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Sbp4p, an essential putative RNA helicase, is required for a late step in the assembly of 60S ribosomal subunits in *Saccharomyces cerevisiae*

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ABSTRACT

Sbp4p is a putative ATP-dependent RNA helicase that is required for synthesis of 60S ribosomal subunits. Polysome analyses of strains genetically depleted of Sbp4p or carrying the cold-sensitive *spb4-1* mutation revealed an under-accumulation of 60S ribosomal subunits. Analysis of pre-rRNA processing by pulse-chase labeling, northern hybridization, and primer extension indicated that these strains exhibited a reduced synthesis of the 25S/5.8S rRNAs, due to inhibition of processing of the 27SB pre-rRNAs. At later times of depletion of Sbp4p or following transfer of the *spb4-1* strain to more restrictive temperatures, the early pre-rRNA processing steps at sites *A*₀, *A*₁, and *A*₂ were also inhibited. Sucrose gradient fractionation showed that the accumulated 27SB pre-rRNAs are associated with a high-molecular-weight complex, most likely the 66S pre-ribosomal particle. An HA epitope-tagged Sbp4p is localized to the nucleolus and the adjacent nucleoplasmic area. On sucrose gradients, HA-Sbp4p was found almost exclusively in rapidly sedimenting complexes and showed a peak in the fractions containing the 66S pre-ribosomes. We propose that Sbp4p is involved directly in a late and essential step during assembly of 60S ribosomal subunits, presumably by acting as an rRNA helicase.

Keywords: DEAD-box proteins; pre-rRNA processing; ribosome biogenesis; yeast

INTRODUCTION

The synthesis of eukaryotic ribosomes is a complex process that takes place largely, although not exclusively, in a specialized subnuclear compartment termed the nucleolus (Mélèse & Xue, 1995). There, the rDNA is transcribed as precursors (pre-rRNAs), which undergo processing, cleavage, and covalent modification. Concomitantly, these pre-rRNAs assemble with the ribosomal proteins (r-proteins) (reviewed in Woolford & Warner, 1991; Eichler & Craig, 1994; Tollervey & Kiss, 1997).

In *Saccharomyces cerevisiae*, three of four rRNAs (18S, 5.8S, and 25S) are produced as a single 35S pre-rRNA by RNA polymerase I, whereas the fourth rRNA (5S) is transcribed independently by RNA polymerase III (Woolford & Warner, 1991). In the 35S pre-rRNA, the mature rRNA sequences are separated by two internal transcribed spacers, ITS1 and ITS2, and flanked by two external transcribed spacers, 5′ ETS and 3′ ETS (see Fig. 1A). Maturation of this 35S pre-rRNA requires different trans-acting factors, including small nucleolar RNAs (snorRNAs), endonucleases, exonucleases, and RNA modifying enzymes (van Nues et al., 1995; Venema & Tollervey, 1995; Tollervey & Kiss, 1997). Although the pre-rRNA processing pathway has been well-characterized (see Fig. 1B and its legend), the assembly process of the rRNAs and the r-proteins into mature ribosomal subunits is still poorly understood (Woolford & Warner, 1991). In the nucleolus, the pre-rRNAs associate with many of the r-proteins to form pre-ribosomal particles. The first pre-ribosome that can be detected is called 90S and contains the 35S pre-rRNA. From this particle, 66S and 43S pre-ribosomes are formed, containing the 27S and 20S pre-rRNAs, respectively (Trapman et al., 1975). The 66S particle remains in the nucleus for further process-

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FIGURE 1. Pre-rRNA processing in *S. cerevisiae*. **A:** Structure of the 35S pre-rRNA and processing sites. This precursor contains the sequences for the mature 18S, 5.8S, and 25S rRNAs that are separated by the two internal transcribed spacers ITS1 and ITS2, and flanked by two external transcribed spacers, 5’ ETS and 3’ ETS. The location of various probes (numbered from 1 to 9) used in this study are indicated. Bars represent mature rRNA species and lines the transcribed spacers. **B:** Pre-rRNA processing pathway. The 35S pre-rRNA is cleaved at site A0 by the endonuclease Rnt1p, generating the 33S pre-rRNA. This molecule is subsequently processed at sites A1 and A2, resulting in the separation of the pre-rRNAs destined for the small and large ribosomal subunits. The early pre-rRNA cleavages A0 to A2 are proposed to require a large snoRNP complex, which may be assisted by the putative ATP-dependent RNA helicases Dbp4p, Fal1p, Rok1p, and Rrp3p. The final maturation of the 20S precursor takes place in the cytoplasm, where endonucleolytic cleavage at site D yields the mature 18S rRNA. The 27SA2 precursor is processed by two alternative pathways that both lead to the formation of mature 5.8S and 25S rRNAs. In the major pathway, the 27SA2 precursor is cleaved at site A3 by the RNase MRP complex. The putative ATP-dependent RNA helicase Dpb3p assists in this processing step. The 27SA2 precursor is exonucleolytically digested 5’→3’ up to site B1L to yield the 27SBs precursor, a reaction requiring the exonucleases Xrn1p and Rat1p. A minor pathway processes the 27SA2 molecule at site B1S, producing the 27SBs pre-rRNA. While processing at site B1S is completed, the 3’ end of mature 25S rRNA is generated by processing at site B2. The subsequent ITS2 processing of both 27SB species appears to be identical. Cleavage at sites C1 and C2 releases the mature 25S rRNA and the 7S pre-rRNA. The latter undergoes exosome-dependent 3’→5’ exonuclease digestion to the 3’ end of the mature 5.8S rRNA. It has been proposed that Dob1p/Mtr4p, a putative ATP-dependent RNA helicase, assists the exosome activity. The data presented in this study suggest that Spb4p is required for a late step in the assembly of 60S ribosomal subunits, a process that may also involve three other putative ATP-dependent RNA helicases Dpb6p, Dpb7p, and Drs1p.
ing, whereas the 43S pre-ribosome is exported rapidly to the cytoplasm, where the final maturation step in the synthesis of the 18S rRNA takes place (Udem & Warner, 1973; Trapman & Planta, 1976). A large number of r-proteins associate with the nucleolar pre-ribosomes at early steps in ribosome maturation, whereas another group assembles later (Kruiswijk et al., 1978). The available data are, however, far from being sufficient to establish any clear assembly pathway for the different r-proteins. In addition to r-proteins, the nucleolar pre-ribosomes have long been known to contain non-r-proteins (Trapman et al., 1975); the identity of these proteins has not been clearly established, but they presumably correspond to trans-acting factors required for pre-rRNA processing and modification or are involved in the assembly of the pre-rRNAs with the r-proteins.

