A 100-kD complex of two RNA-binding proteins from mitochondria of *Leishmania tarentolae* catalyzes RNA annealing and interacts with several RNA editing components

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ABSTRACT

A stable 100-kD complex from mitochondria of *Leishmania tarentolae* containing two RNA-binding proteins, Ltp26 and Ltp28, was identified by cross-linking to unpaired 4-thiouridine nucleotides in a partially duplex RNA substrate. The genes were cloned and expressed and the complex was reconstituted from recombinant proteins in the absence of RNA or additional factors. The Ltp26 and Ltp28 proteins are homologs of gBP27 and gBP29 from *Crithidia fasciculata* and gBP25 and gBP21 from *Trypanosoma brucei*, respectively. The purified Ltp26/Ltp28 complex, the individual recombinant proteins, and the reconstituted complex are each capable of catalyzing the annealing of complementary RNAs, as was previously shown for gBP21 from *T. brucei*. A high-molecular-weight RNP complex consisting of the Ltp26/Ltp28 complex and several 55–60-kD proteins together with guide RNA could be purified from mitochondrial extract of *L. tarentolae* transfected with Ltp28-TAP. This complex also interacted in a less stable manner with the RNA ligase-containing L-complex and with the 3′ TUTase. The Ltp26/Ltp28 RNP complex is a candidate for catalyzing the annealing of guide RNA and pre-edited mRNA in the initial step of RNA editing.

Keywords: RNA editing; RNA annealing; RNA binding proteins; trypanosomes

INTRODUCTION

The “enzyme cascade” model (Blum et al. 1990) of U-insertion/deletion RNA editing that occurs in mitochondria of kinetoplastid protozoa predicts the involvement of at least four enzymatic activities (endoribonuclease, 3′ TUTase, RNA ligase, 3′-5′ exonuclease) and most likely several structural or RNA-binding proteins within large ribonucleoprotein complexes (Pollard et al. 1992; Peris et al. 1994; Corell et al. 1996). The only enzymatic components isolated to date and shown to be involved in editing by gene knockout or RNAi are the larger RNA ligase (REL1; Schnaufer et al. 2001), a putative RNA helicase which may be required for the 3′ to 5′ processivity of editing or for gRNA-mRNA hybrid unwinding (Missel et al. 1997), and a 3′ terminal uridylyltransferase (TUTase; Aphasizhev et al. 2002). TbMP81 (Drozdz et al. 2002) and band III protein (Huang et al. 2002) are likely to be structural components of core RNA editing complexes in *Trypanosoma brucei*, as also shown by gene knockout studies. A smaller RNA ligase (REL2) and several unidentified proteins have also been isolated as components of a large “editing complex” (Panigrahi et al. 2001). A gRNA-directed mitochondrial endoribonuclease has been partially purified from *T. brucei* (Seiwert et al. 1996; Rusche et al. 1997) and a 3′-5′ U-specific mitochondrial exonuclease has been partially purified from *Leishmania tarentolae* (Aphasizhev and Simpson 2001). The REAP-1 mitochondrial protein from *T. brucei* was isolated as a component of high-molecular-weight complexes, and an inhibition of in vitro editing by anti-REAP-1 antibodies was reported (Madison-Antenucci et al. 1998), but the specific role of this protein remains to be identified.

One approach for the detection of components of the editing machinery has been to search for gRNA-binding or
mRNA-binding proteins. This approach has had varied success. For example, in *L. tarentolae*, three proteins, p18, p51, and p110, were identified by RNA binding and then discovered to be the mitochondrial metabolic enzymes ATPase subunit b, aldehyde dehydrogenase, and glutamate dehydrogenase, respectively (Bringaud et al. 1995, 1997), and a double knockout of the glutamate dehydrogenase gene had no effect on editing (Estévez et al. 1999). TBRG1, a mitochondrial protein from *T. brucei* that binds poly(U) and gRNAs, was cloned by homology to nucleolin and shown to cosediment with in vitro editing activity, but no role in editing has been ascribed to this protein (Leegwater et al. 1995; Vanhamme et al. 1998). RBP16 from *T. brucei* which binds in vitro to the gRNA 3' oligo[U] tail contains a cold-shock domain, but its role is uncertain (Hayman and Read 1999; Pelletier et al. 2000).

