-1a (MAT_{α}), D55-7b (MAT_a), and Le788 (MAT_{α}), which bear various genetic markers (pha2, petx, met2, and met4) on chromosome XIV (Table 1) (29). Tetrad analysis allowed us to localize RPC31 on the left arm of the chromosome, between MET4 (23 parental ditypes, 22 tetratypes, and 3 recombinant ditypes) and PETX (8 parental ditypes and 5 tetratypes). The MAK26 gene also maps in that region (49), but its precise position relative to that of RPC31 has not been determined.

Amino acid sequence analysis. The calculated isoelectric point of the C31 subunit (assuming no polar modification of amino acids) is 4.5. This is largely due to a highly acidic C-terminal domain, which mainly consists of Asp and Glu downstream from position 202 (Fig. 3). We compared the amino acid sequence of C31 to the most recent versions in the SWISSPRO, GENPRO, and NBRF data banks with the FASTA alignment program of Pearson and Lipman (34). The best alignment was obtained with the δ subunit (24) of the *Bacillus subtilis* RNA polymerase (initial score of 109 against an average score of 23.9 ± 6.46). Animal HMG1 proteins were also found among the 20 best alignments, with a rather modest initial score of 73. All of the 20 best alignments corresponded to proteins with extensive stretches of acidic amino acids (Asp and Glu) and were limited to these stretches. Data bank searches done with the first 201 amino acids of RPC31 (without the acidic domain) as a probe yielded no appreciable homology to any known protein. We also noted that animal HMG1 proteins were strongly invariant except in their acidic tail, which is almost exclusively formed of Asp residues in the trout protein (35) but contains numerous Glu residues in the mammalian ones (32). No cross-antigenicity was detected between the C31 protein and animal HMG1 proteins by Western blot (immunoblot) analysis (data not shown). Finally, the cruciform DNA-binding activity typical of animal HMG1 proteins (3) was not detected on an in vitro translated C31 protein produced from plasmid pBSC31-5, with in vitro translated rat HMG1 as a positive control (data not shown).

Redundancy of the 3'-terminal part of RPC31 on the yeast genome. Southern hybridization of an RPC31 probe corresponding to the first 189 amino acids (without the acidic tail) yielded a single signal against the S. cerevisiae genome digested with various restriction enzymes (Fig. 4B), establishing that RPC31 is a single-copy gene. There was no cross-hybridization against the S. pombe genome under low-stringency conditions (Fig. 4C), indicating that RPC31 is at best poorly conserved between these two species, unlike the RPC160 gene encoding the largest subunit of RNA polymerase C (data not shown). In contrast, a probe corresponding to the acidic tail of RPC31 yielded numerous cross-hybridization signals against the S. cerevisiae and S. pombe genomes (Fig. 4D), indicating that both yeasts probably harbor numerous genes encoding proteins with highly acidic domains rich in Asp and Glu residues. In retrospect, this observation explains the fortuitous cloning of *RPC31* (ACP2) by Haggren and Kolodrubetz (15), who used a probe encoding a stretch of Asp and Glu residues to screen a yeast genomic library.

Mutagenesis of RPC31. The heterozygous diploid $rpc31\Delta$:: LEU2/RPC31 was constructed by one-step disruption (42) of strain YNN281 × YNN282. The disrupted rpc31 allele (detected by its Leu⁺ phenotype) invariably cosegregated with a lethal phenotype (at most one or two cell divisions upon spore germination on the complete medium YPD at 16 and 30°C) in tetrad analysis. Viable haploid segregants of the $rpc31\Delta$::LEU2 allele were recovered by transforming the heterozygous diploid with a centromeric plasmid bearing the 1.8-kb SpeI-EcoRI (pC50) or the 2.5-kb EcoRI-EcoRI (pC329 or pC326) fragment encompassing the RPC31 gene (Fig. 1B and C). These segregants could not lose the complementing plasmid, showing that the lethal phenotype reflects an inability to undergo vegetative growth and division and not just defective spore germination.

The $rpc31\Delta::LEU2$ allele deletes about 200 bp upstream from the *RPC31* coding region and may prevent the transcription of an essential gene adjacent to *RPC31*, with *RPC31* itself encoding a nonessential product. To eliminate this possibility, we inactivated *RPC31* in gene disruption experiments with the rpc31::Tn10-13 and rpc31::HIS3a insertions (Fig. 1C), which again yielded a recessive lethal phenotype cosegregating with the mutated allele. Further-