Among the different trans-acting factors predicted to function enzymatically in ribosome biogenesis are the ATP-dependent RNA helicases. RNA helicases have been found in all organisms studied and are involved in many RNA metabolic processes, including translation initiation, pre-mRNA splicing, ribosome biogenesis, RNA degradation, and poly(A)⁺ RNA export (Schmid & Linder, 1992; Fuller-Pace, 1994; Jacobs Anderson & Parker, 1996; Snay-Hodge et al., 1998; Staley & Guthrie, 1998; Tseng et al., 1998). For several of these proteins, an RNA-dependent ATPase activity has been found (for examples see Kim et al., 1992; O’Day et al., 1996b) and a small number of them have been shown to have an ATP-dependent RNA unwinding activity (for examples, see Rozen et al., 1990; Czapinski et al., 1995; Schwer & Gross, 1998; Tseng et al., 1998; Wang et al., 1998). This class of proteins is therefore generally regarded as putative ATP-dependent RNA helicases. In yeast, the putative RNA helicases Dpb4p, Fal1p, Rok1p, and Rrp3p are required for 18S rRNA synthesis (O’Day et al., 1996a; Kressler et al., 1997; Liang et al., 1997; Venema et al., 1997), whereas Dsb3p, Dpb6p, Dpb7p, and Dsr1p (Ripmaster et al., 1992; Weaver et al., 1997; Daugeron & Linder, 1998; Kressler et al., 1998) have been reported to be involved in the 25S/5.8S rRNA maturation and Dob1p/Mtr4p is required for correct 3’ end processing of the 5.8S rRNA (de la Cruz et al., 1998). In no case is the precise function of these enzymes understood, but, considering their presumed ATP-dependent RNA unwinding activities, RNA helicases are expected to be involved in structural rearrangements within the pre-ribosomal particles during pre-rRNA processing and assembly.

Spb4p is a putative RNA helicase that is essential for cell viability (Sachs & Davis, 1990). The cold-sensitive (cs) mutation spb4-1 was isolated as one of seven different extragenic suppressor of the thermosensitive (ts) phenotype of a poly(A)-binding protein (Pab1p) mutant (pab1-F364L) (Sachs & Davis, 1989). Each of the spb (suppressor of poly(A)-binding protein) mutations, including spb4-1, led to decreased steady-state levels of free 60S ribosomal subunits (Sachs & Davis, 1989). In the case of spb4-1, this was due to reduced synthesis of the 25S rRNA, suggesting that Spb4p could be involved in the biogenesis of 60S ribosomal subunits (Sachs & Davis, 1990). However, the function of Spb4p has not been characterized further.

To learn more about the function of Spb4p, we have studied the subcellular localization of an HA-tagged Spb4p, and we have performed the detailed characterization of both a strain that expresses Spb4p conditionally and the spb4-1 mutant. Our results indicate that Spb4 is a nucleolar protein required for normal processing of the 27SB precursors. We conclude that Spb4 is involved in the correct assembly of pre-ribosomal particles. In the absence of functional Spb4p, maturation of 60S ribosomal subunits is arrested at a late stage, leading to impaired production of mature 25S and 5.8S rRNAs from the 27SB pre-rRNAs, and finally to the depletion of mature 60S relative to 40S ribosomal subunits.

RESULTS

Conditional systems for the phenotypic analysis of Spb4p

The putative RNA helicase Spb4p is essential for cell viability (Sachs & Davis, 1990). Thus, two different conditional systems for phenotypic analysis were used to characterize its function. First, the SPB4 ORF was placed under the control of an inducible GAL promoter, which allows expression of the gene in culture medium that contains galactose (YPGal) but is repressed in culture medium containing glucose (YPD). The resulting plasmid, pAS24-SPB4, expressed an N-terminally HA-tagged Spb4 fusion protein that fully complemented the spb4 null strain (JDY8-1A) on YPGal and resulted in a severe slow-growth phenotype on YPD plates (Fig. 2A). In liquid YPGal medium, there was no difference in the growth rate between the GAL::SPB4 (JDY8-1A pAS24-SPB4) and an isogenic wild-type SPB4 strain (JDY8-1A YCplac111-SPB4). After transfer of the GAL::SPB4 strain from liquid YPGal to YPD medium, the growth rate remained similar to that of the wild-type control strain for the first 12 h (doubling time 1.7 h), but it then decreased progressively to a doubling time of more than 10 h after 36 h in YPD (Fig. 2B). Western blot analysis (Fig. 2C) showed that the decrease in growth rate is preceded by depletion of HA-Spb4p.

In addition, we analyzed the spb4-1 cs mutant (YAS168) (kindly provided by A. Sachs, University of California, Berkeley). This mutant showed a growth defect at 30°C (doubling time 2.9 h for YAS168 versus 1.7 h for JDY8-1A YCplac111-SPB4 in YPD medium) that was enhanced at 18°C (doubling time of more than 12 h for YAS168 versus 5.0 h for JDY8-1A...
YCplac111-SPB4 in YPD medium). Transformation of YAS168 with a CEN-SPB4 plasmid restored a wild-type growth rate, showing the spb4-1 mutation to be recessive (data not shown).

