Xlrbpa, a Double-stranded RNA-binding Protein Associated with Ribosomes and Heterogeneous Nuclear RNPs

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Abstract. We have cloned and characterized Xlrbpa, a double-stranded RNA-binding protein from *Xenopus laevis*. Xlrbpa is a protein of 33 kD and contains three tandemly arranged, double-stranded RNA-binding domains (dsRBDs) that bind exclusively to double-stranded RNA in vitro, but fail to bind either single-stranded RNA or DNA. Sequence data and the overall organization of the protein suggest that Xlrbpa is the *Xenopus* homologue of human TAR-RNA binding protein (TRBP), a protein isolated by its ability to bind to human immunodeficiency virus (HIV) TAR-RNA. In transfection assays, TRBP has also been shown to inhibit the interferon-induced protein kinase PKR possibly by direct physical interaction. To determine the function of Xlrbpa and its human homologue we studied the expression and intracellular distribution of the two proteins. Xlrbpa is ubiquitously expressed with marked quantitative differences amongst all tissues. Xlrbpa and human TRBP can be detected in the cytoplasm and nucleus by immunofluorescence staining and Western blotting. Sedimentation gradient analyses and immunoprecipitation experiments suggest an association of cytoplasmic Xlrbpa with ribosomes. In contrast, a control construct containing two dsRBDs fails to associate with ribosomes in microinjected *Xenopus* oocytes. Nuclear staining of *Xenopus* lampbrush chromosome preparations showed the association of the protein with nucleoli, again indicating an association of the protein with ribosomal RNAs. Additionally, Xlrbpa could be located on lampbrush chromosomes and in snurposomes. Immunoprecipitations of nuclear extracts demonstrated the presence of the protein in heterogeneous nuclear (hn) RNP particles, but not in small nuclear RNPs, explaining the chromosomal localization of the protein. It thus appears that Xlrbpa is a general double-stranded RNA-binding protein which is associated with the majority of cellular RNAs, ribosomal RNAs, and hnRNAs either alone or as part of an hnRNP complex.

Most cellular RNAs associate with a variety of proteins that are required for several important cellular functions. Heterogeneous nuclear (hn) RNP proteins, for instance, package newly transcribed hnRNA into hnRNP particles (Piñol-Roma et al., 1988). Splicing factors, in combination with splicing small nuclear (sn) RNPs, on the other hand, mediate the removal of introns from pre-mRNAs (reviewed by Nilsen, 1994). Other modifying proteins are required for capping, polyadenylation, or editing of mRNA (Adam et al., 1986; Christofori and Keller, 1989; Hamm and Mattaj, 1990; Wahle, 1991; Bass, 1993; Polson and Bass, 1994).

Structural RNAs such as ribosomal RNAs or snRNAs also have to be processed, modified, and complexed with proteins to become functionally active (Reddy and Bush, 1988; Tyc and Steitz, 1989; Savino and Gerbi, 1990; Wool et al., 1990; Peculis and Steitz, 1993; Mougey et al., 1993). RNA-binding proteins can even be involved in the control of gene expression, such as the iron regulatory factor that controls the stability of human transferrin receptor mRNA and the translatability of ferritin mRNA by binding to these RNAs (Leibold and Munro, 1988; Müllner et al., 1989). Other RNA-binding proteins are required for the proper localization of certain RNAs thereby controlling cell polarity or body axis and pattern formation during development (for review see St Johnston, 1995).

Some RNA-interacting proteins bind RNA directly while others contact RNA only as part of a larger complex (Piñol-Roma et al., 1988). RNA–protein interaction is frequently mediated through specific RNA-binding domains (RBDs). These domains can be quite similar to each other in very different proteins, both in their primary amino acid sequence and at the structural level, and can thus be defined by specific consensus sequences. So far, several con-
served RBDs could be defined by their homologies to each other. Among these, the 90–100-amino acid RNA recognition motif is the most abundant and probably best-characterized RBD (Bandziulis et al., 1989; Nagai et al., 1990; Wittenkind et al., 1992). Other well-characterized RBDs include the zinc-finger motif, the arginine-rich motif, the RGG box, and the KH box (for review see Burd and Dreyfuss, 1994).

A new RNA-binding motif that binds exclusively to double-stranded RNA or RNA–DNA hybrids is the so-called double-stranded RNA-binding domain (dsRBD) (St Johnston et al., 1992; Green and Mathews, 1992; Bass et al., 1994). This protein domain is ~70 amino acids in length with several basic amino acids clustered at its carboxy terminus. While most dsRBDs fit a consensus sequence quite well over their entire length, some dsRBDs appear less conserved at their amino terminus than at their basic carboxy-terminal end (St Johnston et al., 1992; Krovat and Jantsch, 1996). Like other RBDs, the dsRBD can sometimes be found in multiple copies within a single protein. So far, dsRBDs could be identified in almost 30 putative or known RNA-binding proteins from *Escherichia coli* to man (Kharrat et al., 1995; Krovat and Jantsch, 1996). Recently the in-solution structure of two different dsRBDs have been determined by nuclear magnetic resonance (NMR) analysis (Bycroft et al., 1995; Kharrat et al., 1995). Both studies determined virtually identical structures for both dsRBDs, suggesting a conserved structural organization of all dsRBDs. While some dsRBD containing proteins associate only with specific RNAs in vivo, no sequence-specific binding of dsRBDs could so far be demonstrated in vitro.

From a screen for RNA-binding proteins we have isolated at least four different dsRBD-containing proteins from a *Xenopus* ovari-specific cDNA library. One of these proteins, named *Xenopus laevis* RNA-binding protein A (Xlrba), shows significant homology to human trans-activation–responsive (TAR)–RNA binding protein (TRBP) (Gatignol et al., 1991) and seems to be its *Xenopus* homologue. Human TRBP has been isolated by its ability to bind human immunodeficiency virus 1 (HIV-1) TAR-RNA, and has been postulated to be involved in the transcriptional activation of several viral promoters (Gatignol et al., 1991). TRBP has also been shown to act as an inhibitor of the cellular interferon-induced kinase PKR when overexpressed in COS-1 cells (Park et al., 1994). PKR is activated by double-stranded RNAs resulting in autophosphorylation and consecutive phosphorylation of translation initiation factor eIF2a, which leads to inhibition of protein synthesis. Inhibition of PKR by TRBP has thus been attributed to competitive binding of double-stranded RNAs by TRBP, therefore preventing their binding to PKR and subsequent activation of kinase activity (Park et al., 1994). A possible function of TRBP could therefore be the complexation of structured cellular RNAs to prevent inadvertent activation of PKR. TRBP has also been shown to dimerize with PKR through RNA bridges (Cosentino et al., 1995). In contrast, a recent study demonstrates physical interaction of the two proteins in an RNA-independent manner, suggesting a regulation of PKR by TRBP in a more direct way (Benkirane et al., 1997).