The RNA-binding mitochondrial protein gBP21, from *T. brucei*, has been extensively studied (Köller et al. 1994; Allen et al. 1998; Lambert et al. 1999; Muller et al. 2001) and has several properties that make it a very likely candidate for indirect involvement in editing. These involve high-affinity binding to gRNA and coimmunoprecipitation with in vitro editing activity. An RNA annealing activity was also demonstrated, and it was proposed that this protein played a role in the initial interaction of mRNA and the cognate gRNA (Muller et al. 2001). However, a targeted knockout of the corresponding gene had no apparent affect on editing in vivo (Lambert et al. 1999). In *Crithidia fasciculata*, gBP29, an ortholog of gBP21, and a second related RNA-binding protein, gBP27, were cloned. In addition, an ortholog of gBP27 from *T. brucei*, gBP23, was identified (Leegwater et al. 1995; Blom et al. 2001). These proteins from *C. fasciculata* coimmunoprecipitated and were present in the form of variable-sized high-molecular-weight complexes in the mitochondrial extract.

Here we present the isolation and characterization of a stable heterotetrameric 100-kD complex of two RNA-binding mitochondrial proteins, Ltp28 and Ltp26 from *L. tarentolae*. The Ltp28 protein is an ortholog of the *T. brucei* gBP21, and the Ltp26 protein is an ortholog of the *T. brucei* gBP25. Both the complex and each recombinant protein catalyze the annealing of RNA. The complex is present in the extract as a high-molecular-weight RNP complex, which interacts with the two mitochondrial RNA ligases and the 3' TUTase.

**RESULTS**

Crosslinking of protein to unpaired 4-thiouridine nucleotides in a model substrate RNA

As shown in 1986 (Favre et al.), 4-thiouridine forms efficient and stable crosslinks when RNA and protein molecules are positioned in close proximity and the crosslinking is activated by long-wavelength UV. RNA-I, shown in Figure 1A, was radioactively labeled and 4-thioU residues site-specifically inserted during T7 transcription by incorporation of a mixture of [α-32P]UTP and 4-thiouridine. The partially complementary RNA-II which hybridized downstream and upstream of the 4-thio Us was also produced by T7 transcription and subsequent annealing of the two RNAs, yielding the substrate RNA shown in Figure 1A, which emulates an mRNA-gRNA substrate with four unpaired U (and thio-U) residues at the initial U-deletionsite. It should be noted that we have not tested this substrate for activity in an RT-PCR editing assay (Byrne et al. 1996) or a precleaved editing assay (Igo et al. 2000).

The labeled RNA substrate was incubated with mitochondrial extract which had been preincubated with 100-fold mass excess of nonspecific RNA, and then subjected to 300-nm UV irradiation. One major crosslink was observed on an SDS gel (Fig. 1B). The formation of crosslink I was dependent on the an-
nealing of RNA-II and was inhibited more than 90% by a twofold mass excess of heparin or a 3–5-fold excess of a poly[U] homopolymer. Poly[C] only achieved this level of inhibition at 10-fold excess, and poly[A] and nonspecific RNA did not inhibit the formation of the crosslink until more than a 500-fold excess (not shown).

**Isolation of Ltp26 and Ltp28 proteins and cloning of genes**

Crosslink I was preparatively gel-isolated and digested with RNase, and the labeled proteins separated in an SDS gel (data not shown). Label transfer after RNase digestion was observed mostly to the upper 28 kD band (data not shown).

Based on this information, a purification method was developed using the generation of crosslink I as an analytical assay. As described in Materials and Methods, a fractionation method employing a 40%–55% ammonium sulfate cut of the mitochondrial extract, followed by heparin affinity chromatography, and finally by size fractionation in a Superose 12 column led to the isolation of a fraction that was active in the generation of crosslink I. A single 100-kD protein peak was seen in the final Superose 12 gel fractionation (see Fig. 5 below). Interestingly, an enrichment for two major bands of 26 and 28 kD and two minor bands of 21 and 23 kD occurred during this fractionation, especially after the heparin affinity chromatography (Fig. 1C).

