

## RNA-Binding Domain Proteins in Kinetoplastids: a Comparative Analysis†

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**RNA-binding proteins are important in many aspects of RNA processing, function, and destruction. One class of such proteins contains the RNA recognition motif (RRM), which consists of about 90 amino acid residues, including the canonical RNP1 octapeptide: (K/R)G(F/Y)(G/A)FVX(F/Y). We used a variety of homology searches to classify all of the RRM proteins of the three kinetoplastids *Trypanosoma brucei*, *Trypanosoma cruzi*, and *Leishmania major*. All three organisms have similar sets of RRM-containing protein orthologues, suggesting common posttranscriptional processing and regulatory pathways. Of the 75 RRM proteins identified in *T. brucei*, only 13 had clear homologues in other eukaryotes, although 8 more could be given putative functional assignments. A comparison with the 18 RRM proteins of the obligate intracellular parasite *Encephalitozoon cuniculi* revealed just 3 RRM proteins which appear to be conserved at the primary sequence level throughout eukaryotic evolution: poly(A) binding protein, the rRNA-processing protein MRD1, and the nuclear cap binding protein.**

Organisms of the order Kinetoplastida are flagellated protists with a microscopically visible complex of mitochondrial DNA. They include a number of pathogens of animals and plants. Among these organisms are *Leishmania major*, which causes a range of diseases in the tropics and Europe; *Trypanosoma brucei*, which causes sleeping sickness in Africa; and *Trypanosoma cruzi*, which is the etiological agent of Chagas' disease in Latin America. All three parasites undergo a “digenetic” life cycle involving transmission from one mammal to the next by an arthropod vector.

Some aspects of RNA metabolism in trypanosomes and leishmanias are similar to those of other eukaryotes, whereas others are quite deviant. As in other organisms, multiple stable and structural RNAs must be processed, modified, and assembled into ribonucleoprotein complexes. Some of these complexes are themselves involved in further RNA-processing reactions. Perhaps the most remarkable aspect of trypanosomatid RNA metabolism is the editing of kinetoplast mRNA transcripts through the addition and deletion of uridine residues (46). Almost as surprising, however, is the dependence of the parasites on post-transcriptional mechanisms to control the levels and translation efficiencies of mRNAs. Trypanosomatid protein-coding genes are arrayed in long polycistronic transcription units, for which specific promoters have proved elusive (34, 35). Monocistronic mature RNAs are generated by *trans* splicing (at the 5' end) and polyadenylation (at the 3' end) (30). The cotranscription of many open reading frames, irrespective of the cell's requirement for differing levels of the final gene products, results in an almost exclusive dependence upon posttranscriptional mechanisms for the regula-

tion of gene expression (8). Although control could theoretically act during mRNA processing or export from the nucleus, most mRNAs tested so far have been found to be regulated primarily at the levels of mRNA degradation and translation; the signals determining the regulation are usually located in the 3' untranslated regions (UTRs) (8).

Posttranscriptional processing and degradation of RNAs are effected by enzymes and ribonucleoprotein complexes and regulated by *trans*-acting factors that bind the RNAs (14). In mammalian cells, several hundred unstable mRNAs contain AU-rich elements (AREs) in their 3' UTRs. Proteins implicated in the regulation of ARE-RNA stability fall into several classes, depending on the RNA-binding domains: these include KH domains, zinc fingers, and RNA recognition motif (RRM) domains (45). Similar AREs are present in various regulated mRNAs in kinetoplastids, and there is evidence that the mechanisms of regulation, including the involvement of RRM-containing regulatory proteins, may resemble those in mammals (12, 44).

The RRM comprises about 90 amino acid residues. It often contains the signature RNP1 sequence motif, (K/R)G(F/Y)(G/A)FVX(F/Y). Proteins with RRMs are involved in a large number of processes through specific interactions with RNA (14), but a subset of RRMs can also interact with other proteins (27). The signature sequence RNP1, together with a second one known as RNP2, form the central part of the RRM beta-sheet which is involved in RNA binding. In this article, we describe the proteins containing RRM domains from the three sequenced trypanosomatid genomes and compare the gene set with those from other eukaryotes, particularly the minimal eukaryote *Encephalitozoon cuniculi*.