However, association of TRBP with cellular RNAs has not been investigated to this point. We therefore studied the expression and cellular distribution of Xlrba and its human homologue, TRBP, both cytologically and at the biochemical level. We also determined the association of these proteins with RNA. Both proteins seem ubiquitously expressed and are localized in the cytoplasm and nucleus. In the cytoplasm, most of these proteins seem associated with ribosomes. Additionally, lambrush chromosome preparations of *Xenopus* germinal vesicles revealed the association of Xlrba with nascent hnRNAs on transcribing chromosome loops. Both observations, the association with ribosomes and hnRNAs, were confirmed by biochemical experiments.

**Materials and Methods**

**Isolation of Xlrba cDNA**

Xlrba cDNA was isolated from a *Xenopus laevis* ovari cDNA library by Northwestern screening with 32P-labeled U1 or U2 snRNA. The screen was performed according to the procedure of Vinson et al. (1988), except that 8 M urea replaced 6 M guanidine hydrochloride as the chaotropic agent. The binding buffer contained 50 mM NaCl, 10 mM MgCl2, 10 mM HEPES, pH 7.5, 0.1 mM EDTA, and 1 mM DTT. The probe was a mixture of 32P-labeled transcripts produced by T7 polymerase from cloned U1 and U2 snRNA genes. The Xlrba insert was sequenced after conversion of the λZAP bacteriophage into the corresponding pBluescript phagemid (Stratagene, LaJolla, CA). These sequence data are available from EMBL/GenBank/DDBJ under accession No. M96570.

**Antibody Production**

A BamHI fragment encoding the carboxy-terminal two thirds of the Xlrba cDNA was cloned into the GST fusion vector pGEX (Pharmacia Diagnostics AB, Uppsala, Sweden). Plasmids containing the proper insert were transformed into *Escherichia coli* BL21. 3 ml starting cultures were grown overnight and used to inoculate larger 300-ml cultures the next morning. When the cultures reached an OD600 of 0.7, protein production was induced by the addition of isopropyl β-D-thiogalactoside (IPTG) to 1 mM final concentration. After an additional 4 h of incubation, cells were harvested by centrifugation and lysed by sonication in PBS, 1% Triton X-100. Cell lysate was cleared by centrifugation and the supernatant was loaded on a glutathione Sepharose 4B column (Pharmacia Diagnostics AB). The column was washed extensively with PBS, 1% Triton X-100 to remove unbound protein. Bound glutathione-S-transferase (GST) fusion protein was eluted by washing the column with 5 mM glutathione in 50 mM Tris-HCl, pH 8.0. Typical yields were 3–5 mg of pure protein per 500 ml culture.

**Peptide Coupling**

Peptide MJ2 (CDYVKMLKDVAEELDF) was coupled to keyhole limpet hemocyanin (KLH) by ω-maleimidobutyric acid N-hydroxysuccinimide ester cross-linking according to Harlow and Lane (1988).

**Antibodies**

Two rabbits (Rb3 and Rb5) were immunized with KLH-coupled peptide MJ2 while two rabbits (Rb6 and Rb7) were immunized with GST fusion protein by subcutaneous injections after the immunization schemes given by Harlow and Lane (1988). 2 wk after the first booster injection, test sera were collected from the ear vein and tested for their ability to detect the GST fusion protein in Western blots. Since all animals had developed antibodies against Xlrba, two further booster injections were administered before animals were sacrificed, at which stage all blood was collected. Sera were prepared according to Harlow and Lane (1988) and stored frozen at −70°C.

mAb 4F4 recognizes human hnRNPs C1 and C2 (Choi and Dreyfuss, 1984). Human autoimmune sera 13751 and 92751 recognize ribosomal phosphoproteins P0, P1, and P2 as determined by comparison with reference serum SD1 obtained from CDC (Atlanta, GA) (Steiner, G., personal communication). mAb Y12 recognizes the Sm epitope of snRNPs (Lerner et al., 1981).
Western Blotting

Sample preparation and SDS gel electrophoresis was performed according to standard procedures. After electrophoresis, proteins were transferred to Immobilon-PVDF (polyvinylidene difluoride) membranes (Millipore Corp., Waters Chromatography, Bedford, MA). For Western blotting, peptide antiserum were used at 1:300 dilutions while antiserum against GST fusion proteins were used at a dilution of 1:500. Bound antibodies were detected with 125I-labeled protein A and autoradiographed.

Oocyte Sections

Small pieces of *Xenopus* ovary containing oocytes of different developmental stages were quick-frozen by immersion in isopentane at −180°C, quickly transferred to ethanol at −70°C, and stored for 2–3 d at that temperature. Dehydrated tissue was subsequently warmed to room temperature (3 h at −20°C, 3 h at 4°C, 3 h at 20°C), and finally embedded in histocote at 65°C after a passage through paraformaldehyde.

4-μm sections were cut with a glass knife on a Reichert ultramicrotome and mounted on coated slides. Sections were deparaffinized in xylene, rehydrated in a descending alcohol series, and used for immunostaining after a final wash in PBS. Primary antibodies were detected with secondary, alkaline phosphatase–labeled antibodies (Harlow and Lane, 1988).

Immunoprecipitation

For precipitation of hnRNP particles, HeLa nucleoplasms was prepared according to Choi and Dreyfuss (1984). For precipitation of ribosomal proteins, small pieces of ovary were homogenized by sonication in ribosomal buffer D (0.1 M KC1, 5 mM MgCl2, 1 mM β-mercaptoethanol, 0.1 M EDTA, and 50 mM Tris-HCl, pH 7.8) containing 0.1% deoxycholic acid, 1 mM vanadyl–ribonucleoside complex, 100 μg/ml PMSF, 2 μg aprotinin, and 2 μg/ml pepstatin (Brown et al., 1974). The lysate was cleared from insoluble material by two rounds of centrifugation at 4°C in a table top centrifuge. Avoiding the fatty surface layer, the supernatant was layered over a linear 10% to 30% (wt/vol) sucrose gradient prepared in ribosomal buffer E (50 mM KC1, 2 mM MgCl2, 1 mM β-mercaptoethanol, 50 mM Tris-HCl, pH 7.8) and centrifuged for 4.5 h at 26,000 rpm in a Beckman SW28 rotor (Beckman Instruments, Inc., Fullerton, CA) at 4°C (Brown et al., 1974). Gradients were fractionated in a cold room. 1-ml fractions were taken from the bottom of the gradient and absorbance at 254 nm was monitored with an UV-M-II optical unit (Pharmacia Diagnostics AB).

Immunofluorescence Staining

Cells grown on acid-etched coverslips were fixed and permeablized according to Fu and Maniatis (1990), with the exception that methanol replaced acetone in the permeablization step. After fixation, cells were dehydrated through an ethanol series, permeablized with methanol at −20°C for 10 min, washed in 70% ethanol, and transferred to PBS. Lampbrush chromosome preparations were performed as described (Gall et al., 1991). Immunofluorescence staining was performed as described by Wu et al. (1991).

