Cotranscriptional recruitment of the nuclear poly(A)-binding protein Pab2 to nascent transcripts and association with translating mRNPs

Caroline Lemieux and François Bachand*

RNA Group, Department of Biochemistry, Université de Sherbrooke, Québec, Canada

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ABSTRACT

Synthesis of the pre-mRNA poly(A) tail in the nucleus has important consequences on the translational activity of the mature mRNA in the cytoplasm. In most eukaryotes, nuclear polyadenylation of pre-mRNAs is thought to require the nuclear poly(A)-binding protein (PABP2/PABPN1) for poly(A) tail synthesis and ultimate length control. As yet, however, the extent of the association between PABP2 and the exported mRNA remains poorly understood. Here, we used chromatin immunoprecipitation (ChIP) assays to show that the fission yeast ortholog of mammalian PABP2 (Pab2) is cotranscriptionally recruited to active genes. Notably, the association of Pab2 to genes precedes that of a typical 3'-processing/polyadenylation factor, suggesting that Pab2 recruitment during the transcription cycle precedes polyadenylation. The inclusion of an RNase step in our ChIP and immunoprecipitation assays suggests that Pab2 is cotranscriptionally recruited via nascent mRNA ribonucleoprotein (mRNPs). Tandem affinity purification coupled with mass spectrometry also revealed that Pab2 associates with several ribosomal proteins as well as general translation factors. Importantly, whereas previous results suggest that the nuclear poly(A)-binding protein is not present on cytoplasmic mRNAs, we show that fission yeast Pab2 is associated with polysomes. Our findings suggest that Pab2 is recruited to nascent mRNPs during transcription and remains associated with translated mRNPs after nuclear export.

INTRODUCTION

Two evolutionarily conserved poly(A)-binding proteins (PABPs) have been characterized with some details: PAPBC in the cytoplasm and PABP2/PABPN1 in the nucleus (1,2). Consistent with its cytosolic localization, PAPBC (Pab1 in yeast) stimulates translation initiation by mediating contacts between the mRNA 5'- and 3'-ends via interactions between PAPBC and components of the translational machinery (3,4). PAPBC also appears to act as an antagonist of nonsense-mediated decay (5–7), a pathway of mRNA surveillance that targets transcripts with premature termination codons. Studies in budding yeast and mammals indicate that Pab1 and PAPBC, respectively, shuttle between the nucleus and cytoplasm (8–10) and that Pab1 facilitates the biogenesis and the export of mRNAs (9–11). Consistent with an evolutionarily conserved nuclear function for the cytosolic PABP, intron-containing RNAs can be copurified with mammalian PAPBC (12).

The nuclear counterpart of PAPBC, PABP2, is structurally different from PAPBC and thought to function during polyadenylation of pre-mRNAs. Polyadenylation of most eukaryotic pre-mRNAs consists of a two-step reaction involving endonucleolytic cleavage and poly(A) tail addition. An exhaustive list of evolutionarily conserved proteins responsible for specific and efficient 3'-end processing have been characterized (13–15). These conserved proteins form large multisubunit complexes that bind different cis-acting elements in the 3'-end of pre-mRNAs and determine the site of endonucleolytic cleavage. It has now become clear that 3'-end processing events are tightly integrated to the transcription cycle [reviewed in (13,14,16,17)] through the carboxy-terminal domain (CTD) of RNA polymerase II (Pol II). The Pol II CTD consists of evolutionarily conserved heptad repeats that are thought to act as a platform for the recruitment of various trans-acting factors required for pre-mRNA maturation (18). Notably, this includes many factors involved in 3'-end processing/polyadenylation that are recruited late during the transcription cycle and near the polyadenylation site of yeast (19–21) and human (22,23) genes.

Following cleavage, the nuclear poly(A) polymerase is responsible for the synthesis of poly(A) tails with average length of 70 and 300 nt in yeast and mammals, respectively.

*To whom correspondence should be addressed. Tel: +1 819 820 6868 (12733); Fax: +1 819 564 5340; Email: f.bachand@usherbrooke.ca

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suggest that PABP2 has a dual role in 3'-end formation: (i) PABP2 stimulates processive poly(A) synthesis by direct and simultaneous interactions with the growing poly(A) tail and the poly(A) polymerase (25) and (ii) PABP2 promotes the transition from processive to distributive poly(A) synthesis once a specific length is reached (26). Whereas the genome of the yeast Saccharomyces cerevisiae does not encode for an ortholog of mammalian PABP2, we have recently reported the identification of the PABP2 ortholog in the yeast Schizosaccharomyces pombe (27). Deletion of S. pombe PAB2 results in the expression of RNAs with hyperadenylated tails, indicating that factors other than Pab2 stimulate poly(A) polymerase processivity. Therefore, the precise role of the nuclear poly(A)-binding protein in pre-mRNA polyadenylation in vivo remains unclear.