### MATERIALS AND METHODS

**Databases.** The databases utilized in this work were obtained from sequencing projects of the three trypanosomatids. The main databases used were the *T. cruzi* database (TcBr version 2.0), the *T. brucei* database (Tb927 versions 2.1 and 3), and the *L. major* database (LmjF version 3), but the final results were updated

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TABLE 1. RRM-containing proteins in selected genomes

Genome	No. of RRM proteins	No. of RRM domains	Total no. of ORFs	% proteins with RRM	RRM no./ORF no.
<i>Encephalitozoon cuniculi</i>	18	30	1,909	0.94	0.016
<i>Schizosaccharomyces pombe</i>	67	107	4,956	1.35	0.022
<i>Plasmodium falciparum</i>	61	95	5,251	1.16	0.018
<i>Saccharomyces cerevisiae</i>	50	86	6,218	0.80	0.014
<i>Leishmania major</i>	78	120	8,277	0.94	0.014
<i>Trypanosoma brucei</i>	67	100	8,599	0.78	0.012
<i>Brachydanio rerio</i>	150	273	9,249	1.62	0.030
<i>Drosophila melanogaster</i>	177	304	16,175	1.09	0.019
<i>Caenorhabditis elegans</i>	130	223	21,926	0.59	0.010
<i>Homo sapiens</i>	308	542	35,844	0.86	0.015

using the July 2005 versions, including homologues derived from aligning the genomes of all three species (5, 15, 16, 24). The European Molecular Biology Open Software Suite (EMBOSS) (<http://www.ebi.ac.uk/emboss/>) and various other programs available from GeneDB were used to extract sequences from the databases.

**Sequence searches.** A selection of RRM-containing sequences from trypanosomes, *Saccharomyces cerevisiae*, and *Homo sapiens* were used in BLAST searches against trypanosomatid databases. BLAST hits having log E values of <9 were used in further analysis. The motif search utility within GeneDB was also used to detect RNA-binding proteins. The searches were continued until no more potential proteins with RRM domains could be located.

All RRM-containing protein sequences were run against the Pfam (<ftp://ftp.sanger.ac.uk/pub/databases/Pfam/>) set of domains, hidden Markov model profiles, and Prosite databases ([ftp://bo.expasy.org/databases/prosite/tools/ps\\_scan/](ftp://bo.expasy.org/databases/prosite/tools/ps_scan/)). Potential RNA-binding proteins were identified using the following RNA recognition motifs: Pfam PF00076, Smart SM00360, Prosite PS50102, and Prosite PS00030. All of the *T. brucei* protein sequences were then compared against the entire EMBL database using BLASTP and against the *Saccharomyces cerevisiae* genome. A functional assignment was made for a predicted protein only when a BLAST search using the *T. brucei* sequence gave the corresponding functionally characterized protein from another eukaryote with a low (usually less than  $10^{-13}$ ) P value and a reciprocal BLAST search with the characterized protein sequence also gave the *T. brucei* protein as the best match. Matches within the RRM alone usually gave probabilities in the range of  $10^{-11}$  to  $10^{-12}$  and were not used for functional assignments; matches outside the RRM, and the presence of appropriate domains, were also required. We did not infer function because of matches with proteins for which function had been predicted on the basis of homology or was designated "putative."

Proteins with clear functional assignments were named according to the homologue in yeast or mammals or using published names. Other proteins were named according to the following scheme, where x is a number: One RRM, RBPx; two RRM domains, DRBDx; three RRM domains, TRRMx; RRM(s) and an SR domain(s), RBSRx. Since the number of RNA-binding domains recognized depended on the cutoff used, and sometimes the apparent numbers of domains were not the same in all three organisms, these names are only an approximate guide.