The GAL::SPB4 and spb4-1 strains are deficient in 60S ribosomal subunits

To determine the effects of Spb4p depletion on ribosome metabolism, cell extracts were prepared from the GAL::SPB4 and SPB4 strains and subjected to sucrose gradient centrifugation. Polysome profiles and ratios of total ribosomal subunits were analyzed. As shown in Figure 3, depletion of Spb4p is accompanied by an overall decrease in polysomes, a deficit in free 60S ribosomal subunits, and the appearance of half-mer polysomes (Fig. 3A,B,C,D). Quantification of total ribosomal subunits in low Mg²⁺ sucrose gradients indicated a deficit of total 60S ribosomal subunits relative to 40S subunits; an A₂₅₄ 60S-to-40S ratio of around 2 was observed for SPB4 in YPD or for GAL::SPB4 grown in YPGal. This ratio decreased to around 1.2 after transfer of GAL::SPB4 to YPD for 24 h (data not shown).

Polysome and total ribosomal subunit analyses were also performed with cell extracts prepared from spb4-1 cells grown in YPD at permissive temperature (30 °C) or 12 h after transfer to the nonpermissive temperature (18 °C). The spb4-1 mutation also led to a deficit of free 60S ribosomal subunits and the appearance of half-mer polysomes, the defect being more drastic after the shift to 18 °C (Fig. 3E,F). This defect can be fully reversed at either temperature by complementation of the spb4-1 strain with a CEN plasmid harboring the wild-type SPB4 gene (data not shown). When A₂₅₄ 60S-
to-40S ratios were determined, the spb4-1 mutant gave values of 1.5 (30°C) and 1.3 (18°C) compared to a value of 1.9 for the spb4-1 strain complemented by a CEN-SPB4 plasmid at either temperature.

Taken together, polysome and ribosomal subunit analyses indicated that the depletion of Spb4p and the spb4-1 mutation led to similar reductions in the levels of 60S ribosomal subunits.

Formation of the mature 25S and 5.8S rRNAs is impaired upon Spb4p depletion

To further characterize the role of Spb4p in the metabolism of 60S ribosomal subunits, we first analyzed the effects of Spb4p depletion on synthesis and processing of pre-rRNA by [methyl-3H]methionine pulse-chase labeling experiments. For this purpose, JDY8-1A pAS24-SPB4 and JDY8-1A YCplac111-SPB4 were pre-grown in YPGal, then transferred to YPD for 13 h, and finally grown for 9 h (to an OD600 of around 1) in synthetic medium lacking methionine (SD-Met). At this time point, the GAL::SPB4 strain was doubling every 4.5 h compared with 2.5 h for the wild-type strain. The cells were pulse-labeled for 1 min, then chased for 2, 5, and 15 min with an excess of cold methionine and total RNA was extracted and analyzed. In the wild-type SPB4 strain, the 35S precursor was processed rapidly into 32S pre-rRNA and then into 27S and 20S species, the immediate precursors of the mature 25S and the 18S rRNA, respectively (Fig. 4A, lanes 1–4). In the Spb4p-depleted strain, we observed that processing of the 35S precursor was delayed (Fig. 4A, lane 5). Less 27SA species was formed, and the 27SB precursors per-
sisted after the 15-min chase time point (Fig. 4A, lanes 5–8). As a consequence, almost no labeled mature 25S rRNA was detected (Fig. 4A, lane 8). The processing pathway leading to the formation of 18S rRNA was also mildly impaired, as revealed by the lower levels of mature 18S rRNA (Fig. 4A, lanes 7, 8) and its 20S precursor (Fig. 4A, lanes 5, 6). An aberrant 23S species was also detected (Fig. 4A, lane 6). However, the kinetics of processing of the 20S precursor to the mature 18S rRNA was not affected.

To exclude a defect in RNA methylation and to monitor the synthesis of low-molecular-weight rRNAs, cells were also pulse-labeled with [5,6-3H]uracil. Strains JDY8-1A pAS24-SPB4 PRS416 and JDY8-1A YCp50-SPB4 were pre-grown in SGal-Ura and then transferred to SD-Ura for 22 h (to an OD600 of around 1).

At this time point, the GAL::SPB4 strain was doubling every 3.6 h compared with 2 h for the isogenic SPB4 strain. The cells were pulse-labeled for 2 min and then chased for 15, 30, 50, and 60 min with an excess of cold uracil. Analysis of high-molecular-weight RNAs gave results comparable to those shown in Figure 4A (data not shown). Analysis of low-molecular-weight RNAs showed that the synthesis of mature 5.8S rRNAs was both substantially delayed and strongly reduced following Spb4p depletion. In contrast, synthesis of the 5S rRNA and tRNAs was comparable both in kinetics and efficiency in wild-type and Spb4p-depleted cells (Fig. 4B).

These results indicated that the deficit in 60S ribosomal subunits following Spb4p depletion was due to the inhibition of processing of the 27SB precursors to mature 25S and 5.8S rRNAs. A defect in earlier processing reactions on the pathway of 18S rRNA synthesis was also observed.