Both major bands were subjected to N-terminal and internal protein microsequencing, and N-termini were also determined for the minor 21- and 23-kD bands. The Ltp28 protein was blocked at the N-terminus, but the Ltp26 protein did yield an N-terminal sequence, which was used to establish the site of cleavage of the mitochondrial signal sequence (see below). Degenerate oligonucleotides based on the peptide sequences were used to amplify genomic fragments, which were employed as probes to clone the genes from a subgenomic library. The genes were expressed in *E. coli*, and the recombinant proteins were used to generate polyclonal antisera.

The amino acid sequences of the Ltp28 and Ltp26 genes (Fig. 2, peptides indicated by lines) contained the peptides obtained from the microsequencing of p23 and p21, respectively, confirming that these represented proteolytic fragments. An N-terminal mitochondrial targeting sequence in the Ltp28 gene was recognized by the SignalP and Mitoprot programs with high confidence levels. The cleavage site of the N-terminal targeting sequence in Ltp28 was determined by the N-terminal amino acid sequence of the mature protein (Fig. 2A, arrow). The 46-aa signal peptide in Ltp26 probably accounts for the observed difference in the electrophoretic mobility of Ltp28 and Ltp26, given that the calculated molecular weight of the Ltp28 preprotein is only 0.25 kD larger that Ltp26.

A BLAST search revealed a strong homology of Ltp28 with *T. brucei* gBP21 (Koller et al. 1997) and *C. fasciculata* gBP29 (Blom et al. 2001). The *L. tarentolae* Ltp26 sequence showed strong homology with *C. fasciculata* gBP27 (Blom et al. 2001). The *L. tarentolae* Ltp26 protein also produced a significant hit with an EST clone (A1881040) in the *T. brucei* Genome Database. This sequence was PCR-amplified and used to clone the ortholog from *T. brucei*, which proved to be gBP25, a gene previously cloned from *T. brucei* (Blom et al. 2001).

An alignment of these RNA-binding proteins from *T. brucei*, *C. fasciculata*, and the Ltp26 and Ltp28 proteins from *L. tarentolae* is shown in Figure 2A,B. This analysis extends the findings of Blom et al. (2001) with sequences from three organisms and indicates several highly conserved motifs in each ortholog group. A limited conservation of three motifs can be seen in the alignments of all six sequences in Figure 2C. No functional motifs were recognized by Frosite or Profile analysis, including any known RNA-binding motifs.

**Mitochondrial localization of the Ltp26/Ltp28 complex**

In situ immunofluorescence analysis using the antisera against the purified 100-kD complex, which recognizes both proteins, showed that the proteins are confined to the circular mitochondrion in log-phase cells (Fig. 3A–C) and the asymmetric mitochondrion in stationary-phase cells (Fig. 3D–F; Simpson and Kretzer 1997). Interestingly, there was a predominant localization in the kinetoplast region (K) of the mitochondrion in addition to a diffuse localization throughout the tubular mitochondrial structure (Fig. 3C,F).

**Ltp26 and Ltp28 are components of a stable 100-kD complex**

As mentioned above and shown in Figure 5A below, the purification procedure yielded a single peak in a Superose 12 fractionation with an apparent molecular weight of 100 kD. This peak contained both Ltp26 and Ltp28 in approximately equimolar ratio. This purified complex was used to generate polyclonal antisera, which showed reactivity against both Ltp26 and Ltp28 with approximately equal affinities. The equal intensity of the Ltp26 and Ltp28 bands in gels stained with Coomassie Blue (Fig. 1C, lane 4) and Sypro Ruby (see Fig. 5B below) suggested that the proteins are present at a 1:1 ratio. The yields of tryptic peptides from the two bands for protein sequencing were also close to this ratio (Harvard Microchemical Facility, pers. comm.). The 1:1 ratio and the 100-kD molecular weight of the complex argue for an α2β2 organization.