Translocation of the mRNA ribonucleoprotein (mRNP) complex from the nucleus to the cytoplasm is linked to remodeling events mediated by a myriad of different proteins (28, 29). As yet, the status of the association between PABP2 and nascent mRNPs during and after transit from the nuclear pore complex remains poorly understood. Although mammalian PABP2 shuttles between the nucleus and the cytoplasm (30), earlier results suggest that PAB2 is restricted to nuclear transcripts. Specifically, it has been shown that human PABP2 copurifies with a subunit of the nuclear cap-binding complex, but not with the general translation initiation factor, eIF4E (31). On the basis of these results and the different steady-state distribution of PABPC and PABP2, it has been suggested that poly(A)-bound PABP2 is replaced by PABPC upon transit of the mRNP to the cytosol. The mechanism and cellular compartment of such a substitution between PABP2 and PABPC remain elusive, however.

To further characterize the role of the nuclear poly(A)-binding protein during mRNA synthesis, we performed a comprehensive analysis of Pab2 during mRNP formation in fission yeast. Using chromatin immunoprecipitation (ChiP) assays, our results suggest that Pab2 associates with pre-mRNAs cotranscriptionally prior to 3'-end processing/polyadenylation. Furthermore, tandem affinity purification and mass spectrometry revealed that Pab2 associates with proteins involved in diverse RNA-related functions, including several proteins involved in cytoplasmic translation. Notably, we show that Pab2 is a shuttling protein and present strong evidence that Pab2 associates with translating mRNPs. Our data suggest that Pab2 is recruited early during the transcription cycle and remains associated with translated mRNPs after nuclear export.

**MATERIALS AND METHODS**

**Strains, growth media and genetic methods**

The strains used in this study are listed in Table 1. Schizosaccharomyces pombe was grown at 30°C in yeast extract medium with amino acid supplements (YES) and Edinburgh minimum medium (EMM) containing appropriate amino acid supplements. Schizosaccharomyces pombe cells were transformed with plasmids and PCR products by the lithium acetate method. The strains expressing TAP-tagged version of Pab2 and Pcf11 were constructed by PCR-based gene targeting as described previously (32, 33). The oligonucleotide sequences used for the construction of these strains are available upon request. Appropriate tagging of strain was confirmed by PCR and immunoblotting.

**Plasmids**

Cloning of Pab2 in fusion with the green fluorescent protein (GFP) in a S. pombe-based plasmid was previously described (27). To generate a S. cerevisiae-based plasmid that expresses GFP-Pab2, the DNA encoding the GFP-Pab2 cassette was amplified by PCR using oligonucleotide sequences containing XhoI and EcoRI restriction sites. The resulting PCR product was then cloned into p416ADH (34) that was previously digested with EcoRI and XhoI restriction enzymes to create plasmid pFB197. The ADH1 promoter of pFB197 was replaced by the CYC1 promoter (406 bp) via PCR amplification from S. cerevisiae genomic DNA using oligonucleotide sequences containing SacI and SpeI restriction sites. Following digestion of the PCR product, the DNA was cloned into the SacI and SpeI site of pFB197 to create plasmid pFB235. The open reading frame encoding for S. pombe Pab1 was amplified by PCR from genomic DNA using oligonucleotide sequence containing NotI and BglII restriction sites. Following digestion of the PCR product with NotI and BglII, the DNA was cloned into the NotI and BglII site of pSLF273 (35) to create plasmid pFB243.

**Antibodies**

Mouse monoclonal antibody specific to recognize the carboxy-terminal heptapeptide repeat present on the largest subunit of RNA Pol II (Rpb1; 8WG16) was from Covance Research Products. Rabbit anti-Protein A was from Sigma. Mouse monoclonal antibody specific to HA (clone 12CA5) was from Roche Applied Science. Rabbit polyclonal antibodies specific to fission yeast 40S ribosomal protein S2 (Rps2) were raised at Covance Research Products against GST fusion proteins purified from Escherichia coli.

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ChIP assays

ChIP assays were performed as previously described (36). For RNA Pol II ChIPs, a mouse monoclonal antibody specific for the C-terminal domain of Pol II (8WG16) was used. For the analysis of the NMT1 gene, thiamine was added to the culture medium at a final concentration of 30 μM for 3 h. Determination of RNA-dependent ChIP signals was based on a previously described procedure (37). Briefly, the cross-linking time was reduced to 5 min and the lysis buffer was adjusted to a final SDS concentration of 0.05%. Chromatin preparation were then treated or not treated with a cocktail of RNases to obtain 7.5 U of RNase A and 300 U of RNase T1. Subsequent steps were as previously described (36).