Construction of Control Construct MP-2

Control construct MP-2 was constructed by PCR amplification of the two dsRBDs of a clone related to 4f1 (Bass et al., 1994). Primers were designed in a way to allow in-frame cloning of the amplified fragment upstream of six myc tags, followed by the polyadenylation (polyA) tail of NO38 cDNA. The resulting clone encodes a protein with a calculated molecular weight of 32 kD consisting of two dsRBDs followed by six COOH-terminal myc tags.

Protein Expression and Northwestern RNA-binding Assay

To determine the RNA-binding ability of construct MP-2, the coding region of this construct was cloned in-frame into the protein expression vector pET7b. For a comparison of RNA-binding of MP-2 and wild-type Xlrbpa, the entire coding region of Xlrbpa was cloned into the protein expression vector pET 3b, resulting in construct pET3.5. For protein expression, both clones were transformed into *E. coli* BL21(DE3). Small cultures were grown to mid log phase at which stage protein expression was induced by the addition of 1 mM isopropyl β-D-thiogalactoside (IPTG). After 2 h of induction, cells were harvested by centrifugation and cell lysates were prepared by sonication in SDS sample buffer. RNA-binding assays were performed as described, using radiolabeled poly rI/rC as a double-stranded RNA substrate (St Johnston et al., 1992).

Multimerization Assay

To determine a possible multimerization of Xlrbpa, the full-length protein was expressed from construct pET3.5 (see above) in *E. coli* BL21(DE3). Overexpressed protein was purified by two rounds of cation-exchange chromatography resulting in at least 90% pure protein as judged by SDS gel electrophoresis. Purified Xlrbpa was RNA-binding as determined by Northwestern assay.

The multimerization potential of Xlrbpa was tested by chromatography of purified protein on a Sepharose CL-4B sizing column (Pharmacia Diagnostics AB) next to suitable sizing standards. On these columns Xlrbpa eluted right at its calculated molecular weight of 32 kD, giving no indication for a potential multimerization of the protein. As a second approach to determine potential multimerization of Xlrbpa in solution, glutaraldehyde cross-linking of the protein was performed (Thompson et al., 1991). Transcription factor GCN-4 and carbonic anhydrase were used as positive and negative controls in the same assay. Cross-linking products were analyzed by SDS gel electrophoresis and silver staining of gels. No cross-linking of Xlrbpa could be detected in this assay. Finally, purified Xlrbpa was subjected to sedimentation gradient analysis. Again, there was no indication for a possible multimerization of Xlrbpa, as the protein always migrated at the top of the gradient.

Tissue Culture Transfection Assays

To determine the cellular localization of construct MP-2, the coding region, including the six myc tags and the polyA tail of MP2 were cloned into the eukaryotic expression vector pcDNA (Invitrogen, San Diego, CA). The resulting construct was used for transient transfection of coverslip-grown HeLa cells. Immunofluorescence staining of transfected cells was performed as described (Jantsch and Gall, 1992).
Oocyte Injection

Construct MP-2 was linearized at a unique BamHI site downstream of the NO38 polyA tail. Capped in vitro–transcribed RNA was produced using T3 RNA polymerase as described (Jantsch and Gall, 1992). Oocytes were injected with ~50 ng RNA per oocyte and incubated overnight at 16°C to allow protein synthesis. For Western blotting, ~10 germinal vesicles (GV) or three cytoplasts were used, per lane. For sedimentation gradient analysis, the homogenate of ~100 microinjected oocytes were used for each gradient. The MP-2 myc fusion protein was detected by Western blotting using mAb 9E10 directed against the myc tag, and a secondary anti–mouse antibody labeled with alkaline phosphatase for enzymatic detection (Evan et al., 1985).

Results

Isolation of Xlrbpa cDNA

By Northwestern screening of a Xenopus laevis oocyte cDNA library with radiolabeled U1 and U2 snRNAs we have previously isolated several clones that encode proteins which bind to these RNAs. Sequence analysis and deletion mapping of the RNA-binding region within these clones led to the identification of the dsRBD in several cDNAs (St Johnston et al., 1992).

One group of the isolated cDNAs encodes a protein showing strong homology to human TRBP (Gatignol et al., 1991), both at the primary amino acid level and in its overall organization (Fig. 1). We called the protein encoded by these cDNAs Xenopus laevis RNA-binding protein A (Xlrbpa).

At the amino acid level Xlrbpa shows >70% homology and >55% identity to human TRBP. The two proteins are of similar size with calculated molecular weights of 32.8 and 36.9 kD for Xlrbpa and human TRBP, respectively. Additionally, both proteins are very similar in their general organization. Both contain three tandemly arranged dsRBDs, each separated by short spacers of variable length (Fig. 1). In both proteins, the third dsRBD shares a lower degree of homology with the overall dsRBD consensus sequence while the first and second dsRBDs are highly homologous to it (St Johnston et al., 1992; Krovat and Jantsch, 1996).

Taken together, the data suggest that Xlrbpa is the Xenopus homologue of human TRBP. This view is supported by two additional facts. First, in pairwise alignments of 20 dsRBDs from different proteins, the corresponding dsRBDs in Xlrbpa and human TRBP always turn out to be more closely related to each other than to any other dsRBDs in the entire alignment (St Johnston et al., 1992; Krovat and Jantsch, 1996). Second, antibodies directed against Xlrbpa recognize in vitro–translated human TRBP and a protein of identical molecular weight to TRBP in HeLa cells (see below).

Expression of Xlrbpa

To study the distribution of Xlrbpa both at the level of the entire organism and at the cellular level, we produced antibodies against an Xlrbpa fusion protein and a peptide located in the third dsRBD of Xlrbpa. All antisera recognized Xlrbpa expressed from E. coli and the endogenous protein in Xenopus oocytes; however, the antisera directed against the fusion protein (Rb6 and Rb7) gave stronger signals than peptide sera and were thus used for most parts of this study.

On Western blots, all antisera recognized a single band of 33 kD both in oocytes and XIA6, a Xenopus cell line (Fig. 2 a), while preimmune sera showed no signals on these blots (data not shown). The observed band of 33 kD corresponds well with a calculated molecular weight of 32.8 kD for Xlrbpa. In addition, we tested whether any of our antisera would recognize human TRBP, the putative homologue of Xlrbpa. All antisera directed against Xlrbpa fusion protein (Rb6 and Rb7) were able to specifically immunoprecipitate 35S-labeled, in vitro–translated human TRBP while the corresponding preimmune sera failed to do so (data not shown). Additionally, antisera were tested on Western blots for their ability to recognize human TRBP in HeLa cell extracts. Of all sera tested, Rb6 directed against the Xlrbpa fusion protein gave the strongest signal on these blots. The band observed in HeLa extracts migrates a little slower than the corresponding protein in Xenopus oocyte extracts (Fig. 2 b), which is in good agreement with the slightly larger molecular weight of 36 kD of human TRBP. The corresponding preimmune serum showed no signal in HeLa cells (data not shown). Also, the antiserum could be blocked by preincubation with E. coli–produced Xlrbpa, suggesting that serum Rb6 recognizes authentic human TRBP in HeLa cells (data not shown).