**Alignments and orthologues.** Each of the sequences obtained for one kinetoplastid species was used as a query in reciprocal multiple BLAST searches against the two other genomes in order to identify orthologues. The assignments were confirmed manually by examining the protein lengths and domain structures; then, orthologous proteins were assigned similar names. Multiple sequence alignments were performed using CLUSTALX v1.81 (47) with the default alignment parameters, and this information was also used in orthologous identification. The orthologue assignment was then checked against orthologues assigned based on aligning all three genomes (clusters of orthologous groups [COGs]) (15). Nearly all assignments were consistent. The exceptions were found exclusively among families of genes encoding proteins of unknown function with a single RRM, such as the RBP7 family; often, the more divergent orthologue had acquired additional sequence.

**NJ trees.** Protein distances were calculated by Clustal algorithms using the DNASTar package (species-specific trees) or the CLUSTALX v1.81 program (comparative tree). For the calculation, the neighbor-joining (NJ) method was used. Percent divergences between all pairs of sequence from the multiple

alignments were calculated, and the NJ method was applied to the distance matrix. The unrooted tree obtained was plotted using Treeview software (<http://taxonomy.zoology.gla.ac.uk/rod/treeview.html>) (39).

## RESULTS

**Overview of RRM-containing proteins in trypanosomes.** Predicted proteins containing RRM domains were identified from the complete predicted proteomes of *T. brucei*, *T. cruzi*, and *L. major* (tri-tryp). A total of 139 RRM-type proteins were found in *T. cruzi*, 75 in *T. brucei*, and 80 in *L. major*. When these were classified as orthologues by sequence similarity and by comparing the positions of the genes on all three genomes (see Materials and Methods), 77 different RRM proteins were found. In most cases, the *T. cruzi* genome contains two or more paralogues, presumably because the CL Brener clone used for sequencing has a hybrid genotype (31). In addition, some genes were present in two or more copies in either *T. brucei* or *L. major* as a consequence of tandem duplications. A complete list is provided in the supplemental material. Two genes were unique to *T. cruzi*, three were unique to *L. major*, and seven were found in *T. cruzi* and *T. brucei* but not *L. major*.

We compared the protein set from *T. brucei* with those of several other organisms, relying mainly on annotation available from the Pfam database (Table 1 and Fig. 1). Overall, the number of proteins with RRM domains—and the number of domains in the proteins—increased with genome complexity. We have previously noted that in comparison with *S. cerevisiae*, trypanosomes have more proteins that are less than 200 residues long and have a single RRM (24). The broader comparison, however, suggests that trypanosomes are not remarkable in their RRM protein content. The only irregularity visible in

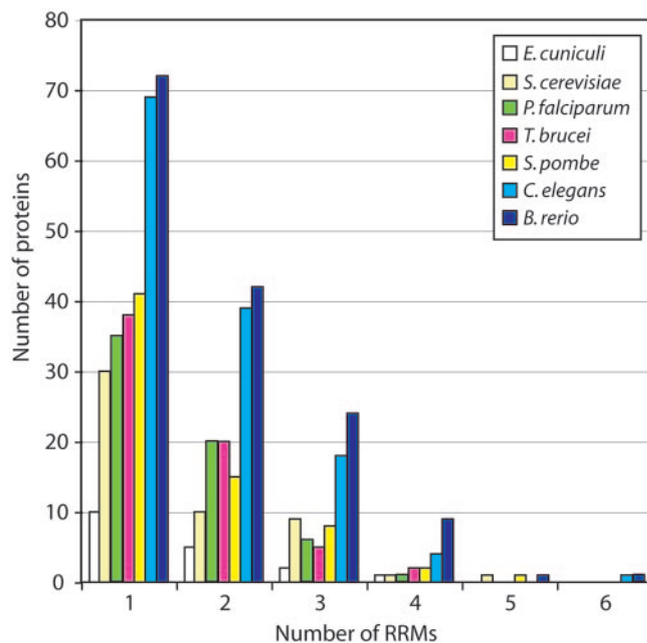


FIG. 1. Comparison of RRM proteins in *Encephalitozoon cuniculi*, *Trypanosoma brucei*, *Saccharomyces cerevisiae*, *Plasmodium falciparum*, *Schizosaccharomyces pombe*, *Caenorhabditis elegans*, and *Brachydanio rerio*. The proteins were classified into groups according to the number of RRM domains.