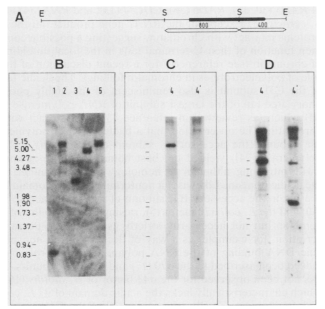


FIG. 4. Southern blot analysis of S. cerevisiae YNN281 and S. pombe h⁻⁹⁷² DNAs against RPC31. (A) Simplified restriction map of the RPC31 region (2.5-kb EcoRI fragment; see Fig. 1B for restriction site symbols). The 800-bp Sall fragment contains the first 189 codons of RPC31 (without the acidic tail). The 400-bp Sall-EcoRI fragment includes the acidic tail (see Fig. 3 for the position corresponding to the SalI restriction site in the amino acid sequence). (B) Stringent hybridization of the S. cerevisiae genome digested with different restriction enzymes against the 800-bp Sall probe. Lane numbers refer to the restriction enzymes used: 1, Sall; 2, NcoI; 3, EcoRI; 4, BamHI; 5, HpaI. (C) Hybridization of S. cerevisiae (lane 4) and S. pombe (lane 4') DNAs (after BamHI digestion) against the 800-bp Sall probe under low-stringency conditions. Filters were washed twice in 4× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate) for 10 min at room temperature and then for 1 h at 55°C. The weak bands observed in the S. pombe DNA were due to a background signal corresponding to the rDNA restriction fragments. (D) Hybridization of the S. cerevisiae (lane 4) and S. pombe (lane 4') DNAs (after BamHI digestion) against the 400-bp SalI-EcoRI probe containing the acidic tail of RPC31 under low-stringency conditions. Numbers at left are in kilobases.

more, we inactivated RPC31 on plasmid pC329 by two additional insertions (rpc31::Tn10-33 and rpc31::Tn10-38) (Fig. 1C) and by six nonsense mutations (Fig. 3) scattered throughout RPC31. The corresponding mutations (including an rpc31-Tyr4 (UAA) nonsense allele at the fourth codon of *RPC31*) all failed to complement $rpc31\Delta$::LEU2, although they were themselves recessive, as established by a plasmid shuffle assay with plasmid pC50 (RPC31) as the complementing plasmid. This result established that the lethal phenotype of the $rpc31\Delta$::LEU2 allele is entirely due to its defective RPC31 gene product. Finally, we noted that the C-terminal rpc31-TYR230 (UAG) allele (which deletes about half of the acidic C-terminal domain) also had a lethal phenotype, whereas the last six amino acids of the acidic tail were not essential, as they could be replaced by six unrelated amino acids by rpc31-HIS3b gene disruption without affecting the growth rate.

Defective tRNA synthesis in a leaky mutant. Two centromeric plasmids (pC329 and pC50) bearing *RPC31* were mutagenized with hydroxylamine, and *RPC31* function was tested by genetic complementation of the $rpc31\Delta::LEU2$ allele in a plasmid shuffle test. We found no conditional (cold- or heat-sensitive) mutant under conditions that yield-

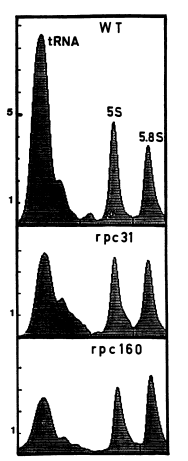


FIG. 5. Synthesis of small RNAs in a defective rpc31 mutant. The incorporation of ³H-labeled uracil into small RNAs was measured after 15 min of in vivo labeling at 30°C in strains YNN281 (wild type [WT]), D27-7c (rpc160-41), and DM824 (rpc31-Tyr4 SUP11-0). RNAs (20 µg) were separated on a 6% polyacrylamide gel and autoradiographed. Uracil incorporation was quantified by densitometry of the autoradiogram after 3 to 10 days of exposure. The proportionality of the signals as a function of the exposure time was checked to avoid artifacts due to signal saturation. The heights of the peaks are given in arbitrary units and are normalized to the 5.8S rRNA signal.

ed 225 fully defective mutants and thus probably corresponded to saturation mutagenesis. These data strongly suggest that temperature-sensitive alleles of *RPC31* are genuinely rare, at least after hydroxylamine mutagenesis. A leaky mutant strain was constructed by cloning the *rpc31*-*Tyr4* (UAA) insert on the centromeric plasmid pSE359, which bears the tyrosine-inserting suppressor SUP11-0. The resulting plasmid (pSE359-Tyr4) generated a wild-type protein with a reduced intracellular concentration due to the low efficiency of nonsense suppression. The resulting *rpc31*\Delta:: *LEU2* (pSE359-Tyr4) mutant (strain DM824) had a growth rate of 6 h on minimal medium supplemented with tryptophan and uracil at 30°C, as compared with 2.5 h for our wild-type strain, YNN281, and with 3.0 h for the *rpc31*\Delta:: *LEU2* (pSE359-RPC31⁺) strain DM824.