Quantification of the immunoprecipitated DNA was done by quantitative real-time PCR (Rotor-gene 3000, Corbett life sciences) using gene-specific primer sets. Specific dilutions of coimmunoprecipitated DNA as well as of sonicated and reverse cross-linked input DNA were used to determine the percentage of input DNA in each immunoprecipitate. To calculate the increase in signal for the different gene regions, the percentage input values obtained by quantitative PCR were normalized to the percentage input value obtained with the nontranscribed intergenic region, arbitrarily set to 1. The 104-bp PCR amplicon corresponding to the intergenic region lies on the left arm of chromosome I (nucleotides 3009380 to 3009484). Each PCR was run in triplicate, and all ChIPs were repeated at least three times using independent chromatin extracts.

Protein purification

Coimmunoprecipitation assays of RNA Pol II were based on a previously described procedure (38). Briefly, cells were grown in YES at 30°C to mid-log phase and lysed in ice-cold lysis buffer (25 mM NaPO4 at pH 6.8, 0.1 M KOAc, 2 mM MgOAc, 10% glycerol, 1 mM PMSF, 3 ng/ml pepstatin, 3 ng/ml leupeptin, 3 ng/ml aprotonin, 3 ng/ml chymostatin, 0.2 mM Na3VO4, 5 mM β-glycerophosphate, 1 mM NaF) with a Fastprep FP120 using 0.5 mm glass beads. The lysate was transferred to IgG-Sepharose buffer. The lysate was transferred to IgG-Sepharose buffer and incubated with 22 U/ml RNase A for 10 min at room temperature before ultracentrifugation onto 5–45% (w/w) sucrose gradients. KCl treatment of extracts was performed by the addition of salt to a final concentration of 500 mM in polysome lysis buffer and the sucrose gradient. Analysis of formaldehyde cross-linked ribosome profiles was done as previously described (41). Briefly, shaved ice and formaldehyde were added to cultures at final concentrations of 25% and 1%, respectively. After gentle mixing, cells were left on ice for 15 min and 0.1 M glycine was added to stop the cross-linking reaction. Cell lysis and ultracentrifugation were as previously described (39) except that KCl was adjusted to 500 mM in lysis buffer and sucrose gradients. For puromycin treatment experiments, a mixture of 1 mM puromycin/2 mM GTP was added to the culture for 15 min at 30°C before cell lysis. Lysis buffer was also adjusted to 1 mM puromycin/2 mM GTP. Sucrose gradients were fractionated by upward displacement with 55% (w/w) sucrose using a gradient fractionator (Brandel Inc.) connected to a UA-6 UV monitor (Teledyne Isco) for continuous measurement of the absorbance at 254 nm. Twenty fractions of 600 μl were collected and the proteins were TCA-precipitated. Proteins from same amounts of each fraction were separated onto 12% SDS–PAGE, and analyzed by immunoblotting.

Nuclear export assays

Nuclear export assays in S. cerevisiae were done as previously described (40) with some modifications. Briefly, GFP-Pab2 (pFB235), NLS-LacZ-GFP (pFB145) and Nub2-GFP (pFB147) expression constructs were transformed into S. cerevisiae strain FY152 that harbors the nup49-313 temperature-sensitive allele. Cells were grown to OD600 0.1 at 25°C in synthetic medium and separated into two different cultures that were grown for 5 h at the permissive (25°C) or nonpermissive (37°C) temperature. To prevent the synthesis of new GFP-tagged proteins in the cytoplasm, cycloheximide (100 μg/ml) was added to each sample for the last hour to inhibit translation. Cells were examined for GFP fluorescence signal by live fluorescence.

Polysome assay

Polysome profiles were generated from S. pombe as previously described (39). To disrupt ribosomes by chelating Mg2+ ions, EDTA was added to final concentration of 20 mM to the lysis buffer. The lysate was loaded onto 5–45% (w/w) sucrose gradient containing 20 mM EDTA. For RNase disruption of polysomes, the lysate was incubated with 22 U/ml RNase A for 10 min at room temperature before ultracentrifugation onto 5–45% (w/w) sucrose gradients. KCl treatment of extracts was performed by the addition of salt to a final concentration of 500 mM in polysome lysis buffer and the sucrose gradient. Analysis of formaldehyde cross-linked ribosome profiles was done as previously described (41). Briefly, shaved ice and formaldehyde were added to cultures at final concentrations of 25% and 1%, respectively. After gentle mixing, cells were left on ice for 15 min and 0.1 M glycine was added to stop the cross-linking reaction. Cell lysis and ultracentrifugation were as previously described (39) except that KCl was adjusted to 500 mM in lysis buffer and sucrose gradients. For puromycin treatment experiments, a mixture of 1 mM puromycin/2 mM GTP was added to the culture for 15 min at 30°C before cell lysis. Lysis buffer was also adjusted to 1 mM puromycin/2 mM GTP. Sucrose gradients were fractionated by upward displacement with 55% (w/w) sucrose using a gradient fractionator (Brandel Inc.) connected to a UA-6 UV monitor (Teledyne Isco) for continuous measurement of the absorbance at 254 nm. Twenty fractions of 600 μl were collected and the proteins were TCA-precipitated. Proteins from same amounts of each fraction were separated onto 12% SDS–PAGE, and analyzed by immunoblotting.
RESULTS