To investigate the topography of protein–protein interactions within the Ltp26/Ltp28 complex, chemical cross-linking was performed, a fraction of which had undergone an additional purification step on Phenyl Superose (see Materials and Methods). Incubation of the complex with ho-
mobifunctional crosslinking reagents with different chain lengths produced three closely migrating crosslinks with molecular weights that are commensurate with all pairwise combinations, Ltp26-Ltp28, Ltp26-Ltp26, and Ltp28-Ltp28 (Fig. 4A). The low efficiency and the absence of a crosslink which would correspond to three or four subunits together may be due to the low lysine content in these proteins (four in Ltp26 and seven in Ltp28). No endogenous RNA component could be detected by phenol-extracting 50 µg of the purified Ltp26/Ltp28 complex, ethanol-precipitating, and incubating the pellet with RNA ligase and [α²P]PCP or with poly[A] polymerase and [α²P]ATP (not shown). This is not surprising however, because the complex was subjected to heparin affinity chromatography and high salt conditions during the isolation.

**Yeast two-hybrid and coimmunoprecipitation evidence of an interaction of Ltp26 and Ltp28**

To assess the relative affinity of protein–protein interactions within the Ltp26/Ltp28 complex, yeast two-hybrid analysis (Chien et al. 1991) was performed by cotransfecting a yeast strain deficient for several nutritional markers with Ltp28 and Ltp26 fusion constructs followed by selection on the appropriate media (see Materials and Methods). The survival of the cells on selective media depended on the pres-
ence of both Ltp26 fused with the gal4 binding domain (= bait) and Ltp28 fused with the gal4 activation domain (= prey), or vice-versa (Fig. 4B). No interaction of Ltp26 with Ltp26 or Ltp28 with Ltp28 could be detected under the same stringency conditions. In addition, cotransformants of yeast with a fusion construct of the \textit{L. tarentolae} 3/H11032 TUTase and a fusion construct of Ltp26 or Ltp28 did not grow on selective medium, possibly suggesting a lack of direct interaction between Ltp26 or Ltp28 and TUTase, although other explanations are of course possible for this negative result in a heterologous system.

Interactions were also assayed by coimmunoprecipitation. S\textsuperscript{35}-labeled HA-tagged or Myc-tagged proteins were synthesized in vitro transcription-translation and then mixed, and physical interactions were assayed by coimmunoprecipitation using an anti-Myc monoclonal antibody (Fig. 4C). The anti-Myc antibody pulled down Myc-tagged S\textsuperscript{35}-labeled Ltp28 (lane 3) but only pulled down HA-tagged S\textsuperscript{35}-labeled Ltp26 in the presence of unlabeled Myc-tagged Ltp28 (lanes 1 and 2), thus confirming the in vivo yeast two-hybrid results. In addition, this procedure could not demonstrate an association between TUTase and Ltp26 (lanes 4 and 5), which is consistent with the negative yeast two-hybrid results (Fig. 4B). The reciprocal pull down of Ltp28 by Ltp26 was not technically possible as in vitro-synthesized Ltp28 (unlike TUTase or Ltp26) partially precipitated out of solution during the immunoprecipitation. Together these data provide evidence of a strong interaction of Ltp26 and Ltp28.

**In vitro reconstitution of Ltp26/Ltp28 complex with recombinant Ltp26 and Ltp28 proteins**

The Ltp26 and Ltp28 proteins were expressed in \textit{E. coli} with His-tags at the N-termini of the mature proteins. The Ltp26 protein was purified under native conditions by metal and heparin affinity chromatography to more than 95% purity, as estimated from Sypro Ruby staining of an SDS gel. The Ltp28 protein formed inclusion bodies in \textit{E. coli} and was therefore purified by metal affinity chromatography under denaturing conditions; the denatured Ltp28 protein was then refolded while attached to the matrix and eluted (see Materials and Methods). Subsequent heparin column purification of Ltp28 resulted in a fully soluble preparation of more than 95% purity, as estimated by SDS gel analysis.