We measured the incorporation of ³H-labeled uracil into small RNAs in strain DM824 after a short pulse of 15 min (Fig. 5; similar results were obtained after a 30-min pulse [data not shown]). The *rpc160-41* mutant strain D27-7c (a bona fide RNA polymerase C mutant with a growth rate of 7 h at 30°C [14]) was used as a control. Since tRNAs and rRNAs are comparatively stable molecules, this procedure essentially measures the rate of tRNA synthesis in vivo. To eliminate possible secondary effects of the reduced mutant growth rate per se on RNA polymerase C, we also reduced the growth of the wild-type control strain, YNN281, with glucosamine to compete for glucose uptake (27). The rpc31and rpc160 mutants both had reduced rates of tRNA synthesis (relative to 5.8S rRNA synthesis [Fig. 5]) that were correlated with their reduced growth rates. This result supports the hypothesis that the C31 polypeptide is a functional subunit of RNA polymerase C. In keeping with earlier data on rpc160-41 (14), there was at best a minor effect on the rate of synthesis of 5S rRNA.

DISCUSSION

We cloned a 13.5-kb fragment of the yeast genome that includes the single-copy gene *RPC31*. DNA and protein sequence data demonstrated that it codes for the C31 subunit of RNA polymerase C. Null mutations were constructed, and their lethality established that the integrity of this protein is essential for cellular growth and/or division. Furthermore, the reduced rate of tRNA synthesis of a partially defective mutant supported the hypothesis that the C31 protein functionally belongs to RNA polymerase C. Unlike the genes of the large RNA polymerase subunits (51), *RPC31* is not strongly conserved between the *S. cerevisiae* and *S. pombe* genomes. The gene was precisely located on the left arm of chromosome XIV, very close to an 8-bp *Not*I site. The only gene identified so far in that region (*MAK26* [49]) controls the maintenance of the yeast RNA killer virus.

RPC31 codes for a 251-amino-acid protein with a predicted M_r of 27,707 and an isoelectric point of 4.5. This acidic character is largely due to a C-terminal domain which is characterized by stretches of Asp and Glu residues from positions 202 to 248, which may be partly organized in a helix, and which is likely to protrude from the C31 protein because of its hydrophilic character and C-terminal position. Using an oligonucleotide probe rich in Asp and Glu codons, Haggren and Kolodrubetz (15) isolated the same gene (termed ACP2, for acidic protein 2) which they interpreted as coding for an HMG1 protein. HMG1 proteins are functionally ill-defined components of chromatin which specifically bind cruciform DNA (3), a property that is not shared by the C31 protein. Furthermore, animal HMG1 proteins and the C31 subunit are antigenically unrelated, and there is no sequence homology outside a C-terminal run of Asp and Glu residues which creates a highly acidic domain also present in a number of other proteins (see below). This domain is itself not particularly invariant among animal HMG1 proteins. Taken together, these data strongly argue against the idea that C31 is an HMG1 protein. Interestingly, two S. cerevisiae genes (NHP6A and NHP6B [22]) which code for polypeptides of about 10 kilodaltons were recently shown to have notable homology to the middle region of the animal HMG1 protein amino acid sequence. These polypeptides have no acidic tail and are in fact highly basic. The structure of yeast HMG1 proteins (and their functional relationship to their animal counterparts) remains, therefore, an open question.

The highly acidic C-terminal tail of the C31 protein is functionally or structurally essential, since the elimination of half of this domain by a nonsense mutation inactivated the protein in vivo. A C-terminal acidic domain is present on a variety of proteins, such as the spermine-binding protein (6), the murine homeodomain protein (20), and the gene products of RAD3 (30, 37), RAD6 (36), CDC34 (13), and RNA1 (48) in S. cerevisiae. Except for the RNA1 gene product, all these proteins interact with chromatin, suggesting a possible common function of their C-terminal tails in the local unfolding of chromatin (see reference 7 for a recent discussion of the role of polyelectrolytes in chromatin folding). The acidic tail of the C31 subunit is also reminiscent of the highly phosphorylated tail of the largest subunit of RNA polymerase B (II) which (as reviewed in reference 43) may interact with chromatin. The observation that a bacterial RNA polymerase subunit (the accessory δ subunit of *B*. subtilis RNA polymerase [24]) yielded the best homology score against C31 is intriguing, although the homology was again limited to the acidic domain. Likewise, a nonterminal acidic domain is present in the accessory B32 subunit of RNA polymerase B (50) and the E. coli σ 70 factor (5), raising the possibility of a common but not necessarily strictly essential role in transcription, for example, as a way of decreasing the nonspecific DNA binding of the RNA polymerase. This role has indeed been ascribed to the σ 70 (17) and δ 21 (47) subunits. It has not been observed for the $\sigma 43$ factor of *B*. subtilis (16), which characteristically lacks the acidic domain of its E. coli σ 70 counterpart. Finally, the acidic tail of the C31 protein may directly interact with a basic component of the RNA polymerase C transcription machinery, such as the TFIIIA transcription factor (12).

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