Cotranscriptional recruitment of Pab2 to active genes precedes 3'-end processing/polyadenylation signals

Electron microscopy has previously detected PABP2 in the vicinity of transcription complexes using salivary glands from the insect Chironomus tentans (42). To investigate further the mechanism by which the nuclear poly(A)-binding protein associates with the transcriptional machinery, we used ChIPs to examine whether S. pombe Pab2 is recruited during the transcription cycle of RNA Pol II. ChIP assays have been used extensively to determine the position of mRNA processing factors along genes and infer the steps at which they are recruited during the transcription cycle. Three Pol II-transcribed genes (ADH1, PYK1 and PGK1) were first examined because of their relatively strong transcription levels (43). To perform ChIP assays, we used a strain in which a Pab2-TAP fusion protein was expressed from the endogenous PAB2 promoter. The Pab2-TAP strain did not confer cold sensitivity in contrast to PAB2-null cells (27) and showed doubling times similar to a wild-type control (data not shown), suggesting that TAP-tagged Pab2 is functional. The amount of genomic DNA associated with a TAP-tagged version of Pab2 was determined by real-time PCR using primer sets located at the 5'-end, the middle and the 3'-end of the ADH1, PYK1 and PGK1 genes (Figure 1A).

ChIP experiments using antibodies specific for the large subunit of RNA Pol II (Rpb1) demonstrated similar cross-linking levels across the ADH1-coding region (Figure 1B). In contrast, ChIP assays using extracts of cells that expressed a TAP-tagged version of the cleavage/polyadenylation factor Pcf11 showed robust cross-linking near the polyadenylation site of the ADH1 gene (Figure 1C). The ChIP patterns for fission yeast Rpb1 and Pcf11 are consistent with similar analyses reported in the budding yeast S. cerevisiae (20,37,44) and demonstrate the validity of our ChIP assays. Pab2 associated mainly with the middle and the 3'-end of the ADH1, PYK1 and PGK1 genes, although some levels of cross-linking were also detected at the 5'-end of these genes (Figure 1D). To confirm and better visualize the gradient of association of Pab2 from 5' to 3', we used the 2.3-kb long MET26 gene (Figure 1A). MET26 was chosen because the larger size of this gene allows for a better resolution of gene segments. Consistent with the data obtained for the ADH1, PYK1 and PGK1 genes, the enrichment of Pab2 at the 5'-end of the MET26 gene (Figure 1D, regions 1 and 2) clearly increased in the middle and at the 3'-end. Further analysis of Pab2 cross-linking using intron-containing genes demonstrated ChIP profiles similar to intronless genes (Figure 2). The results of these ChIP experiments using six different genes indicate a general 5' to 3' enrichment of Pab2.

To determine whether the association of Pab2 to Pol II-specific genes is dependent on active transcription, real-time PCR was performed on Pab2-enriched genomic DNA using primer sets spanning the NMT1-coding region (Figure 3A). Expression of the S. pombe NMT1 gene is strongly repressed following addition of thiamine (45). As can be seen in Figure 3B, Pab2 immunoprecipitates prepared from extracts of cells that were previously grown in the absence of thiamine (active conditions) showed a ChIP pattern on the NMT1 gene consistent with the cross-linking profiles detected for the ADH1, PYK1, PGK1 and MET26 genes. In contrast, the levels of Pab2 cross-linking across the NMT1 gene were similar to the intergenic control region after repression of the NMT1 promoter (with thiamine). These results indicate that recruitment of Pab2 to genes requires active transcription by RNA Pol II.