The reconstitution of high-molecular-weight complexes by the recombinant proteins after incubation for 16 h at 2 mg/mL protein concentration was monitored by Superose 12 chromatography (Fig. 5A). As mentioned above, the Ltp26/Ltp28 complex purified from the mitochondrial extract migrated as a sharp 100-kD peak. Recombinant Ltp28 migrated as an apparent monomer, which is consistent with previously described properties of the \textit{T. brucei} ortholog, gBP21 (Koller et al. 1997). Recombinant Ltp26 migrated as an apparent homodimer, with a small shoulder migrating at a possible homotrimer position. The interaction of Ltp26 with itself was not detected in the above yeast two-hybrid assay, suggesting that this interaction may be less stable than the Ltp26/Ltp28 interaction. Incubation of Ltp28 and a two-fold molar excess of Ltp26 yielded a substantial peak comigrating with the purified 100-kD Ltp26/Ltp28 complex in addition to a major dimer peak and a minor trimer peak. The molar excess of Ltp26 was necessary to achieve incorporation of Ltp28 into the tetrameric complex (data not shown). Figure 5B shows the SDS gel profiles of the proteins from chromatographic fractions 4–14. The stoichiometric ratio of Ltp28:Ltp26 in the reconstituted tetramer peak in fractions 5–8 was approximately 1:2. We attribute this deviation from a 1:1 ratio to cross contamination of the tetramer peak with the overlapping Ltp26 trimer peak. Consistent with this interpretation, the ratio of Ltp26 to Ltp28 in the leading edge of the tetramer peak (fraction 4) is close to 1:1.

**RNA-binding properties of the purified and reconstituted Ltp26/Ltp28 complex and the recombinant Ltp26 and Ltp28 proteins**

The equilibrium dissociation constants (K\textsubscript{d}) for binding single-stranded (denatured RNA-III, Fig. 7 below) and double-stranded RNA (annealed RNA-III+RNA-V) were determined for the purified native Ltp26/Ltp28 complex, the individual recombinant proteins, and the reconstructed Ltp26/Ltp28 complex. As shown in Table 1, binding to denatured RNA-III and to annealed RNA-III+RNA-V was ap-
approximately the same, with a possible slight preference of the native and reconstituted complexes and the Ltp28 for the double-stranded RNA. However, the Kd values were significantly lower for both the native and the reconstructed complexes, indicating a higher affinity of binding of the complex than either of the component proteins. The Kd for the binding of Ltp28 with single-stranded RNA was approximately 10-fold higher than that reported for the binding of *T. brucei* gBP21 to single-stranded RNA (Muller et al. 2001), which may reflect a species variation or differences in the recombinant protein preparation. However, the Kd of binding of the Ltp26/Ltp28 complex to single-stranded RNA (9.6 nM) is nearly identical to that of gBP21 (Muller et al. 2001).

The interaction of the purified native Ltp26/Ltp28 complex with in vitro transcribed gRPS12-I gRNA (see Materials and Methods) was assayed by gel filtration chromatography. At a 10-fold excess of protein, the formation of an ~700-kD complex was observed, which corresponds to approximately 5–6 100-kD complexes bound to one RNA molecule (Fig. 6A). A 10-fold molar excess of RNA resulted in the formation of an ~200-kD complex, which may correspond to a single complex bound per RNA molecule. The 100-kD complex alone and the labeled RNA alone are also shown. It should be noted that the use of the T7 transcribed gRNA as substrate in this assay does not imply any specificity of binding for gRNA, but was used since it represents a natural ligand for the complex (see below).

Native gel analysis of increasing amounts of Ltp26/Ltp28 complex mixed with 20 nM labeled gRNA showed a series of discrete higher-molecular-weight protein/RNA complexes (Fig. 6B). This method, however, does not allow determination of accurate molecular weights due to the high isoelectric point (pH 10.1) of the Ltp26/Ltp28 complex, because in the absence of RNA these proteins do not enter the native gel at pH 8.4. The two RNA bands of unequal intensity in the “no protein” control lane probably represent folded and unfolded RNA molecules.