Pre-rRNA processing is inhibited by depletion of Spb4p

To define the pre-rRNA processing steps that are affected in the conditional spb4 strains, steady-state levels of mature rRNAs and pre-rRNA intermediates were assessed by northern and primer extension analyses. Total RNA was isolated from the GAL::SPB4 and an isogenic SPB4 strain at various time points after transfer from YPGal to YPD and analyzed by northern hybridization. As shown in Figure 5A, depletion of Spb4p resulted in a slight decrease in 18S rRNA and in a more drastic decrease in 25S rRNA. Oligonucleotides, hybridizing to sites in the 35S pre-rRNA transcript (see Fig. 1A), were used to identify the processing intermediates. Slight differences in the levels of the pre-rRNAs observed between the 0 h and 36 h time points for the wild-type strain are due to the effects of a nutritional up-shift from galactose to glucose medium; comparable increases in the levels of several pre-rRNA species are seen in the GAL::SPB4 strain at early times (6 h and 12 h) after transfer to glucose medium. Consistent with the pulse-chase labeling results, the GAL::SPB4 strain accumulated the 35S pre-rRNA; clear accumulation was observed 12 h after transfer to glucose and increased with time (Fig. 5B,C,D,E,F, lanes 5–8). The 27SB pre-rRNA was strongly accumulated at 12 h and at later time points (shown for probe 8 in Fig. 5F; probe 7 gave identical results, data not shown). The aberrant 23S RNA also appeared, commencing 18 h after transfer to glucose medium (Fig. 5B,C,D, lanes 6–8). This species extends from the 5’ ETS to site A3, as shown by its hybridization with probes 1 and 4 (Fig. 5B,D), but not with oligonucleotide 5 (Fig. 5E). The levels of the 20S (Fig. 5C, lanes 7, 8) and 27SA2 pre-rRNAs (Fig. 5D,E, lanes 7, 8) are reduced in the Spb4p-depleted strain, commencing 18 h after transfer to glucose medium. However, 12 h after transfer to glucose, 27SA pre-rRNAs accumulated (Fig. 5D,E, lane 5). Analysis of low-molecular-weight rRNA species revealed a slight decrease in the steady-state levels of the 7S pre-rRNA and 5.8S rRNAs, but no alteration in the levels of 5S rRNA in the Spb4p-depleted strain (data not shown). The ratio of 5.8Sa and 5.8Sz rRNAs was not affected by Spb4p depletion (data not shown).

We conclude that, approximately 12 h after transfer of the GAL::SPB4 strain to glucose medium, pre-rRNA processing was inhibited at sites C1 and C2. Approximately 18 h after transfer to glucose, cleavage started to be inhibited at the early processing sites A0, A1, and A2, with increasing inhibition at later time points.

Because northern hybridization does not distinguish between the 27SA2 and 27SA3, and between the 27SB1 and 27SB2 precursors, we assessed the levels of these species by primer extension analyses, using oligonucleotides 5 and 8 as primers (see Fig. 1A). The primer extension stops at sites B1L and B1S increased at 12 h and later times after transfer to glucose medium (Fig. 6, upper panel), indicating that both 27SB1 and 27SB2 were accumulated upon depletion of Spb4p. The level of the primer extension stop at site A2, the 5’ end of the 27SA1 pre-rRNA, increased 12 h after transfer to glucose medium (Fig. 6, upper panel, lane 5), but was reduced strongly at later time points (Fig. 6, upper panel, lanes 7, 8), fully consistent with the results obtained by northern hybridization. A similar observation was made for site A3, the 5’ end of the 27SA3 pre-rRNA (Fig. 6, bottom panel, lanes 5, 7, and 8). The level of the 27SA3 pre-rRNA was, however, reduced to a lesser extent than 27SA2. Finally, primer extension analysis also showed that processing at all these sites was accurate at the nucleotide level during the time course of Spb4p depletion.

Similar hybridization and primer extension analyses were performed on total RNA extracted from the spb4-1 mutant and from a SPB4 control strain grown at 30°C or shifted for 12 h to 18°C. These analyses revealed that the spb4-1 mutation led at 30°C and at 18°C to
effects similar to those observed after 12 h and 24 h of Spb4p depletion, respectively. At 30 °C, there was an accumulation of the 27SA and 27SB pre-rRNA species in spb4-1 cells, but no clear alteration in other precursors. After a shift to 18 °C for 12 h, the spb4-1 strain showed an accumulation of the 35S and 23S pre-rRNAs and strong accumulation of both 27SB pre-rRNAs. The levels of the 20S and 27SA precursors were reduced markedly, whereas levels of the 7S pre-rRNA and the 5′S and 25S rRNAs were only reduced mildly (data not shown).

Together, these data demonstrated that depletion of Spb4p and the sbp4-1 mutation led to similar pre-rRNA processing defects. At early times of depletion or with the spb4-1 strain at 30 °C, processing of the 27SB pre-rRNAs at C1 and C2 was inhibited, leading to depletion of the mature 25S and 5.8S rRNAs. At later times of depletion or with the spb4-1 strain at 18 °C, pre-rRNA cleavages at sites A0, A1, and A2 were also inhibited.

The 27SB pre-rRNAs accumulate in pre-ribosomal particles

In wild-type cells, the 27SB pre-rRNAs are present in 66S pre-ribosomal particles (Trapman et al., 1975). We therefore analyzed whether these particles assemble in the absence of functional Spb4p. Cells from spb4-1 and SPB4 strains were grown in YPD at 30 °C and shifted for 12 h to 18 °C. Cell extracts were fractionated on sucrose gradients containing a low concentration of Mg2+, in which ribosomes and polysomes dissociate into free ribosomal subunits (Foiani et al., 1991). RNA was recovered from each fraction and pre-rRNAs and mature rRNAs were analyzed by northern hybridization. In both wild-type and spb4-1 cell extracts, the mature 25S and 18S rRNAs were found at positions on the gradients corresponding to the 60S and 40S peaks, respectively (Fig. 7A,B, fractions 5 and 7–8, respectively). The 20S pre-rRNA co-fractionated with the ma-
Role of Spb4p in 60S ribosomal subunit synthesis

Depletion of Spb4p leads to higher steady-state levels of the 27SB precursors. The strains JDY8-1A YCplac111-SPB4 (SPB4) and JDY8-1A pAS24-SPB4 (GAL::SPB4) were grown in YPGal and shifted for up to 36 h to YPD. Cells were harvested at the indicated times and total RNA was extracted. Primer extension with oligonucleotide 5′ to site A2 and reveals the processing at this site. Arrows indicate the positions of the primer extension stops corresponding to the different pre-rRNA species analyzed.