Pab2 interacts with the nascent mRNP in the vicinity of the transcription complex

The requirement for active transcription to detect Pab2 cross-linking along the coding region of the NMT1 gene suggested that Pab2 might physically interact with components of the transcriptional machinery. We therefore affinity purified TAP-tagged Pab2 from cell extracts using IgG-sepharose and analyzed the precipitated proteins by immunoblotting. Consistent with a previous study in which the communoprecipitation of PABP2 and RNA Pol II from insect cells was demonstrated (42), our experiments also indicated that the large subunit of yeast RNA Pol II (Rpb1) copurifies with TAP-tagged Pab2, but not with a control purification (data not shown). Importantly, we further examined whether the association between Pab2 and the transcriptional machinery is mediated by the nascent mRNP. To test this possibility, extracts were treated with a cocktail of RNases before ChIP analysis. Whereas the ChIP signals for RNA Pol II were not perturbed upon RNase treatment, the level of Pab2 cross-linking over the entire ADH1 coding region was reduced after the samples were treated with RNases (Figure 4A). These results indicate that the nascent transcript is important for the association between Pab2 and genomic DNA in the ChIP assays. Furthermore, the level of Rpb1 that copurified with Pab2-TAP was reduced after RNase treatment (Figure 4B, lanes 2–3), consistent with the importance of RNA for the association between Pab2 and the transcription machinery. Taken together, these immunoprecipitation experiments suggest that recruitment of Pab2 to transcribed genes is mediated by nascent mRNPs.

Pab2 associates with cellular components involved in diverse RNA-related functions

To date, the protein interaction network of the nuclear poly(A)-binding protein remains largely unknown. To get further insights into the mechanism by which Pab2 is cotranscriptionally recruited to nascent mRNPs, a tandem affinity purification approach was used to identify proteins that associate with fission yeast Pab2. Extracts were prepared from 8 l of cells that expressed a TAP-tagged version of Pab2 as well as from untagged control cells. Following two rounds of purification over IgG-sepharose and analyzed the precipitated proteins by SDS–PAGE and visualized by silver staining. As can be seen in Figure 5A, all of the associated proteins were in substoichiometric amounts relative to Pab2. Analysis of the eluted proteins by mass spectrometry
Figure 1. Association of Pab2 to active genes precedes that of the characterized 3'-processing factor, Pcf11. (A) Schematic diagram of genes used for ChIP assays. Boxes represent open reading frames and nucleotides numbers are relative to the initiation codon. Arrows indicate the position of the poly(A) site.
The abundance of the different gene segments from nontranscribed intergenic region was expressed as fold enrichment relative to the nontranscribed intergenic region value, arbitrarily set to 1. The values correspond to the means of at least three independent experiments.

identified unique peptides that corresponded to gene products that are involved in a wide range of RNA-related functions (Figure 5B and Table 2). Notably, peptides from 32 ribosomal proteins, 4 general translation factors, as well as from the cytosolic poly(A)-binding protein (Pab1) were identified (Table 2 and Figure 5B). Peptides that correspond to the evolutionarily conserved nuclear poly(A) polymerase were also detected (Figure 5B), consistent with the role of Pab2 in polyadenylation (27). Other proteins identified that participate in RNA metabolism included the RNA helicase Mtr4, the 5'-to-3' exonuclease Exo2, the mRNA decapping subunit Dcp2 and the nuclear cap-binding protein Cbp80 (Figure 5B). Protein factors for which some peptides were detected in the control purification, no peptides were detected in the control for any of the proteins indicated in Figure 5B and Table 2. In conclusion, our proteomic analysis suggests that Pab2 is associated with factors involved in several steps of the mRNA life cycle.

**Pab2 is a shuttling protein that associates with translating mRNPs**

Fission yeast and human nuclear PABPs are predominantly nuclear at steady state (27,46). Yet, PABP2 was detected in the cytoplasm of human cells by electron microscopy (46) and has been shown to shuttle between the nucleus and cytoplasm (30). To test the capacity of fission yeast Pab2 to shuttle, we used a NUP49-based assay in *S. cerevisiae* (47) as such an *in vivo* export assay does not exist in *S. pombe*. The *nup49-313* allele of *S. cerevisiae* expresses a temperature-sensitive nucleoporin that exhibits defective nuclear import at the nonpermissive temperature without perturbing export of protein and RNA from the nucleus (48). Accordingly, the *nup49-313* allele has been used to assay the export of several nuclear RNA-binding proteins (40,47,49).
Consistent with previous reports indicating that *S. cerevisiae* Nab2 is a shuttling protein (40,50), cytoplasmic accumulation of Nab2 was observed at the non-permissive temperature (Figure 6G and H), whereas Nab2 was predominantly nuclear at the permissive temperature (Figure 6E and F). The expression of GFP-tagged Pab2 in *S. cerevisiae* resulted in nuclear localization (Figure 6I and J), consistent with the steady-state localization of Pab2 in fission yeast (27). In contrast, significant GFP signal was detected in the cytoplasm of Pab2-expressing cells that were shifted to the non-permissive temperature (Figure 6K and L). As a control, a GFP-tagged LacZ protein that includes a strong nuclear localization signal remained in the nucleus at both permissive and non-permissive temperatures (Figure 6A–D). This control also demonstrated that the cycloheximide treatment before imaging prevented *de novo* synthesis of GFP-tagged proteins in the cytoplasm. The results of the nuclear export assay indicate that Pab2 can shuttle between the nucleus and the cytoplasm.