To determine the expression of Xlrbpa in adult Xenopus we performed Western blots of various tissues. Xlrbpa could be detected in all tissues investigated, including oocyte, spleen, liver, kidney, heart, nerve, brain, and XIA6, indicating that Xlrbpa is a ubiquitously expressed protein.


Intracellular localization of Xlrbpa

Human TRBP has been suggested to be involved in the transcriptional activation of HIV TAR-RNA and also has been shown to increase the expression of reporter genes when expressed under the control of other viral promoters (Gatignol et al., 1991). However, an RNA-independent interaction of TRBP with the double-stranded RNA-activated kinase PKR has also been reported recently (Benkirane et al., 1997). Direct transcriptional activation could only take place in the nucleus while interaction of TRBP with the cytoplasmic kinase PKR would most likely occur in the cytoplasm. We therefore wanted to determine the intracellular localization of Xlrbpa and its human homologue TRBP. To do this, immunostainings of oocyte sections and coverslip-grown HeLa tissue culture cells were performed. Additionally, we made Western blots of hand-enucleated oocytes and isolated germinal vesicles.

Surprisingly, most of the protein could be detected in the cytoplasm of stained oocyte sections (Fig. 3). Upon close investigation, however, minor staining could also be detected in the nucleus where staining appeared in small dots, possibly representing nucleoli or chromatin. The same result was obtained on Western blots of hand-isolated GVs and enucleated oocytes (Fig. 2 d).

For comparison, we stained coverslip-grown HeLa tissue culture cells with the anti-Xlrbpa antiserum Rb6. As mentioned, this antiserum recognizes human TRBP (Fig. 2 b). Similar to the situation in oocytes, human TRBP in HeLa cells seemed to be located in both the cytoplasm and the nucleus, resulting in an almost homogeneous staining pattern (Fig. 4). However, when compared to oocytes, the concentration of TRBP in HeLa cells seemed relatively high in the nucleus. In the cytoplasm, the staining appeared in minute dots possibly colocalized with the ER. Colocalization of TRBP with the ER could be indicative of an association of the protein with ribosomes. Therefore, to compare the cytoplasmic distribution of TRBP with that of ribosomes in HeLa cells we performed double immunofluorescence staining with Rb6 and serum 13751, an anti-ribosomal autoimmune serum that recognizes ribosomal proteins P0, P1, and P2 (Steiner, G., unpublished results), which are the putative eukaryotic homologues of bacterial ribosomal proteins L10, L7, and L12, respectively (Santos and Ballesta, 1994). In these experiments both sera showed similar staining patterns, consistent with an association of TRBP with the ER (Fig. 4 g).

Xlrbpa in Ribosomes

To determine whether the cytoplasmic localization of Xlrbpa and its human homologue TRBP could be due to a physical association of these proteins with ribosomes, we performed sedimentation gradient analyses of Xenopus oocyte extracts. In sucrose gradients, the majority of Xlrbpa cosedimented with the peak of 80S ribosomes. Only minor amounts could be found in the pellet and supernatant corresponding to polysomes and hnRNAs, as noted, however, that it was almost impossible to obtain protein extracts completely free of degradation products, indicating that Xlrbpa is very sensitive to proteolytic degradation.
well as free protein, respectively (Fig. 5 A). To confirm these results and to exclude the possibility that Xlrbp was only trapped in the ribosomal fraction, the peaks of 80S ribosomes from several gradients were concentrated by microfiltration and rerun on a second sucrose gradient. Again, the majority of Xlrbp was cosedimenting with 80S ribosomes, confirming the association of Xlrbp with ribosomes (Fig. 5 A, right). A ribosomal association of Xlrbp is in agreement with our recent study which demonstrated binding of Xlrbp to rRNA in vitro (Krovat and Jantsch, 1996). The observed association of Xlrbp with 80S ribosomes was unchanged by puromycin treatment of oocyte extracts, but was sensitive to RNase A treatment (data not shown). It should be noted, however, that the association of Xlrbp with ribosomes was sensitive to high salt washes. When ribosomal pellets were exposed to salt concentrations higher than 300 mM, a large fraction of Xlrbp dissociated from the ribosomes resulting in reduced amounts of Xlrbp cosedimenting with ribosomes and larger amounts of Xlrbp migrating as free protein at the top of the gradient (data not shown).

Xlrbp has a molecular weight of 33 kD. The free protein should therefore be able to pass through a microfiltration membrane with a molecular weight cutoff of 50,000. However, if Xlrbp was part of a larger particle, such as a ribosome, it should remain in the top chamber of the microconcentration device. We therefore tested the material from the top and bottom chamber of the Centricon device for the presence of Xlrbp. As can be seen in Fig. 5 A, all Xlrbp remained in the top chamber during microconcentration, indicating the association of Xlrbp with a physical particle >50 kD.

Purified full-length Xlrbp does not form protein multimers as judged by sizing columns, sedimentation gradients, and glutaraldehyde cross-linking experiments (data not shown). The possibility that Xlrbp does not pass through the microfiltration column due to the formation of protein multimers could therefore be excluded. However, RNA-dependent multimerization of TRBP and the dsRBD-containing fragment of PKR kinase has been observed in the yeast two-hybrid system and in vitro (Cosentino et al., 1995). Even direct interaction of PKR and TRBP has been reported recently (Benkirane et al., 1997). An association of Xlrbp with ribosomes could therefore be mediated by an interaction of the protein with the ribosome-associated kinase PKR.

Nonetheless, to exclude the possibility that Xlrbp would only stick to ribosomes in a nonspecific manner due to the protein’s intrinsic RNA-binding ability or, alternatively, would form multimers through a common RNA linker, we determined whether an unrelated control construct containing two dsRBDs would also associate or cosediment with ribosomes when expressed in Xenopus oocytes. The Xenopus 4f1 protein seemed suitable for this purpose. Xenopus 4f1 contains two dsRBDs, and is also very abundant in Xenopus oocytes (Fig. 6 A) (Bass et al., 1994). The 4f1 protein is homologous to the human NF90 transcription factor (Kao et al., 1994). A clone encoding an isoform of this protein was previously isolated by us (Jantsch, M., unpublished results). Since full-length 4f1 protein contains an NLS and is ~98 kD in size, we wanted to express a part of the protein encoding the two dsRBDs only. An ~600-bp region, including the two dsRBDs, was therefore amplified by PCR...
from our 4f1-related cDNA and cloned in-frame upstream of six myc tags which are followed by the 3′ UTR and polyA tail of *Xenopus* NO38 in a pBluescript SK vector (Fig. 6 B) (Jantsch and Gall, 1992). The resulting construct, designated as MP2, was sequenced to confirm the presence of the coding region for the two dsRBDs and the six myc tags (Fig. 6 C). The putative translation product of MP2 has a calculated molecular weight of ~31 kD and should thus, like Xlrbpa, be able to diffuse into the nucleus. To determine whether MP2 was indeed able to bind dsRNA, the MP2 myc fusion and full-length Xlrbpa were expressed in *E. coli* and tested for their RNA-binding ability in a Northwestern assay. As expected, both proteins showed strong in vitro RNA-binding ability (Fig. 7, A and B).