FIGURE 6. Depletion of Spb4p leads to higher steady-state levels of the 27SB precursors. The strains JDY8-1A YCplac111-SPB4 (SPB4) and JDY8-1A pAS24-SPB4 (GAL::SPB4) were grown in YPGal and shifted for up to 36 h to YPD. Cells were harvested at the indicated times and total RNA was extracted. Primer extension with oligonucleotide 5′ to site A2 and reveals the processing at this site. Arrows indicate the positions of the primer extension stops corresponding to the different pre-rRNA species analyzed.

were present in the form of high-molecular-weight complexes, most likely corresponding to the previously identified 90S, 66S, and 43S pre-ribosomal particles, respectively (Trapman et al., 1975). In the spb4-1 mutant, the 27SB pre-rRNAs accumulated in a pre-ribosomal particle, which did not clearly differ in sucrose gradient mobility from the wild-type particle. It has been suggested that the 90S pre-ribosomes are split into the 66S and 43S pre-ribosomal particles after cleavage of the 35S pre-rRNA at site A2 (van Nues et al., 1995). No co-sedimentation of the 20S and 27SA/B pre-rRNAs was observed, indicating that the 66S and 43S pre-ribosomal particles dissociate rapidly on cleavage of the 35S pre-rRNA. Our results also showed that the aberrant 23S rRNA is found in a particle of higher sedimentation coefficient than 43S, but lower than 90S, presumably indicating that this particle remains associated with processing and/or assembly factors that dissociate from the 20S pre-rRNA following cleavage.

Spb4p is required for processing of the 27SB pre-rRNAs within the 66S pre-ribosomal particles. We therefore determined whether Spb4p is also a component of these particles. For this purpose, Spb4p was HA-tagged at its N-terminus and expressed from its authentic promoter on a CEN plasmid (YCplac111-HA-SPB4). This plasmid complemented the spb4 null allele to the wild-type extent at all temperatures tested (18, 30, and 37°C), indicating that the HA-Spb4 fusion protein was fully functional. Western-blot analysis with a mouse monoclonal anti-HA antibody detected a single protein with a molecular mass of 85 kDa in a total cell extract from the strain expressing the HA-tagged Spb4p (JDI8-1A YCplac111-HA-SPB4), but not from the SPB4 strain (JDI8-1A YCplac111-SPB4) (data not shown). Sucrose gradient fractionation was performed with cell extracts from the strain expressing HA-Sbp4p and fractions were analyzed by western blot analysis. Ssm1p, which is a 60S r-protein (Petitjean et al., 1995), was used as a marker for the sedimentation of 60S ribosomal subunits. The small nucleolar ribonucleoprotein (snoRNP) protein Nop1, which is expected to be associated with the pre-ribosomal particles, was also used as a control. As shown in Figure 8, Ssm1p was found exclusively associated with fractions corresponding to the mature 60S ribosomal subunits and the putative 66S pre-ribosomal particle (Fig. 8, fractions 6–8). Little HA-Sbp4p or Nop1p sedimented at the positions expected for the free proteins (Fig. 8, fractions 1 and 2), although overexposure of the blots indicated the existence of very minor pools of both proteins (data not shown). A peak of HA-Sbp4p was found in fractions corresponding to the 66S pre-ribosomal particle (Fig. 8, fractions 7 and 8); these fractions also contained the peak of the 27SB pre-rRNAs (data not shown). Lower amounts of HA-Sbp4p were detected lower on the gradient, in fractions that contain the 35S pre-rRNA, presumably corresponding to the 90S pre-
ribosomes (Fig. 8, fractions 9–11). Both HA-Spb4p and Nop1p were also found in fractions above the 40S subunits. In the case of Nop1p, this presumably represented the population of snoRNPs that was not engaged with the 35S pre-rRNA at the moment of cell lysis.

We conclude that HA-Spb4p is almost entirely present in relatively large complexes in cell lysates. The sedimentation of these overlapped the positions of the 60S ribosomal subunits and 66S pre-ribosomes, and is similar to that of Nop1p. These data are consistent with the

**FIGURE 7.** The 27SB pre-rRNAs accumulate in 66S pre-ribosomal particles. Wild-type (YAS168 YCplac111-SPB4) (A) and spb4-1 (YAS168) cells (B) were grown at 30 °C in YPD and shifted to 18 °C for 12 h. Cells were harvested at an OD<sub>600</sub> of 0.8. Cell extracts were resolved in 7–50% sucrose gradients containing a low concentration of Mg<sup>2+</sup> to dissociate ribosomes into subunits. The A<sub>254</sub> was measured continuously. Sedimentation is from left to right. The peaks of free material and total 40S and 60S ribosomal subunits are indicated. Fractions were collected and RNA was extracted from each fraction. T stands for total extract and numbers indicate the fraction numbers. Equal volumes were resolved on a 1.2% agarose–6% formaldehyde gel and transferred to a nylon membrane for northern hybridization. The same filters were hybridized consecutively with different probes to detect the different pre-rRNAs and mature rRNAs indicated by the arrows.
Role of Spb4p in 60S ribosomal subunit synthesis

To determine whether Spb4p was also associated with the mature 60S ribosomal subunits, we identified its subcellular localization. Indirect immunofluorescence was performed with JDY8-1A YCplac111-HA-SPB4 by using anti-HA antibodies (Fig. 9B). For subnuclear localization, the nucleolus was visualized with anti-Nop1p antibodies (Fig. 9A) and the nucleoplasm was visualized by staining the DNA with DAPI (Fig. 9C). Anti-Nop1p antibodies gave the crescent-shaped staining characteristic of nucleolar proteins (Fig. 9A), which was largely excluded from the DAPI stained area (Fig. 9F). The HA-tagged Spb4p was restricted to the nucleus (Fig. 9B) and detected in the nucleolus as shown by its colocalization with Nop1p (Fig. 9D, overlap in yellow). However, in contrast to Nop1p, HA-Spb4p was also detected throughout the nucleoplasm, as shown by its colocalization with the DAPI staining (Fig 9E, overlap in magenta). No anti-HA signal was observed using a nontagged control strain (JDY8-1A YCplac111-SPB4) (data not shown).