FIG. 3. Domain structures of *T. brucei* proteins found mainly in the lower half of Fig. 2. Locus numbers are in the left column, and protein names are on the right. Different domains are specific colors as shown on the adjacent blocks; the strongest RNA-binding-domain matches are black, and the weakest are light gray. Proteins encoded by adjacent genes are bracketed.

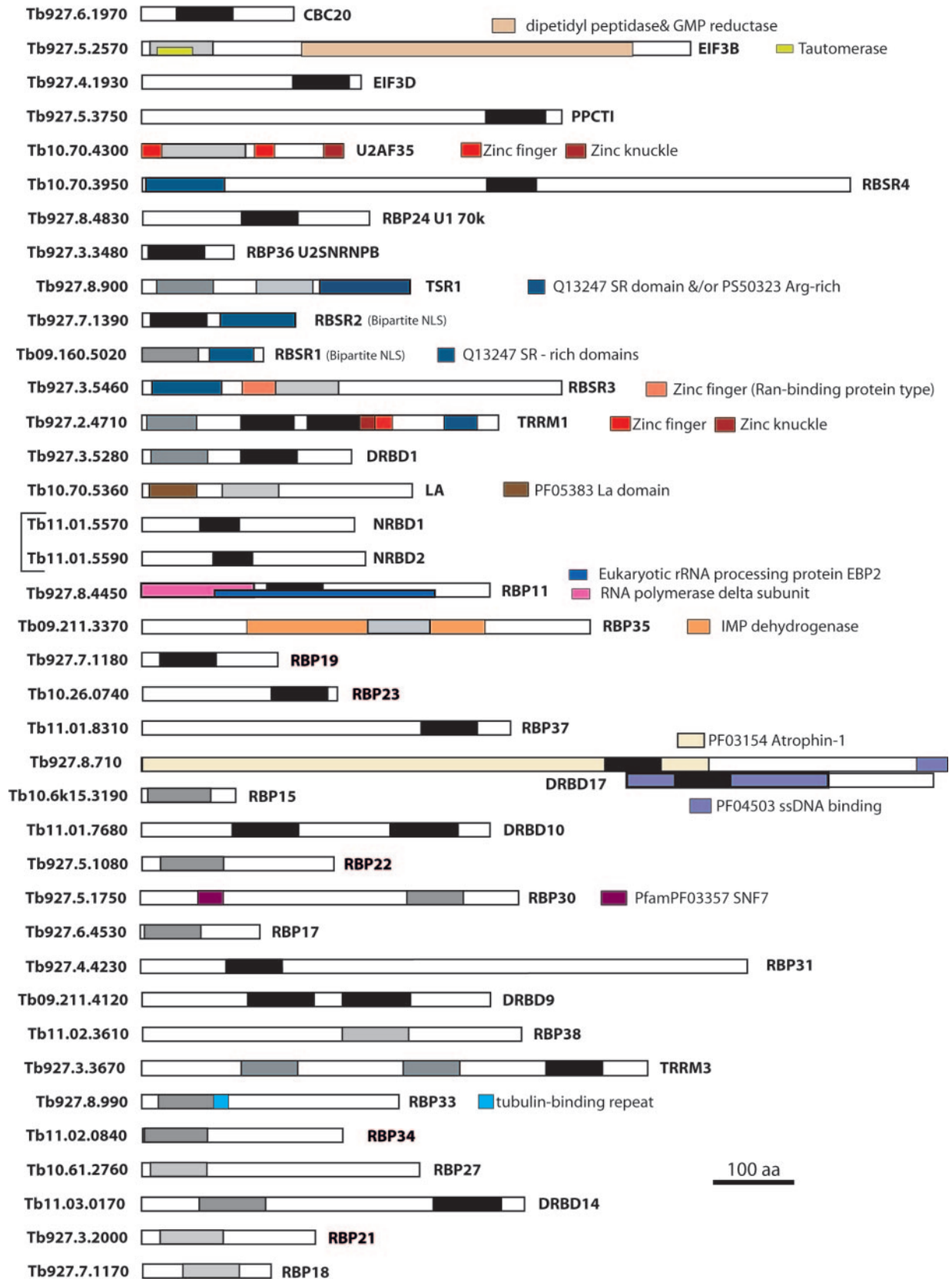


FIG. 4. Domain structures of *T. brucei* proteins found mainly in the upper half of Fig. 2. The key is as for Fig. 3. DRBD17 did not fit and so is shown broken into two portions.