For expression of MP2 in *Xenopus* oocytes, in vitro transcripts of the construct were injected into oocytes. Western blots of nuclear and cytoplasmic extracts were detected with mAb 9E10 directed against the myc tag (Evan et al., 1985). Although the protein migrated at a slightly aberrant molecular weight of ~38 kD, MP2 could be detected in the cytoplasm and the nucleus, indicating that MP2 carried no cryptic NLS but could freely diffuse into the nucleus (Fig. 7 C).

To determine whether MP2 would associate with ribosomes, we performed sedimentation gradient analyses of oocytes expressing MP2. 100 oocytes were injected with MP2 RNA. After overnight incubation, to allow protein synthesis to occur, the oocytes were homogenized and loaded on a sedimentation gradient. Fractions of these gradients were then analyzed for the presence of MP2 protein (Fig. 5 B). As expected, no MP2 protein could be found cosedimenting with ribosomes. Instead all MP2 protein remained at the top of the gradient indicating that MP2 protein, although RNA-binding, failed to form multimers or to associate with ribosomes.

Colocalization in situ and cosedimentation of Xlrbpa with ribosomes is indicative, but does not provide absolute proof for the physical association of the protein with ribosomes. Therefore, to test the association of Xlrbpa and TRBP with ribosomes in a more direct manner, we performed
coimmunoprecipitation experiments. Two different anti-
ribosomal autoimmune sera were used to precipitate ribo-
somes from oocyte extracts. The precipitated material was 
then tested for the presence of Xlrbpa by Western blot-
ting. Both antisera, serum 9275 and serum 13751 were able
to coprecipitate Xlrbpa with ribosomes, while all Xlrbpa 
remained in the supernatant when no antibody was added 
to the beads (Fig. 8 A). In contrast, when MP2-injected oo-
cytes were used in the same experiment, no coprecipita-
tion of MP2 protein with ribosomes could be detected 
(Fig. 8 B, and data not shown). These data suggest that 
Xlrbpa and possibly its human homologue, TRBP, are 
closely associated with ribosomes. Our control experiments 
also indicate that ribosomal association of these two pro-
teins is specific and not merely an effect of the intrinsic 
RNA-binding ability of any dsRBD-containing protein.

Figure 5. Sedimentation gradient analysis of Xlrbpa and control 
construct MP2. (A) Oocyte lysates were fractionated on sucrose 
gradients and tested for the distribution of Xlrbpa (left). UV ab-
sorbance at 260 nm was recorded for all fractions (top) indicating 
the peak of 80S ribosomes in fractions 13–23. Representative frac-
tions were tested by Western blotting with serum Rb7 for the 
presence of Xlrbpa (A). The majority of Xlrbpa could be found in 
the ribosomal peak fractions (fractions 16 and 18). In these frac-
tions, degradation bands of lower molecular weight could also be 
detected. Minor amounts of Xlrbpa could be found at the top of 
the gradient (fraction 34), corresponding to free protein. How-
ever, the free protein was mostly degraded. Unfractionated oo-
cyte extract was loaded in the first two lanes (Oo). Position of 
full-length Xlrbpa is indicated by an arrow. Ribosomal peaks 
from several primary gradients were concentrated in a Centricon 
microconcentration device (exclusion limit = 50 kD) and rerun on 
a second sucrose gradient (right). Supernatant (cen) and flow-
through (fl) from the microconcentration step was also moni-
tored for the presence of Xlrbpa. Xlrbpa was exclusively present 
in the top chamber, indicating the presence of the protein in a 
particle larger than 50 kD. UV absorbance of the gradient rerun 
was monitored (top) and representative fractions were tested for 
the presence of Xlrbpa (A). Again, the majority of Xlrbpa could 
be found in the ribosomal peak fractions. No free protein could 
be found in the supernatant fraction (34). (B) Oocyte lysates of 
MP2-injected oocytes were fractionated on sucrose gradients. 
Fractions were monitored for the presence of MP2 protein by 
Western blotting with mAb 9E10. MP2 protein could be found in 
the ribosomal peak fractions. No free protein could be found 
in the ribosomal peak fractions. Instead all MP2 remained at the top of the gradient. Un-
fractionated extracts of injected oocytes (Oo) were loaded as a 
control.

Figure 6. Schematic representation of control construct MP2. (A) 
Clone 4f1 contains two dsRBDs (gray boxes) and a NLS (hatched 
box). PCR primers used to amplify the sequence encoding the two 
dsRBDs are indicated by arrows. (B) The PCR amplified frag-
ment was cloned upstream of six tandemly arranged myc tags (open 
box) and the NO38 3' UTR and poly(A) + tail in pBluescript KS 
to give construct MP2. T3 transcripts of the construct could be 
made after linearization at a unique restriction site downstream 
of the NO38 A + tail. (C) Conceptual translation product of MP2. 
Gray boxes, the two dsRBDs. Underlining, the myc epitopes.

Figure 7. RNA-binding abil-
ity of construct MP2 and de-
tection of the protein by myc 
antibodies in oocytes. (A) Con-
trol construct MP2 and 
full-length Xlrbpa were ex-
pressed in Escherichia coli. 
RNA-binding abilities of 
both constructs were deter-
mined by Northwestern prob-
ing of E. coli lysates. Both 
proteins show strong RNA-
binding activity. (B) Detec-
tion of epitope-tagged MP2 
protein by mAb 9E10 di-
rected against the myc tag. E. 
coli lysates of cells expressing 
either MP2 or Xlrbpa protein 
were probed by Western 
blotting with mAb 9E10. 
MP2 protein is readily de-
tectable by mAb 9E10. (C) 
Detection of MP2 protein in 
microinjected oocytes. 1 cytoplasm (Cytopl) and 10 germinal ves-
cles (GV) of injected (MP2) and uninjectected (Con) oocytes were 
separated by SDS-PAGE and tested for the presence of MP2 
protein by Western blotting with mAb 9E10. MP2 protein could 
be detected in both cellular compartments indicating that MP2 
can freely diffuse into the nucleus. No protein could be detected 
in uninjected oocytes.
Xlrbpa in the Nucleus

Xlrbpa and TRBP could not only be detected in the cytoplasm, but were also present in the nucleus as shown by in situ immunostaining and Western blotting of hand-enucleated oocytes and germinal vesicles. Xlrbpa does not contain an obvious consensus nuclear localization signal (Dingwall and Laskey, 1991). Its presence in the nucleus and the relatively small molecular weight of <33 kD, therefore suggest that the protein can enter the nucleus by diffusion.