The aforementioned results indicating that Pab2 is a shuttling protein suggested that the detection of ribosomal proteins and translation factors in the eluate of the Pab2-TAP purification could be the result of the association between Pab2 and translating mRNPs. We therefore tested whether Pab2 cosedimented with polysomal mRNAs by examining the fractionation profile of Pab2 after ultracentrifugation using sucrose gradients. As can be seen in Figure 7A, western blot analysis indicated that significant amounts of Pab2 were detected in ribosome-containing fractions (lanes 6–19). We also determined the ratio of ribosome-associated proteins for Pab2, Pab1 and a protein of the small ribosomal subunit, Rps2 (Figure 7A). Roughly 36% of cellular Pab1 cosedimented with ribosomes, consistent with earlier studies (51–53). Notably, our sucrose gradient fractionation experiments indicated that 25% of cellular Pab2 was ribosome-associated (Figure 7A). As expected for a ribosomal protein, most of Rps2 was associated with ribosome-containing fractions (lanes 6–19) and free ribosomal subunits (lanes 3–5). These results indicate that Pab2 cosediments with polysomes.

To validate that the sedimentation behavior of Pab2 reflects an association with polysomes, ribosomes were first disrupted into ribosomal subunits by chelating Mg$^{2+}$ ions. As can be seen in Figure 7B, Pab2 was no longer detected in the heavy fractions of the gradient under these conditions, suggesting that the sedimentation of Pab2 in heavy fractions requires the presence of polysomes. Treating cellular extracts with RNase A prior to ultracentrifugation also perturbed the normal distribution of Pab2 after velocity sedimentation on sucrose gradients (Figure 7C). Specifically, RNase treatment resulted in the disruption of polysomal complexes and the concomitant redistribution of Pab2 to lighter fractions. These results are consistent with the association of Pab2 with translating mRNPs.

To confirm that Pab2 associates with polysomes and not with other large RNP, we used the translation inhibitor, puromycin. Puromycin specifically disrupts polysomes by causing premature release of nascent peptides as well as of mRNAs from ribosomes (54). Accordingly, treatment of cells with puromycin resulted in the almost complete disappearance of polysomes and the concurrent accumulation of 80S monosomes (Figure 7D). Importantly, Pab1, Pab2 and Rps2 were removed from polysomes-containing fractions after puromycin treatment (Figure 7D). These results demonstrate that Pab2 is specifically associated with polysomal mRNPs.

Given that Pab2 shows nuclear localization at steady state [Figure 6; (27)], it was important to determine whether the association of Pab2 with polysomes occurs *in vivo*. To test this, cells were cross-linked with formaldehyde prior to lysis, extracts were prepared using high-salt conditions, and separated on sucrose gradients. As can been seen in Figure 7E, a significant amount of Pab2 cosedimented with polysomes under these experimental conditions. Conversely, Pab2 was redistributed to lighter fractions under similar high-salt conditions, but lacking the formaldehyde cross-linking step (Figure 7F). These experiments indicate that the high-salt conditions dissociated proteins that had not been cross-linked to the mRNPs before cell lysis, including Pab2. Our data thus indicate that a fraction of Pab2 is associated to polysomes *in vivo*.  

**Figure 4.** Pab2 interacts with the nascent mRNP in the vicinity of the transcription complex. (A) Rpb1 and Pab2-TAP ChIPs with and without an RNase treatment step were performed using wild-type cells (for Rpb1) as well as using cells that express TAP-tagged Pab2 using methods similar to those described in Figure 1. The coprecipitating DNA was quantified by real-time PCR using *ADH1*-specific primer pairs as shown in Figure 1. The percentage signal remaining after RNase treatment was calculated as the ratio of the fold enrichment for the RNase-treated sample over the fold enrichment of the sample not treated with RNases. Values correspond to the means of at least three independent experiments. (B) Equal amounts of a whole-cell extract (WCE, lane 1) prepared from Pab2-TAP cells were subjected to affinity purification using IgG-sepharose (lanes 2 and 3). The amount of extract loaded in lane 1 represents 2% of the protein used in the immunoprecipitation. Following extensive washing steps, the beads were treated (lane 3) or not treated (lane 2) with RNases. The eluted proteins were analyzed by western blotting (WB) using antibodies specific to Rpb1 (upper panel) and to the TAP epitope (lower panel).
Although experiments using purified versions of the nuclear PABPs (PABP2) have provided insights into its biochemical properties, the functions as well as the network of associations of PABP2 in a cellular context remain poorly understood. We have recently reported the identification of the ortholog of the mammalian nuclear poly(A)-binding protein in *S. pombe* (27). Consistent with a functional role in poly(A) tail synthesis, RNAs from *PAB2*-null cells display hyperadenylated 3’-ends (27). In this study, we investigated the recruitment of Pab2 during mRNP formation and the subsequent association between Pab2 and exported mRNPs. Based on our results, we propose that Pab2 is recruited early during the transcription cycle of RNA Pol II genes and