TABLE 2. Accession numbers of conserved proteins<sup>a</sup>

Protein	<i>Trypanosoma brucei</i>	<i>Brachydanio rerio</i>	<i>Caenorhabditis elegans</i>	<i>Plasmodium falciparum</i>	<i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	<i>Encephalitozoon cuniculi</i>	<i>Entamoeba histolytica</i>
CBC20	Tb927.6.1970	O8IGR6_BRARE	O93594_CAEEL	PDF0750w	YPL178W	SPBC13A2.01c	Q8SW46_ENCCU	100.m00125
hnRNPH	Tb927.2.3880	Q8W3W8_BRARE	Q22708_CAEEL	PFL1170w	YER165W	SPAC57A7.04c	Q8SR30_ENCCU	344.m00050
PABP1	Tb09.211.0930	OQP3L1_BRARE	O9U302_CAEEL					193.m00064
PABP2	Tb09.211.2150	O7T3G5_BRARE	Q19581_CAEEL	PFL0830w	YPR112C	SPBP22H7.02c	Q8SRD9_ENCCU	37.m00220
MRD1	Tb927.8.4170	O6DR16_BRARE	Q9XU67_CAEEL	PEI1175c	YIR001C	SPBC16E9.12c	Q8SWN2_ENCCU	
SGN1	Tb927.6.3870	AAH79522.1	AA46772.1	PFE0885w	YOR361C	SPAC25G10.08		172.m00082
EIF3B	Tb927.5.2570	gi168436531	CAA21681.1	PF08_0086		SPBC17G9.05		43.m00204
PPCT1	Tb927.5.3750	O6PGV9_BRARE	O8I4J2_CAEEL	PF11_0200	YPR107C	SPAP8A3.06	Q8SOL5_ENCCU	13.m00293
U2AF35	Tb10.70.4300	O8JHI3_BRARE	O9U2U0_CAEEL	PFL11705w	YER068W	SPAC16C9.04c	Q8SQS6_ENCCU	57.m00152
MOT2P		O7SYDI_BRARE	O9GYO9_CAEEL					

<sup>a</sup> GenBank numbers except for *T. brucei*, which are GeneDB locus numbers.

(29), and to be associated with importin alpha and three novel subunits, none of which is obviously homologous to CBP80; the trypanosome cap-binding complex was also shown to be essential in early steps of *trans* splicing (29).

At the top of Fig. 3 are two possible hnRNP-like proteins, similar to those found in mammals but not in yeast. The protein which is annotated as hnRNPA (24) has an N-terminal glycine-rich region with a few RGG motifs and a single RRM; mammalian hnRNPA has, in contrast, two RRMs and a more extensive glycine-rich domain that contains proportionately more RGG motifs and is at the C terminus. The hnRNPH homologue is considerably more similar to the mammalian counterparts. Both hnRNPA and hnRNPH family proteins affect several aspects of RNA processing. hnRNPH is required for splicing enhancer activity in some viral messages (7), and both proteins have been implicated as regulators of alternative splicing of endogenous mRNAs (19, 20). Regulated shuttling of hnRNPA into the cytoplasm (1) means that this protein could also be involved in the control of cytoplasmic processes, such as translation (6).

In addition to these proteins, there are several with serine-arginine-rich domains, which are suggestive of an involvement in splicing (22). RBSR3 also has a Ran-binding-type zinc finger. TRRM1, a protein with three RRMs, a zinc finger and a zinc knuckle, and an SR domain, was shown to be located in nuclear speckles (32). RBSR1 and RBSR2 have bipartite nuclear localization signals; RBSR3 has a zinc finger, suggestive of binding to the nuclear import factor Ran. TSR1 was previously shown to be in the nucleus of *T. brucei* and to be capable of interacting with two different parts of the spliced leader RNA in the yeast three-hybrid system (23); its role is as yet unclear, but it is clearly not, as originally thought, a direct homologue of U1 70k.