On the basis of these results, we conclude that Spb4p is a nuclear protein that localizes to the nucleolus and to the adjacent nucleoplasm. No cytoplasmic signal was detected, indicating that Spb4p is not associated with the mature 60S ribosomal subunits. Although the HA-Spb4 fusion protein is expressed under its own promoter from a low copy number CEN plasmid, some overexpression remains possible. A partial loss of function (without negative effects on growth) due to tagging might also account for the nonexclusively nucleolar localization. In any event, the presence of Spb4p in the nucleolus strongly supports a direct role for Spb4p in 60S ribosomal subunit synthesis.

DISCUSSION

In this paper, we describe the functional analysis of Spb4p. Polysome analyses and quantification of total ribosomal subunits in both a GAL::SPB4 and the spb4-1 strain revealed a deficit in 60S ribosomal subunits leading to the appearance of half-mer polysomes. These profiles are characteristic of mutants defective in 60S r-proteins or in trans-acting factors involved in its biogenesis (Kressler et al., 1998 and references therein). Detailed analyses of pre-rRNA processing by pulse-chase labeling, northern hybridization, and primer extension showed that this deficit is attributable to reduced synthesis of mature 25S and 5.8S rRNAs, as a consequence of the inhibition of pre-rRNA processing in the GAL::SPB4 and the spb4-1 strains.

The most striking pre-rRNA processing defect in these strains is the inhibition of the processing of the 27SB, and 27SB₃ pre-rRNAs to mature 25S and 5.8S rRNA. The 27SB pre-rRNAs are normally processed at sites C₁, at the 5' end of the 25S rRNA, and C₂ in ITS2 (see Fig. 1). These processing reactions separate the 25S rRNA from the 7S precursor to the 5.8S rRNA and are the least well understood steps in the yeast pre-rRNA processing pathway. It is not even clear whether processing at both sites is due to endonucleolytic cleavage, or whether one site is processesd by an exonuclease activity from the other. In the Spb4p-depleted strain, the 27SB pre-rRNAs show substantial accumulation as judged by northern hybridization or primer extension and are stabilized as judged by pulse-chase labeling, demonstrating that processing at sites C₁ and C₂ is inhibited. It is not clear whether Spb4p plays a direct role in processing at sites C₁ and C₂ or whether this inhibition is an indirect consequence of defects in the assembly of the pre-ribosomal particle.

In addition to these effects on the synthesis of 25S and 5.8S rRNAs, we also observed an accumulation of the 35S pre-rRNA, the appearance of an aberrant 23S product, and a reduction in the levels of the 27SA and 27SB pre-rRNAs.
20S pre-rRNAs upon depletion of Spb4p or after a shift to 18°C of the spb4-1 mutant. These effects are characteristic of the inhibition of pre-rRNA processing at sites A₀, A₁, and A₂ on the pathway to 18S rRNA synthesis. Similar effects have been reported previously for several other mutants affecting synthesis of the 60S ribosomal subunits. It has been proposed that this inhibition of the early processing steps is the consequence of a feedback mechanism that slows production of the 18S rRNA when the formation of 25S/5.8S rRNA is inhibited (further discussed in Zanchin et al., 1997; Daugeron & Linder, 1998; Kressler et al., 1998). This delay in 35S pre-rRNA processing seems, however, to have only a minor effect on the overall synthesis of 18S rRNA or the steady-state levels of mature 40S ribosomal subunits in the spb4-affected strains. The role of Spb4p in 25S/5.8S rRNA synthesis appears to be specific because neither the spb4-1 mutation nor depletion of Spb4p significantly affects the synthesis of 5S rRNA and tRNAs or rRNA methylation.

Sucrose gradient fractionation of spb4-1 cell extracts revealed that the accumulated 27SB pre-rRNAs are present within pre-ribosomal particles that sediment faster than the mature 60S ribosomal subunits. In wild-type cell extracts, the 27SB pre-rRNAs are found in 60S pre-ribosomes with similar sedimentation rates (Trapman et al., 1975). Sucrose gradient analysis using an HA-Spb4 fusion protein showed that almost all Spb4p in cell extracts is present in high-molecular-weight complexes. The sedimentation of these was consistent with the physical interaction of Spb4p with the 66S and 90S pre-ribosomal particles. The population of HA-Spb4p that is not associated with the pre-ribosomal particles, as shown by its sedimentation above the 40S subunits, is clearly not the free protein. Spb4p may, therefore, be assembled with other pre-rRNA processing factors prior to its association with the pre-ribosomes. The association of Spb4p with the pre-ribosomal particles is supported by indirect immunofluorescence, which revealed that HA-Spb4p is localized to the nucleus with some nucleolar enrichment. Spb4p is not, therefore, a structural component of the 60S ribosomal subunit. The nucleoplasmic signal might represent association of Spb4p with pre-ribosomes in transit to the nuclear pores, but could also be an artifact of the expression of the HA-Spb4 fusion protein.