To investigate the nuclear localization of Xlrbpa and TRBP in more detail we performed immunofluorescence stainings of spread Xenopus germinal vesicles. Because of its size, the amphibian germinal vesicle allows a detailed observation of nuclear organelles like lampbrush chromosomes, amplified nucleoli and snurposomes, spherical organelles enriched in snRNPs, and several accessory splicing factors (Wu et al., 1991). Nascent transcripts on the chromosomes can easily be detected as well as pre-ribosomal RNAs on nucleoli. When oocytes injected with MP2 RNA were stained with anti-Xlrbpa antibodies, indicating the association of the protein with most, if not all nuclear RNAs. This general picture was independent of the antibody used for immunofluorescence staining (Fig. 9).

The staining on the amplified nucleoli suggests that ribosomal transcripts already associate with Xlrbpa before their export from the nucleus and assembly into mature ribosomes. This observation is in good agreement with our finding that Xlrbpa is present in ribosomes. Xlrbpa is a double-stranded RNA-binding protein; ribosomal RNAs are known to contain long, double-stranded stem-loop regions and hairpins (Noller et al., 1981), and could thus provide enough binding sites for Xlrbpa.

In addition to nucleoli, the RNP matrix of lampbrush chromosomes stained homogeneously along the length of the entire chromosome. This was somewhat surprising since it seemed unlikely that all transcripts contain sufficient double-stranded regions to allow binding of significant amounts of Xlrbpa. However, the homogeneous and relatively intense staining of all transcripts could be explained if Xlrbpa associated with nascent transcripts as part of a larger hnRNP complex (Piñol-Roma et al., 1989). In this case, not Xlrbpa itself, but other proteins within the hnRNP complex would make the contact with RNA where Xlrbpa would only bind directly to RNA if sufficient double-stranded regions were present. To test this hypothesis, we wanted to determine whether Xlrbpa was associated with hnRNP particles. We therefore made immunoprecipitations with anti-Xlrbpa antibodies and tested the precipitated material for the presence of known hnRNPs. As a precipitating antibody we used antisera Rb6 or Rb7 and the corresponding preimmune sera as controls. All antisera are able to precipitate in vitro–translated human TRBP (data not shown) and recognize human TRBP on Western blots (Fig. 2 b). For the detection of hnRNP proteins we used mAb 4F4 directed against human hnRNPs C1 and C2 (Choi and Dreyfuss, 1984). Since mAb 4F4 only recognizes human hnRNPs, and not the corresponding proteins from Xenopus, we used HeLa nucleoplasmic extracts as starting material. Both antisera Rb 6 (Fig. 10 A) and Rh 7 (not shown) were able to coprecipitate hnRNPs C1 and C2 very efficiently while no signals were seen when either preimmune serum was used. Coprecipitation of hnRNPs C1 and C2 by anti-Xlrbpa antisera was resistant to predigestion of the nucleoplasmic extracts with RNases, indicating a physical association of Xlrbpa with these hnRNPs. We thus believe that Xlrbpa is part of hnRNP particles, which would also explain its presence on nascent transcripts.

To determine whether the dsRBD-containing control construct MP2 would also associate with hnRNP particles, we performed immunoprecipitation experiments with HeLa cells which were stably transfected with a eukaryotic vector expressing MP2 protein. Immunofluorescence staining of these cell lines with mAb 9E10 showed a homogeneous distribution of MP2 protein throughout the entire cell (data not shown). Immunoprecipitations of stably expressing cell lines were performed with either mAb 4F4 directed against hnRNPs C1 and C2, or with polyclonal serum Rb6 directed against Xlrbpa (TRBP). Immunoprecipitates were then probed with mAb 9E10 for the presence of myc-tagged MP2 protein (Fig. 10 B). In these experiments, MP2 protein could not be coprecipitated with either antibody, indicating that MP2 protein does not associate with hnRNP particles.

The positive staining of snurposomes indicated the presence of Xlrbpa in these structures. Two types of snurposomes (B and C) can be distinguished in Xenopus GV.
The smaller B snurposomes, which can sometimes be found associated with C snurposomes, are highly enriched in the five splicing snRNPs, U1, U2, U5, and U4/U6 (Wu et al., 1991). Besides these snRNPs, accessory splicing factors such as SC-35 or SR proteins can be found in these structures (Roth et al., 1991; Zahler et al., 1992). However, also hnRNPs can be detected in B snurposomes (Wu et al., 1991). In contrast, no hnRNPs could so far be detected in the generally larger C snurposomes. Instead, C snurposomes seem to contain U7 snRNPs as the only snRNP component (Wu and Gall, 1993). The presence of p80 coilin in these structures suggests a homology of C snurposomes and coiled bodies found in somatic cells (Wu et al., 1994; Gall et al., 1995).

In our immunofluorescence stainings, C snurposomes labeled generally more intensely than B snurposomes, suggesting a higher concentration of Xlrbpa in the former structures (Fig. 9). This was somewhat surprising since no hnRNPs could so far be detected in C snurposomes. However, since all snurposomes contain snRNPs, we wanted to determine whether the presence of Xlrbpa in snurposomes could be due to an association of the protein with snRNPs. We performed immunoprecipitation experiments with anti-Xlrbpa and anti-snRNP antibodies and tested the precipitated material for the presence of snRNP proteins or Xlrbpa, respectively. In these experiments, no association of Xlrbpa with snRNPs could be detected (data not shown). It thus seems unlikely that Xlrbpa is associated with snRNAs in snurposomes. Instead, Xlrbpa could be associated with other hnRNPs, at least in B snurposomes.

Discussion

We have cloned and characterized Xlrbpa, a Xenopus double-stranded RNA-binding, 33-kD protein. Our data indicate that Xlrbpa is the putative homologue of human TRBP (Gatignol et al., 1991). Both proteins show a high degree of homology at the primary amino acid level and in their overall organization. They contain three tandemly arranged dsRBDs. In both proteins, the first two dsRBDs show a high degree of homology to a dsRBD consensus sequence while the third dsRBD fits the consensus only at its carboxy-terminal end (St Johnston et al., 1992). The dsRBD has been shown to exclusively interact with double-stranded RNAs or RNA–DNA hybrids (St Johnston et al., 1992; Bass et al., 1994). A preference for these two substrates could be explained by the similar structures exhibited by these double-stranded molecules; both RNA–DNA and RNA–RNA helices assume A form helices, which differ considerably from the B form helices adopted...
by most double-stranded DNAs. Although both substrates, RNA–RNA and RNA–DNA duplexes have been shown to be recognized by the dsRBD in vitro, no example of an RNA–DNA hybrid substrate has been shown to be the natural substrate for any known dsRBD-containing protein. In contrast, in all cases where the substrate for a dsRBD containing protein has been identified, it represents an RNA duplex molecule. Consistent with this idea, we could demonstrate that Xlrbpa and human TRBP are associated with ribosomes and hnRNAs. In both cases RNA–RNA hybrids rather than RNA–DNA hybrids seem the likely substrates of these proteins.