**Processing of stable and nuclear RNAs.** The two related nuclear-RNA-binding proteins NRBD1 and NRBD2 (previously p34 and p37) (51) were shown to be involved in the import and/or assembly pathway of 5S rRNA during ribosome biogenesis in trypanosomes (43). The two genes are neighbors on chromosome 11 and are likely to be products of gene duplication. The multiple RNA-binding domain 1 protein (MRD1) is involved in rRNA processing in yeast (25). The La protein is implicated in the maturation of various RNAs, including the SL RNA and tRNA (17, 33). RBP11 has domains suggestive of involvement in rRNA processing or transcription. Each of three related and chromosomally linked proteins with two RRMs, DRBD6A, 6B, and 11, has a bipartite nuclear localization signal, suggesting a nuclear function.

**Translation.** There are two poly(A) binding proteins (PABPs). PABPs are involved throughout the lifetime of polyadenylated mRNAs, with roles including stabilization and the interaction of the poly(A) tail with the translation initiation complex. A similar function in transcript stabilization has also been demonstrated in trypanosomatid extracts (37); poly(A) binding was demonstrated for *L. major* and *T. cruzi* PABP1 (2, 3) and *T. brucei* PABP2 (21). One of the two *L. major* PABP2 genes is classified as a singleton in the published COG analysis (15), having been isolated on a nonsyntenic chromosome.

We found two possible components of eukaryotic initiation translation factor 3 (EIF3). The EIF3 $\beta$  homologue EIF3B is clear, whereas the putative EIF3D homologue is less convinc-

ing. In humans, EIF3 is a factor composed of 11 subunits which interacts with a 40S ribosomal subunit and is involved in the formation of the 43S preinitiation complex (28, 42). Yeast SGN1 is thought to influence translation (50), so the trypanosome homologue may also have this function. DRBD1 matches TIA-like RNA-binding proteins, but this may be only a consequence of the arrangement of the RRM. TIA proteins have been implicated in both translational silencing and control of alternative splicing (10, 18). Finally, peptidyl-prolyl *cis-trans* isomerase (PPCTI) is involved in protein refolding and is ubiquitous in living organisms, although more than one class of proteins with this function exists (36).

**Possible regulators of mRNA turnover.** Several of the proteins in Fig. 3 are notable for the presence of low-complexity sequences, such as glutamine-rich or glycine-rich regions. The *T. cruzi* RBP family proteins were shown to have binding specificity for G- or U-rich RNA (9), and UBP1 and UBP2 have been implicated in posttranscriptional control of gene expression in both *T. cruzi* and *T. brucei*. AREs and G-rich elements are present in the 3' UTR of the small mucin gene of *T. cruzi* and confer stage-specific mRNA degradation/stability (11). *T. cruzi* UBP1 (TcUBP1) and TcUBP2 bind to the 3' UTRs of small mucin gene RNAs and promote RNA stabilization, forming a ribonucleoprotein complex that includes poly(A) binding protein (13). The UBP1 and UBP2 genes were probably produced by a gene duplication event that occurred before the divergence of trypanosomes and leishmanias, as they are present as neighbors on orthologous chromosomes in all three species. We have recently found that TbUBP1 and TbUBP2 are also implicated in posttranscriptional gene regulation in *T. brucei* (C. Hartmann, unpublished data). In TcUBP1, an extra beta hairpin enlarges the surface of RNA binding in the RRM domain (49). The functions of the other single-RRM proteins in this group (from UBP1 to DRBD13 in Fig. 3) remain to be elucidated. In BLASTP alignments against the EMBL databases, the proteins of known function with the best matches are often from the ELAV family, but this is only because the single RRM of the trypanosome proteins align with the three RRM of the ELAV proteins.