The Ltp26/Ltp28 complex and the recombinant Ltp26 and Ltp28 proteins catalyze the annealing of complementary RNAs

A T1 nuclease protection assay (Portman and Dreyfuss 1994) was used to determine whether the Ltp26/Ltp28 complex catalyzes the annealing of complementary RNAs as has been previously shown for the *T. brucei* gBP21 homolog of the Ltp28 protein (Muller et al. 2001). The substrate consisted of two partially complementary RNAs (RNA-III and RNA-IV in Fig. 7). RNA-IV was uniformly labeled during transcription. T1 nuclease cleaves 3′ of G residues only in single-stranded RNA. As shown in Figure 7, the T1 nuclease-
ase-protected fragment of labeled RNA-IV which is annealed to RNA-III is eight nucleotides shorter than the input RNA.

The increase in rate of annealing in the presence of the Ltp26/Ltp28 complex is shown by comparing the protected fragments produced in the presence of added protein to those produced in the absence of protein (Fig. 7, cf. panels 2–4 to panel 1, lane 3). The annealing of labeled RNA-IV and the partially complementary RNA-III under these conditions in the absence of Ltp26/Ltp28 complex represented approximately 5% (panel 1, lane 3) of that achieved by incubation in the presence of Ltp26/Ltp28 protein or by heating and slow cooling with a 50-fold excess of RNA-IV (panel 1, lane 1). A control experiment showed that annealing of the labeled RNA-IV to the non-complementary RNA-VI did not produce a T1-protected fragment (Fig. 7, panel 1, lane 4).

Nucleotides were not required for the catalysis of RNA annealing by the Ltp26/Ltp28 complex (Fig. 7, panels 2–4), but omitting MgCl₂ greatly reduced the efficiency of annealing (panel 5).

Similar results were obtained with the RNA constructs which were used for the crosslinking experiments in Figure 1, suggesting a lack of RNA sequence specificity of the RNA annealing activity (data not shown).

The second-order association rate constants ($k_2$) for RNA annealing in the absence and presence of the proteins were obtained as described in Materials and Methods. The stimulation of annealing, which was defined as the ratio of $k_2$ (with protein)/$k_2$ (no protein) where $k_2$ in the absence of protein = $1.6 \times 10^4$ M⁻¹ s⁻¹, varied from 20- to almost 70-fold. The recombinant Ltp28 protein showed a threefold higher stimulation of annealing compared to the Ltp26 protein and showed a higher value than the purified Ltp26/Ltp28 complex (Table 1). Recombinant *T. brucei* gBP25 protein showed annealing activity similar to that of Ltp26 (Table 1).

**Interaction of the Ltp26/Ltp28 complex with components of editing machinery in mitochondrial extract**

Affinity-purified polyclonal antibodies raised against the purified Ltp26/Ltp28 complex and the recombinant proteins were used for immunoprecipitation (IP) of mitochondrial extract. The anti-Ltp26 antibody coprecipitated Ltp26 and Ltp28 and the anti-Ltp28 antiserum did the same, confirming that the complex is composed of both proteins (Fig. 8A, panel 1). The specificity of these antisera and the absence of crossreactivity between Ltp26 and Ltp28 was con-

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**TABLE 1.** RNA binding and RNA annealing properties of the Ltp26/Ltp28 100 kDa complex and individual recombinant proteins

<table>
<thead>
<tr>
<th>Protein</th>
<th>Single-stranded RNA Kd (nM)</th>
<th>Double-stranded RNA Kd (nM)</th>
<th>Fold annealing stimulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Native complex</td>
<td>9.6 ± 2</td>
<td>2.6 ± 2</td>
<td>38</td>
</tr>
<tr>
<td>Ltp28</td>
<td>128 ± 21</td>
<td>96 ± 6</td>
<td>67</td>
</tr>
<tr>
<td>Ltp26</td>
<td>225 ± 52</td>
<td>259 ± 65</td>
<td>21</td>
</tr>
<tr>
<td>Reconstituted complex</td>
<td>24 ± 2</td>
<td>4.5 ± 1.2</td>
<td>N/D</td>
</tr>
</tbody>
</table>