**DISCUSSION**

Figure 5. Purification of proteins associated with TAP-tagged Pab2. (A) Proteins copurified with Pab2 by tandem affinity purification (lane 1) were resolved using a Bis–Tris 4–12% gradient SDS–PAGE and analyzed by silver staining. The result for an identically treated extract from control *S. pombe* is shown in lane 2. Molecular weight markers are shown on the right in kilodaltons (kDa). The position of Pab2 is indicated on the left. (B) Summary of the nonribosomal proteins identified by mass spectrometry from the Pab2-TAP purification.
remains associated with translating mRNPs after nuclear export. These findings provide novel insights into the events that govern mRNP remodeling as it transits to the cytoplasm.

During eukaryotic transcription, nascent pre-mRNAs are wrapped in ribonucleoprotein complexes that contain factors required for processing and export (55). It is now established that most factors required for processing and export of pre-mRNAs are loaded onto the nascent mRNP as it comes out of the polymerase RNA exit channel. The cotranscriptional loading of RNA processing factors includes several cleavage/polyadenylation factors that show recruitment near the 3'-end of genes as determined by ChIP assays (19–23). The ChIP assays presented in this study indicated that Pab2 is cotranscriptionally recruited via the nascent mRNP. The physical fragmentation of RNA during the sonication step is likely to be the cause for the plateau in ChIP signal detected for Pab2 beyond the middle of the ADH1, PYK1, PGK1 and MET26 genes, and reflects the size of chromatin fragments to <500 bp. Interestingly, our ChIP assays indicated that Pab2 occupancies upstream of the polyadenylation site are sensitive to RNases (Figure 4). These results could imply that the recruitment of Pab2 is mediated by direct binding to non-poly(A) sequences in the pre-mRNA prior to polyadenylation. We do not favor this interpretation, however, as studies using nuclear PABPs from various organisms indicate poor binding to nonpolyadenylated RNA as compared with poly(A) (27,56,57). Analogous to our ChIP experiments, sensitivity of ChIP signals to RNases was previously reported for the budding yeast proteins Sub2 and Gbp2 (37,58). In this case, results suggest that the TREX complex mediates the cotranscriptional recruitment of Sub2 and Gbp2 to nascent mRNPs. Similarly, we predict a model in which Pab2 is recruited to the nascent mRNP via protein interactions. A structure–function analysis of S. pombe Pab2 should provide useful insights into the mechanism that mediates the cotranscriptional recruitment of Pab2 to nascent mRNPs.

The Pab2 ChIP results described herein are in agreement with earlier electron microscopy studies that have found PAB2 in the vicinity of RNA Pol II along the entire Balbiani ring gene of Chironomus tentans (42). This previous study was limited to a single gene, however, and did not address how PAPB2 is recruited during the transcription cycle of RNA Pol II. Our study goes beyond these previous findings and provides evidence for the transcription-dependent recruitment of Pab2 to the NMT1 gene and to the coding sequences of several other yeast genes that are transcribed constitutively. Furthermore, the inclusion of an RNase step in our experiments supports the recruitment and/or the rapid transfer of Pab2 to the nascent mRNP. It remains unclear, however, whether RNA-independent interactions exist between Pab2 and the transcription machinery. Whereas the level of ADH1 mRNA was reduced after RNase-treatment as compared with a nontreated sample, segments of the ADH1 transcript could still be detected by RT-PCR following treatment with RNases (data not shown). It is therefore likely that the remaining signal detected by our immunoprecipitation assays (Figure 4) is due to the incomplete digestion of RNA.