**The minimal set of RRM proteins.** Our results with the three kinetoplastids had revealed 12 RRM proteins which had clearly been conserved from kinetoplastids to yeast or mammals (Table 2). These were two poly(A) binding proteins; splicing factors U1 70k, U2AF35, and U2AF65; rRNA-processing protein MRD1; peptidyl-prolyl *cis-trans* isomerase; hnRNPH; translation factor SGN1; nuclear cap binding protein; and translation factors EIF3B and EIF3D. This prompted us to ask if these proteins comprised a minimal set required for eukaryotic gene expression. To answer this, we compared the trypanosome gene set with that of *Encephalitozoon cuniculi*, a microsporidian parasite with an obligately intracellular lifestyle and a minimal genome (26). *E. cuniculi* has a quarter of the number of proteins in the other unicellular eukaryotes considered, including *T. brucei*, and only 18 RRM domain proteins. We compared the sequences with those of several other unicellular eukaryotes—*S. cerevisiae*, *Schizosaccharomyces pombe*, *Entamoeba histolytica*, *Plasmodium falciparum*, and *T. brucei*—using GeneDB to screen the individual genomes and applying an E-value cutoff of  $10^{-15}$  in order to exclude matches based only on the RRM (which generally gave values of

about  $10^{-10}$ ). This revealed only three RRM proteins that were present in all organisms tested: the nuclear cap binding protein (Q8SW46\_ENCCU), poly(A) binding protein (Q8SR30\_ENCCU), and Mrd1p (Q8SRD9\_ENCCU). All of the organismal genomes tested, except that of *T. brucei*, also encoded a homologue of *S. cerevisiae* Mot2p (Q8SQS6\_ENCCU for *E. cuniculi*), and Q8SSA1\_ENCCU has a serine-rich region and aligns with an rRNA-processing protein present in the two yeasts and *Entamoeba*. *E. cuniculi* is probably a reduced derivative of a fungal ancestor, and *P. falciparum* contains a plastid of algal origin; consistent with this, a few of the other *E. cuniculi* RRM proteins showed their best matches to *S. pombe*, *S. cerevisiae*, and *P. falciparum*, with E values between  $10^{-10}$  and  $10^{-15}$ . The annotated database does not contain *E. cuniculi* RRM-containing translation elongation factors EIF3B and -D: it may be that the genes have diverged beyond recognition. Although the genome of *E. cuniculi* is annotated to include 13 mRNAs with short introns (26), the putative U2AF35 (Q8SQL5\_ENCCU) lacks classical RRM and the large subunit (Q8SRX8\_ENCCU) has a C-terminal RNA recognition motif that is too diverged to be recognized by Pfam. The *E. cuniculi* peptidyl-prolyl *cis-trans* isomerase is of a prokaryotic type, with no RRM. None of the *E. cuniculi* RRM proteins had zinc fingers or glutamine-rich domains, and no clear SR domains were present.

From these results, it would seem that the minimal set of conserved RRM-containing proteins at the primary sequence level consists of poly(A) binding protein, MRD1, and the nuclear cap binding protein. Even in *E. cuniculi*, over half of the RRM proteins appeared to be organism specific.

## DISCUSSION

Of the 75 *T. brucei* RRM proteins, only 18 can be assigned a function with any confidence. There are two poly(A) binding proteins, U2AF35, MRD1, PPCTI, hnRNPH, SGN1, CBC20, U1 70k (RBP24), and EIF3B, and also possible homologues of EIF3D, U2AF65, and hnRNPA. Experimental functional evidence is available concerning six proteins that are specific to trypanosomatids: NRBD1 and NRBD2, TRRM1, TSR1, and UBP1 and UBP2. This leaves over 50 proteins with no known function. The additional domains, such as zinc fingers, glutamine-rich regions, and SR-rich regions, have diverse functions but can be of only limited utility in predictions; however, it is tempting to speculate that the SR domain proteins might be involved in the regulation of *cis* or *trans* splicing or the selection of alternative splice sites. The comparison with *E. cuniculi* suggested that the presence of organism-specific RRM proteins may be ubiquitous, which implies that the RRM proteins have coevolved with their RNA targets. RNA target evolution will occur most readily in molecules that are relatively free of structural or sequence constraints. Since the kinetoplastids have only two or three *cis*-spliced genes, and regulation of *trans* splicing has not been documented, we hypothesize that many of the kinetoplastid-specific RRM proteins will bind untranslated regions of mRNAs and will be involved in regulating mRNA degradation and translation.

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