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**FIGURE 5.** Reconstitution of Ltp26/Ltp28 100-kDa complex. (A) Superose 12 gel filtration profiles of purified Ltp26/Ltp28 complex, of the material reassembled from the purified recombinant Ltp26 and Ltp28 proteins, and of the individual proteins. Positions of molecular weight markers are indicated. (B) Protein composition of native and reconstructed complexes. Fractions were collected from the same starting point in chromatographic runs (above) and separated on 10%–20% SDS gradient gels which were stained with Sypro Ruby.
firmed by Western blotting analysis of the mitochondrial extract (data not shown).

The anti-Ltp26 antiserum also coprecipitated a fraction of the TUTase from the extract (Fig. 8A, panel 3). A control for specificity of the IP was provided by using antibody against glutamatedehydrogenase (GDH), an abundant mitochondrial matrix protein with a high affinity for RNA but which is apparently not involved in RNA editing (Estévez et al. 1999), and by showing that this protein was present in the extract but not in the anti-Ltp26 IP (Fig. 8A, panel 2). A reciprocal IP with anti-TUTase antibody confirmed these results (Fig. 8A, panel 4). These data suggest an interaction of the Ltp26/Ltp28 complex with TUTase or with a TUTase-containing complex. The latter is suggested by the experiments described in Figure 4C, in which recombinant TUTase was shown not to associate with Ltp26, Ltp28, or the Ltp26/Ltp28 complex in vitro, and by the yeast two-hybrid experiments in Figure 4B in which no direct interaction could be detected in vivo between TUTase and Ltp26 or Ltp28.

The LtREL1 and LtREL2 RNA ligase proteins can be detected by adenylation with [$\alpha$32P]ATP (Sabatini and Hajduk 1995; Peris et al. 1997). Evidence of an interaction of ATP-labeled LtREL1 and LtREL2 in mitochondrial extract with Ltp26 was provided by the co-IP with anti-Ltp26 antiserum shown in Figure 8B, right panel. The co-IP of LtREL1 and LtREL2 (and TUTase) was stable enough to withstand high-stringency washes with 0.5 M NaCl and 0.5% Triton X100 (not shown).

We also investigated the possibility that the Ltp26/Ltp28 complex interacts with endogenous gRNA. Guide RNA in the mitochondrial extract can be specifically 5′ end-labeled with [$\alpha$32P]GTP and guanylyltransferase, as shown for total mitochondrial extract in Figure 8B (left panel, Ex lane). IP of the extract with anti-Ltp26 antiserum was performed and RNA extracted from the washed beads and labeled with GTP. A substantial amount of the gRNA in the extract co-immunoprecipitated with Ltp26 and Ltp28 (Fig. 8B, left panel, IP lane). A control for the specificity of the co-IP of gRNA was shown by the lack of co-IP of the abundant mitochondrial RNAlys by the anti-Ltp26 antiserum (Fig. 8B, lower panel).

To investigate these interactions in more detail, mitochondrial extract was fractionated in a glycerol gradient and the distribution of the Ltp26/Ltp28 complex assayed. The purified Ltp26/Ltp28 complex sedimented as expected in one band, peaking in fractions 4–5 (Fig. 8C, panel 1). However, the Ltp26/Ltp28 complex in the extract showed, in addition to a peak comigrating with the purified complex, another peak sedimenting in approximately the 30S region, suggesting an association of the Ltp26/Ltp28 complex in the extract with RNA and/or protein components. In agreement with this distribution, RNA annealing activity was spread throughout the gradient, with the majority in the ~10–15S region (Fig. 8C, panel 3). The RNA annealing activity in the 10–15S region appears to be higher than the ~30S region, although a similar amount of Ltp26/Ltp28 complex is present, which may indicate the saturation of higher-molecular-weight forms of the complex with substrate RNA. Each gradient fraction was subjected to IP with anti-Ltp26 antiserum and analyzed for co-IP of TUTase and LtREL1 and LtREL2. Co-IP of TUTase (Fig. 8C, panel 4) occurred throughout the gradient, with the relative amount precipitating in different regions, coinciding with the previously reported distribution of TUTase in glycerol gradients (Peris et al. 1997; Aphasizhev et al. 2002). Co-IP of the two RNA ligases was mainly limited to the higher-molecular-weight region of the gradient, coinciding with the second peak of Ltp26/Ltp28 (Fig. 8C, panel 5, indicated by line above fractions 9–13).