**Xlrbpa, a Ribosomal Protein?**

Xlrbpa is ubiquitously expressed in essentially all tissues tested and it seems likely that human TRBP is also ubiquitously expressed in humans (Gatignol et al., 1991). However, we observed marked differences among the tissues tested. Especially nerve and brain tissue showed a very low level of Xlrbpa, while oocytes had the highest concentrations. These differences in concentration could possibly be explained by our finding that Xlrbpa and TRBP associate with ribosomes. It is known that amphibian oocytes contain extremely high levels of ribosomes. In contrast, although neurons contain considerable amounts of ribosomes, nervous tissue as a whole has a relatively low ribosome content. Thus, it appears that Xlrbpa concentration resembles the concentration of ribosomes in a given tissue. Since ribosomal RNAs represent the majority of all RNAs in a cell, it is also obvious that the low amounts of Xlrbpa (or TRBP) assembled with hnRNPs do not fall into account for this comparison.

We could demonstrate the association of Xlrbpa and TRBP with ribosomes by immunofluorescence, cofractionation on density gradients, and coimmunoprecipitation experiments. Additionally, we could purify ribosomal RNAs in immunoprecipitation experiments with anti-Xlrbpa antibodies (data not shown). Sticking, and therefore nonspecific association of Xlrbpa (or human TRBP) with ribosomes, could be excluded by our control experiments in which a dsRNA-binding control construct, containing two dsRBDs failed to associate with ribosomes. Taken together, we are confident that Xlrbpa and TRBP are closely associated with ribosomes. This finding is also supported by our previous observation that Xlrbpa can bind RNA in vitro (Krovat and Jantsch, 1996). A ribosomal association of Xlrbpa or TRBP is also consistent with the recent finding that TRBP can interact with the ribosome-associated kinase PKR (Benkirane et al., 1997). Ribosomes and their components have been intensely studied over the last decades. It is thus surprising that neither Xlrbpa, nor human TRBP have been described as ribosomal proteins, especially when considering their relative abundance. However, most ribosomal preparations include a high salt wash to remove associated proteins. dsRBDs lose their in vitro–RNA-binding ability at salt concentrations above 300 mM (Jantsch, M., unpublished data). Consistently, we observed a dissociation of Xlrbpa from ribosomes in sedimentation gradients at high salt concentrations, which could explain its absence from standard ribosomal preparations. Thus, we believe that Xlrbpa is merely a ribosome-associated factor, but not a necessary component of translationally active ribosomes.

**Xlrbpa on Nascent Transcripts**

Xlrbpa and human TRBP could also be found in the nucleus. Although both proteins lack a clear nuclear localization signal, they seem small enough to enter the nucleus by simple diffusion through nuclear pores. In nuclear spreads of *Xenopus* GVs, we could localize the protein in essentially all RNA-containing structures: amplified nucleoli,
lambrush chromosomes, and snurposomes. The nuclear localization is in agreement with our finding that Xlrbpa and TRBP are associated with ribosomes. It also suggests that the proteins can associate with ribosomal RNAs, possibly even nascent pre-ribosomal RNAs before they are processed, exported to the cytoplasm, and assembled into mature ribosomes.

The localization of Xlrbpa on lambrush chromosomes indicated an association of the protein with essentially all nascent transcripts. Xlrbpa binds exclusively to double-stranded RNAs in vitro (St Johnston et al., 1992; Bass et al., 1994). This raises the question as to how the protein can associate with the many transcripts lacking double-stranded regions. However, the homogeneous distribution of Xlrbpa on essentially all hnRNAs could be explained by our finding that Xlrbpa is associated with hnRNP complexes. In this case, not Xlrbpa itself but the entire hnRNP complex, including Xlrbpa, would associate with the nascent transcripts. Thus, Xlrbpa would only bind RNA when sufficient double-stranded regions are available on the corresponding transcript. In the absence of extensive double-stranded structures, however, other hnRNP proteins within the complex would bind to the RNA. The decision of which RNA-binding protein within an hnRNP complex binds directly to RNA would thereby be governed by competition of the proteins for a suitable binding substrate. The fact that our control construct, MP2, fails to associate with hnRNP particles indicates that the association of Xlrbpa or human TRBP with hnRNPs is specific, and not caused by the intrinsic RNA-binding ability of the two proteins.

An analogous situation has been found for the distribution of snRNPs and splicing components in amphibian GV's. Surprisingly, all components of the splicing machinery can be found on most nascent transcripts irrespective of whether those transcripts contain suitable splice sites or not (Wu et al., 1991). Thus, it seems as if splicing components associate with nascent transcripts, possibly as part of larger RNP complexes, independent of the availability of processing sites. This situation is similar to the one found by us, where Xlrbpa is associated with essentially all nascent transcripts seemingly independent of the availability of secondary structures required for the direct binding of the protein to RNA. Instead, Xlrbpa seems to be associated with transcripts as part of a larger hnRNP complex.

Besides an association of Xlrbpa with nucleoli and nascent transcripts we could find the protein in snurposomes within the *Xenopus* GV. Snurposomes are spherical structures of 1–10 μm diam, which can be distinguished based on their structure and molecular composition in B and C snurposomes. In *Xenopus*, the smaller B snurposomes are highly enriched in the five splicing snRNPs, U1, U2, U5, and U4/U6 as well as several accessory splicing factors (Wu et al., 1991). Additionally, staining with anti-hnRNP antibodies indicates the presence of hnRNPs in B snurposomes (Wu et al., 1991). The generally larger C snurposomes, in contrast, seem to lack splicing snRNPs but contain the U7 snRNP (Wu and Gall, 1993). C snurposomes frequently have B snurposomes attached to their surface or internalized and are thus also called sphere organelles. These structures also contain p80 coilin, suggesting that sphere organelles are homologous to coiled bodies found in several somatic cells (Wu et al., 1994; Gall et al., 1995).

No hnRNPs are detectable in C snurposomes (Wu et al., 1991). The presence of Xlrbpa in B snurposomes and its homogeneous distribution on nascent transcripts resembles that of splicing snRNPs and splicing factors, as well as that of hnRNP proteins in the amphibian GV closely. However, in immunoprecipitations with the anti-Sm antibody, Y12 (Lerner et al., 1981), no association of Xlrbpa with splicing snRNPs could be detected. It is thus likely that Xlrbpa is either present as free protein in B snurposomes or, alternatively, associated with the other hnRNP proteins found in these structures. Since Xlrbpa can be found associated with hnRNP particles, the latter hypothesis appears more likely.

The presence of Xlrbpa in C snurposomes is more problematic. In view of the fact that anti-Sm antibodies were unable to precipitate Xlrbpa, an association of the protein with the U7 snRNP seems very unlikely. Also, no other known hnRNPs could so far be detected in these organelles. Additionally, in HeLa cells we observed no indication for the presence of the Xlrbpa homologue, TRBP, in coiled bodies, which are believed to be homologous to sphere organelles. All these facts make a reasonable explanation for the presence of Xlrbpa in C snurposomes very difficult at this point. However, future studies will be aimed to explain the mechanism that is responsible for the localization of Xlrbpa in these structures.