In many characterized mutants that are defective in synthesis of the 60S subunits, most r-proteins and the 27S pre-rRNAs are degraded rapidly. This is seen on depletion of some 60S subunit r-proteins, e.g., L16 (Moritz et al., 1990), or with mutations in trans-acting factors that are believed to function at early steps in the 60S ribosomal subunit assembly pathway, e.g., Nop4p/Nop77p, Dpb6p or Dpb7p (Bergès et al., 1994; Sun & Woolford, 1994; Daugeron & Linder, 1998; Kressler et al., 1998). In contrast, the accumulation of the 27SB pre-rRNAs observed in the spb4-1 and GAL::SPB4 strains most resembles the phenotypes described for strains depleted of Nop2p, Nop3p, or Nip7p (Russell & Tollervey, 1992; Hong et al., 1997; Zanchin et al., 1997). We speculate that Spb4p, as well as Nop2p, Nop3p, and Nip7p, are required for a late step in the 60S ribo-
somal subunit assembly. In the absence of the activity of these factors, the pre-ribosomes accumulate in a form that is relatively close to the normal structure, leading to their co-sedimentation with the normal 66S pre-ribosomes. Some structural defects, however, prevent the processing reactions at sites C₁ and C₂ from proceeding, preventing synthesis of the 25S and 5.8S rRNAs. In view of the presumed ATP-dependent RNA helicase activity of Spb4p, we assume that it normally functions in the restructuring of the pre-rRNA within the 66S pre-ribosome.

Ten putative ATP-dependent RNA helicases have been reported to function in ribosome biogenesis (see Introduction). Because most of the genes are essential for viability, we conclude that each of these proteins plays a specific, nonredundant function during ribosome biogenesis. This interpretation is supported by the fact that overexpression of other genes encoding putative RNA helicases required for the 60S ribosomal subunit synthesis (DBP6, DBP7, and DRS1) do not suppress either the slow-growth phenotype or the cold-sensitivity of the spb4-1 mutant and do not bypass the lethal phenotype of the spb4 null strain (data not shown). In addition, overexpression of SPB4 does not suppress the slow-growth and the temperature sensitivity of dbp6 (D. Kressler & P. Linder, unpubl.) or dbp7 mutants (Daugeron & Linder, 1998) and double spb4-1 dbp6 or spb4-1 dbp7 mutants are not synthetically lethal (Daugeron & Linder, 1998 and data not shown). During the processing and modification of pre-rRNA and its assembly with the approximately 80 r-proteins, extensive structural rearrangements must occur. Furthermore, the pre-rRNAs must associate with, and dissociate from, a host of snoRNA species, some of which form very extensive base pairing with the rRNA sequences. In most cases, the rRNA–snoRNA interaction is mutually exclusive with the final folding of the rRNA in the ribosome. We therefore anticipate that more RNA helicases required for ribosome synthesis remain to be identified. In no case is the specific substrate known and determining all of their functions will be a significant challenge.

MATERIALS AND METHODS

Strains, media, and microbiological methods

The S. cerevisiae strains YAS168 (MATα, spb4-1 his3 leu2 trp1 ura3) and YAS325 (MATa/MATα, spb4::TRP1/SPB4 ade2/ade2 his3/his3 leu2/leu2 trp1/trp1 ura3/ura3) have been described previously (Sachs & Davis, 1989, 1990). JDY8-1A (MATα, spb4::TRP1) is a meiotic segregant of YAS325 that requires a plasmid-borne copy of SPB4 for cell viability. Genetic manipulations and preparation of standard media were according to established procedures (Ausubel et al., 1994; Kaiser et al., 1994). Yeast cells were transformed using a lithium acetate method (Gietz et al., 1992). Escherichia coli DH10B was used for all recombinant DNA techniques (Sambrook et al., 1989).

Plasmids

Plasmid YCp50-SPB4 (CEN-URA3) has been described previously (Sachs & Davis, 1990). A 2.8-kb BamH I–Sal I fragment from YCp50-SPB4 was subcloned into YCplac111 (CEN-LEU2) and YEplac181 (2μm-LEU2) (Gietz & Sugino, 1988) to generate YCplac111-SPB4 and YEplac181-SPB4, respectively. Plasmid YEplac181-DBP6 (Kressler et al., 1998) and YEplac181-DBP7 (Daugeron & Linder, 1998) have been described. The 2μm-DRS1 plasmid was a generous gift from Dr. J.L. Woolford (Carnegie Mellon University, Pittsburgh, Pennsylvania).

Construction of a GAL::SPB4 allele and in vivo depletion of Spb4p

The SPB4 gene was PCR amplified (Vent Polymerase, New England Biolabs) from YCplac111-SPB4 using the reverse primer (5′-AAC AGC TAT GAC CAT G-3′) and a oligonucleotide introducing the restriction site Sal I (5′-GCA GAA TTC GTC GAG TCA AAG TCA TTG GAA TGG GA-3′); the Sal I site is underlined, and the SPB4 ORF homology, starting with the second codon, is in bold). The PCR product was cut with Sal I and cloned into the Sal I-restricted YCplac111-based plasmid pAS24 (Schmidt et al., 1997). Correct orientation of the Sal I-fragment was verified by restriction analysis. The resulting construct, pAS24-SPB4, contains a GAL1-10 promoter, a start codon followed by a double HA-tag, and the SPB4 ORF and its 3′ contiguous region. This plasmid was transformed into the strain JDY8-1A YCp50-SPB4. The subsequent counter-selection of the URA3 SPB4 harboring plasmid on 5-FOA plates containing galactose resulted in the strain JDY8-1A pAS24-SPB4. The plasmid pAS24-SPB4 complemented the spb4 null strain to the wild-type extent on medium containing galactose (YPGal) at all the tested temperatures (18, 30, and 37 °C). We also refer to this strain as the GAL::SPB4 strain or, if grown in medium containing glucose (YPD), as the Spb4p-depleted strain.

For in vivo depletion of Spb4p, JDY8-1A pAS24-SPB4 was grown in liquid YPGal until mid-exponential phase. Cells were harvested, washed, and used to inoculate YPD cultures. Cell growth was monitored over a period of 36 h, during which the cultures were regularly diluted into fresh YPD medium to maintain exponential growth. As a control, JDY8-1A YCplac111-SPB4 was used. Samples for western blot analyses, polysome analyses, and RNA extraction were taken at different times.