Similar to the ChIP profile of Pab2, other cleavage/polyadenylation factors have also been shown to

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cross-link at promoters and coding regions in yeast and other eukaryotes (19,22,59,60). Although the functional significance for the recruitment of cleavage/polyadenylation factors during the early phase of the transcription cycle remains unclear, the multi-functional nature of some mRNA processing factors might explain this apparent discrepancy. Interestingly, cellular depletion of Drosophila PABP2 using RNAi results in nuclear accumulation of polyadenylated RNAs (61), suggesting a role for the nuclear poly(A)-binding protein in mRNA export pathways. Studies also indicate that a number of proteins important for efficient mRNA export are recruited during the transcription cycle in budding yeast (37,44,62,63) and humans (64,65). Remarkably, the ChIP profile of many mRNA export factors shows enrichment from the 5'-end to the 3'-end, similar to that of Pab2. Whereas these aforementioned evidence may support a role for the nuclear poly(A)-binding protein in mRNA export, it cannot be excluded that the accumulation of poly(A) RNA in PABP2-depleted Drosophila cells is due to defective polyadenylation. Accordingly, we show here that a significant fraction of Pab2 does not get replaced by Pab1 and remains associated with translating mRNPs. Our studies thus provide the first evidence suggesting the association of the nuclear poly(A)-binding protein with translating mRNPs. The presence of peptides corresponding to Pab1 in the eluate of the Pab2-TAP purification (Figure 5) also suggests that both PABPs can coexist on the same poly(A) tail, in agreement with other studies (67,68).

The association of Pab2 with translating mRNPs is in contrast to previous results that suggest that mammalian PABP2 is not associated with general translation. More precisely, biochemical experiments in human cells have suggested that human PABP2 is restricted to a pioneer round of translation (31). In mammalian cells, it has been proposed that PABPC substitutes for PABP2 after nuclear export of mRNPs. How poly(A)-bound PABP2 is replaced by PABPC during or after nuclear export has remained elusive, however. Importantly, we show here that a significant fraction of Pab2 does not get replaced by Pab1 and remains associated with translating mRNPs. This conclusion is supported by the specific cosedimentation of Pab2 with polyribosomes (Figure 7) as well as the copurification of ribosomal proteins and translation factors with Pab2 (Figure 5 and Table 2). Our studies thus provide the first evidence suggesting the association of the nuclear poly(A)-binding protein with translating mRNPs. The presence of peptides corresponding to Pab1 in the eluate of the Pab2-TAP purification (Figure 5) also suggests that both PABPs can coexist on the same poly(A) tail, in agreement with other studies (67,68).

The association of Pab2 with translating mRNPs is in contrast to previous results that suggest that mammalian PABP2 is not associated with general translation. More precisely, biochemical experiments in human cells have suggested that human PABP2 is restricted to a pioneer round of translation (31). In mammalian cells, it has been proposed that aberrant mRNAs containing nonsense codons are recognized during a pioneer round of translation that is defined by mRNAs bound by the nuclear cap-binding complex proteins, Cbp20 and Cbp80 (31,68). The detection of Pab2 in large polysomal fractions reported here (Figure 7) suggests that Pab2 is not restricted to pioneer rounds of translation, but also associates with actively translated mRNAs. It is therefore possible that the nuclear poly(A)-binding protein performs a slightly different role in mRNA translation between yeast

Figure 6. Pab2 shuttles between the nucleus and cytoplasm. The nuclear export assay was performed using S. cerevisiae cells that express the nup49-313 allele as described in the Materials and methods section. The nup49-313 cells expressing either NLS-LacZ-GFP (A–D), Nab2-GFP (E–H) and GFP-Pab2 (I–L) were incubated at 25°C (A, B, E, F, I and J) or shifted to 37°C (C, D, G, H, K and L) before treatment with cycloheximide to block new protein synthesis. Phase contrast (A, C, E, G, I and K) and GFP fluorescence (B, D, F, H, J and L) are shown.
and humans or that technical issues hampered the detection between human PABP2 and translating mRNPs. Our results do not exclude a role for Pab2 in a pioneer round of translation, however. The proteomic analysis of Pab2-associated proteins identified the fission yeast homolog of mammalian Cbp80 (Figure 5), a key constituent of the pioneer round of translation (31,68). Furthermore, two key factors involved in 5′-to-3′ decay of NMD targets, Dcp2 and Exo2, also copurified with Pab2.

The functional significance of the association between Pab2 and translating mRNPs still remains to be determined. Yet, the hypersensitivity of PAB2-null cells to different translational inhibitors (our unpublished data) is consistent with a possible role for Pab2 in translation. Accordingly, a number of predominantly nuclear proteins have recently been shown to cosediment with polysomes and modulate translation in yeast and mammals (53,69–73). Conversely, studies indicate that budding yeast Pab1 (homolog of mammalian PABPC) is a shuttling protein that is important for the proper assembly of mRNPs in the nucleus (9,10). Our findings that Pab2 is associated with translating mRNPs are in concert with the conclusions of the aforementioned studies and illustrate the integrated connections between different steps of translation.
mRNP formation in the nucleus and the fate of the mature mRNA in the cytoplasm. Future studies on Pab2 will certainly provide insight into the mechanisms that coordinate nuclear polyadenylation and translation in the cytoplasm.

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