### Possible Functions of Xlrbpa

Human TRBP has been isolated by its ability to bind HIV TAR-RNA in vitro, and has been suggested to be a human-specific factor specifically required for the replication of HIV in human cells (Gatignol et al., 1991). Binding specificity of TRBP for TAR-RNA has been determined by the introduction of mutations in this RNA which abolished binding of the protein to this sequence (Gatignol et al., 1991). However, several points argue against the notion that human TRBP is a human specific factor necessary for the replication of HIV.

First, TRBP does not seem to be exclusively present in human cells. Our data strongly suggests that Xlrbpa is the *Xenopus* homologue of TRBP, thus demonstrating the presence of a TRBP homologue in an organism incompatible with the replication of HIV.

Second, Xlrbpa and human TRBP contain three tandemly arranged dsRBDs. So far, no sequence-specific binding of dsRBDs could be determined in vitro. In contrast, Xlrbpa has been isolated by its ability to bind U1 or U2 snRNAs in vitro, but can also bind poly rI/rC very efficiently (St Johnston et al., 1992; Krovat and Jantsch, 1996). Also, the third dsRBD in the *Drosophila* Staufen protein binds both U1 snRNA and poly rI/rC in vitro (St Johnston et al., 1992). None of these substrates used for in vitro assays seem to be the natural substrates of these proteins. Thus it is also possible that TAR-RNA is not the in vivo substrate for human TRBP. TAR-RNA forms a stable double-stranded hairpin region. To prove the binding specificity of TRBP for TAR-RNA, several mutations were introduced in this RNA, some of which abolished binding of TRBP. However, all mutations affecting TRBP binding were located in the base-paired region of the TAR-RNA hairpin, thus destabilizing the double-stranded region (Gati-
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It has been shown that chloramphenicol transferase (CAT)-reporter genes under the control of viral promoters are more strongly expressed when TRBP was cotransfected in these cells (Gatignol et al., 1991). This effect has been proposed to be caused by a stimulation of these promoters by TRBP. However, it has not been shown that transcription from the promoter itself was increased by TRBP. Recently, another study reported increased levels of expression of transfected genes upon cotransfection of either TRBP or the vaccinia virus E3L gene, another dsRBD-containing protein (Park et al., 1994). In this case, increased expression of the transfected genes correlated in part with increased mRNA amounts of the corresponding genes. Both effects, increased reporter gene activity and increased mRNA amounts of certain transfected genes, could be explained in part by mRNA stabilization mediated by Xlrbpa or TRBP. We could show that TRBP is part of hnRNP particles. It is thus possible that Xlrbpa or TRBP can stabilize certain mRNAs with extensive secondary structures by binding to them.

It has also been shown that human TRBP acts as an inhibitor of the cellular, double-stranded–activated, interferon-dependent protein kinase PKR (Park et al., 1994). This kinase phosphorylates translation initiation factor, eIF2α, when activated by double-stranded RNAs upon viral infection, resulting in an inhibition of translation (Kaufmann et al., 1989; Manche et al., 1992). Inhibition of PKR by TRBP and other dsRBD-containing proteins has been attributed to the masking of double-stranded RNAs, thereby preventing binding and subsequent activation of PKR by these substrate RNAs (Watson et al., 1991; Langland et al., 1994; Park et al., 1994). Additionally, TRBP has been suggested to be involved in the masking of cellular mRNAs containing extensive double-stranded regions, thus precluding inadvertent activation of PKR by cellular RNAs. This view is consistent with our finding that Xlrbpa or TRBP associate with essentially all cellular double-stranded RNAs. Binding of Xlrbpa or TRBP to double-stranded RNAs might therefore help to prevent the activation of PKR by these RNAs.

However, our finding that Xlrbpa is associated with ribosomes can also be interpreted in an alternative way. Rather than masking ribosomal RNAs to avoid activation of PKR, Xlrbpa or TRBP might compete with the ribosome–associated kinase PKR for suitable substrates directly at the ribosome. Although no substrate specificities of dsRBDs could so far be determined in vitro, biological data suggests that dsRBD-containing proteins can bind in a sequence-specific manner (St Johnston et al., 1991). It is therefore possible that activation of PKR in vivo depends on the availability of high affinity substrates for this kinase. Xlrbpa would thereby act as a competitor to PKR and prevent the binding of low affinity substrates to the kinase by binding to them at the ribosome.

RNA-dependent dimerization of TRBP and PKR has recently been demonstrated in the yeast two hybrid system and in vitro (Cosentino et al., 1995). Direct (RNA-independent) interaction of TRBP and PKR has also been reported recently (Benkirane et al., 1997). The same study shows that TRBP can down-regulate PKR activity in an RNA-independent manner, possibly through direct protein–protein contact (Benkirane et al., 1997). Association of TRBP or Xlrbpa with ribosomes could therefore be mediated by an association of these proteins with PKR, a kinase which has also been shown to be ribosome associated (Levin and London, 1978). However, to demonstrate RNA-independent down-regulation of PKR by TRBP, a mutant version of TRBP (defective in its second dsRBD and thus unable to bind dsRNA) was used (Benkirane et al., 1997). A similar mutation in Xlrbpa, in which the second dsRBD has either been inactivated by mutation or completely deleted, still shows significant RNA-binding activity in our hands (Krovat and Jantsch, 1996). This raises the question of whether down-regulation of PKR by a mutated TRBP is indeed mediated by a completely RNA-independent mechanism or, alternatively, whether residual RNA-binding activity of the mutated TRBP might lead to the observed effect, possibly by competition of TRBP for dsRNAs that activate PKR.

Nonetheless, no PKR homologous kinase has so far been described in Xenopus. Therefore, several other functions can also be envisaged for Xlrbpa. On the one hand, Xlrbpa might act as a general hnRNP whose main function is the complexation and packaging of RNA. Complexation of RNA with hnRNPs has been shown to be important for the proper processing and export of RNA (Bennet et al., 1992; Mayeda and Krainer, 1992; Piñol-Roma and Dreyfuss, 1993). At the same time, hnRNP might stabilize and protect RNA from degradation. On the other hand, Xlrbpa might itself influence the folding state of RNA. Since Xlrbpa binds exclusively to double-stranded regions it might either help in the formation of such structured regions, stabilize such regions or, alternatively, destabilize double-stranded regions. Double-strand annealing and unwinding activities have also been reported for other RNA-binding proteins (Cobianchi et al., 1993; Portman and Dreyfuss, 1994; Xiao et al., 1994). The structure of two dsRBDs from different proteins has recently been solved and showed a conserved structure for those two dsRBDs (Bycroft et al., 1995; Kharrat et al., 1995). Mutational analyses of Xlrbpa suggest that the dsRBDs found in Xlrbpa assume the same conserved tertiary structure (Krovat and Jantsch, 1996). However, little is known about the way double-stranded RNA-binding domains interact with RNA which makes an assessment about other functions of Xlrbpa purely speculative at this point. Nonetheless, studies to determine the mode of interaction of dsRBDs with RNA are under way and will hopefully help to shed some light on these unsolved questions. It will also be interesting to determine whether Xlrbpa and its human counterpart, TRBP, play a role in mRNA turnover and translation of cellular RNAs.

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