Sucrose gradient analyses

Polyribosome preparations, polysome analyses, and ribosomal subunit preparations were done according to Foiani et al. (1991), exactly as described (Kressler et al., 1997). Gradient analysis was performed using an ISCO UV-6 gradient collector and monitored continuously at A_{260}.

For fractionation analyses, low Mg^{2+} gradients identical to those used for ribosomal subunit quantification were pre-
pared. After centrifugation, fractions of 0.5 mL were collected manually. For RNA analysis, 0.4 mL of each fraction was adjusted to a final concentration of 10 mM Tris-HCl, pH 7.5, 10 mM EDTA, and 0.5% SDS. Total RNA was isolated by two consecutive extractions with 10 mM Tris-HCl, pH 7.5, saturated phenol:chloroform:isoamyl alcohol (25:24:1), followed by a chloroform:isoamyl alcohol (24:1) extraction. Then, RNA was precipitated with ethanol in the presence of 0.3 M sodium acetate, pH 5.2. Finally, RNA pellets were dissolved in 20 μL of distilled water and 5 μL was resolved on 12% agarose–6% formaldehyde gels as described (Venema et al., 1998). For western analysis, proteins (0.1 mL of each fraction) were precipitated by addition of trichloroacetic acid to a final concentration of 10%, followed by incubation on ice for at least 10 min. Proteins were pelleted by centrifugation in a microfuge for 10 min at 4 °C. Pellets were washed twice with 1 mL ice-cold acetone and finally resuspended in 20 μL of protein gel loading buffer (Ausubel et al., 1994). Aliquots of 10 μL were loaded onto SDS-polyacrylamide gels and analyzed by western blot analysis according to standard procedures (Sambrook et al., 1989; Ausubel et al., 1994). Monoclonal mouse anti-HA 16B12 (BabCo), monoclonal mouse anti-Nop1p (Aris & Blobel, 1988), and polyclonal rabbit anti-Ssm1p (Petitjean et al., 1995) antibodies were used as primary antibodies. Blots were decorated with alkaline phosphatase conjugated secondary antibodies (Bio-Rad).

**SPB4 HA epitope tagging and cloning under the control of its cognate promoter**

To express an N-terminally HA-tagged Spb4 fusion protein from its cognate promoter at approximately wild-type levels, the *SPB4* promoter was PCR amplified from YCplac111-SPB4 with the universal primer (5’-GTA AAA CGA CGG CCA GT-3’) and an oligonucleotide introducing an Xba I site (5’-CGG TAG TCT AGA TGC CAT TGT TTT TAT TGC TGC TG-3’; the Xba I site is underlined, the reverse complement of the start codon is in bold and underlined, and the reverse complement of the *SPB4* promoter sequence is in bold). The PCR product was digested with EcoRI I and XbaI I, and cloned together with the XbaI BamHI I-released HA tag of plasmid pAS24 into the EcoRI BamHI I-restricted plasmid pAS24-SPB4. Then, the 1.6-kb Spb I fragment from the above plasmid was replaced by the same fragment from YCplac111-SPB4. The *SPB4* ORF sequences still originating from the PCR were verified by sequencing. The resulting plasmid, YCplac111-HA-SPB4, complemented the *spb4* null allele to the wild-type extent at all the tested temperatures (18, 30, and 37 °C) and the HA-tagged Spb4p was detected by western blotting as a band that migrated at the molecular mass of approximately 85 kDa.

**Indirect immunofluorescence**

The strains JDY8-1A YCplac111-HA-SPB4 and JDY8-1A YCplac111-SPB4 were grown up to an OD_{600} of around 0.5 in YPD medium and 2.5 OD_{600} units were harvested by centrifugation. Preparations of yeast cells for immunofluorescence were done according to standard procedures (Pringle et al., 1991). DAPI (4’,6-diamidino-2-phenylindole dihydrochloride, Fluka) was used to stain DNA. Primary monoclonal mouse anti-HA antibodies 16B12, at a dilution of 1:200, and secondary goat anti-mouse rhodamine conjugated antibodies (Pierce), at a dilution of 1:200, were used to detect HA-Spb4p. Polyclonal rabbit anti-Nop1p antibodies at a dilution of 1:500, and secondary goat anti-rabbit fluorescein conjugated antibodies (Pierce), at a dilution of 1:200, were used to detect the nucleolar protein Nop1p (Tollervey et al., 1991). Fluorescently labeled cells were inspected in a Zeiss Axioskop fluorescence microscope using the Plan-NEOFLUAR 100×/1.3 objective, and images were acquired with a cooled CCD camera (Princeton Instruments) controlled by a Power Macintosh 8500/100 computer. Images were adjusted to similar output intensities, pseudo-colors were assigned and images were merged using Adobe Photoshop 3.0. Figures were printed on a Kodak Digital Science 8650 PS Color Printer.

**RNA analyses**

Pulse-chase labeling of pre-rRNA was performed as described previously (Kressler et al., 1998), using 250 μCi [methyl-3H]methionine (Amersham, 70–85 Ci/mmol) or 100 μCi [5,6-3H]uracil (Amersham, 45–50 Ci/mmol) per 40 OD_{600} units of yeast cells. Total RNA was extracted by the acid-phenol method (Ausubel et al., 1994).

Steady-state levels of pre-rRNAs were assessed by northern and primer extension analyses according to Venema et al. (1998). Total RNA was extracted as above from aliquots of 10 OD_{600} of exponentially growing cells. Oligonucleotides 5’A0, 18S, D/A2, A2/A3, A3/B1, 5.8S, E/C2, C1/C2, 25S (as numbered from 1 to 9 according to Fig. 1A), and 5S have been described previously (de la Cruz et al., 1998). Prior to northern hybridization or primer extension analysis, they were end labeled with 30 μCi [γ-32P]ATP (Amersham, 5,000 Ci/mmol) using T4 Polynucleotide kinase (Appligene).

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**REFERENCES**

Role of Sbp4p in 60S ribosomal subunit synthesis