### Isolation of Ltp26/Ltp28-associated complex

In order to more directly assess the Ltp26/Ltp28 interactions, Ltp28 protein with the mitochondrial targeting sequence was ectopically expressed in *L. tarentolae* from the pX vector (LeBowitz et al. 1990) with a C-terminal fusion of a dual TAP epitope tag, which consists of an IgG-binding Protein A domain and a calmodulin-binding domain (Puig et al. 2001). Mitochondria were isolated from transfected cells, and clarified mitochondrial extract was sedimented...
through a glycerol gradient. Each gradient fraction was subjected to affinity binding with IgG Sepharose. As shown in Figure 9A, both the wild-type Ltp28 protein and the Ltp28-TAP protein were present in the IgG “pull down,” indicating that the tagged protein was incorporated into the Ltp26/Ltp28 complex. In addition, as also found above in the co-IP from mitochondrial extract, the IgG “pull down” from the higher-molecular-weight gradient fractions (indicated by line above fractions) contained TUTase and LtREL1 and LtREL2 (Fig. 9A). The identical protein profile was obtained when Ltp26 was used as an entry for the TAP purification system (not shown). GTP-labeled gRNA also sedimented in this region of the gradient (Fig. 9D, lower panel), in agreement with the co-IP results described above.

Adenylation of each fraction and SDS (Fig. 9B) and native gel analysis (Fig. 9D, upper panel) showed the presence of the LtREL1 and LtREL2 RNA ligases sedimenting in the region of the gradient of the purified IgG + calmodulin “pull down” material. The heterogeneous appearance of the adenylated complexes in the native gel is probably a result of breakdown during the electrophoresis and/or to the presence of variable amounts of bound RNA.

Treatment of the mitochondrial extract with RNase A during binding to the IgG matrix led to a disappearance of the high-molecular-weight RNP complex, as shown in Figure 9B for LtREL1 and LtREL2 RNA ligases sedimenting in the region of the gradient of the purified IgG + calmodulin “pull down” material. The heterogeneous appearance of the adenylated complexes in the native gel is probably a result of breakdown during the electrophoresis and/or to the presence of variable amounts of bound RNA.

We have confirmed and extended previous results from *T. brucei* (Koller et al. 1997; Allen et al. 1998; Lambert et al. 1999; Blom et al. 2001; Muller et al. 2001) and *C. fasciculata* (Blom et al. 2001) for the presence of two small mitochon-
Association of Ltp26/Ltp28 complex with components of RNA editing machinery. (A) Co-IP of Ltp26/Ltp28 complex and 3'TUTase. IP from mitochondrial extract was performed using Sepharose G beads coated with affinity-purified mouse polyclonal antibodies against recombinant Ltp26, recombinant Ltp28, and L. tarentolae 3’ TUTase (Ltp26-IP, Ltp28-IP, and TUT-IP lanes). Immunodetection was performed with rabbit antibodies raised against purified Ltp26/Ltp28 complex, recombinant glutamate dehydrogenase (GDH) (Bringaud et al. 1997), and TUTase. Ext, mitochondrial extract; Pre, proteins bound to G-beads coated with preimmune serum; Ab, proteins bound to G-beads coated with specific antibodies. 10%–20% SDS gels blotted to nitrocellulose membranes were incubated with [32P]GTP in the presence of guanylyltransferase and separated on a 12% acrylamide/urea gel. Lower panel: The presence of tRNA<sup>19</sup> was assayed in the same material by Northern blotting with an oligonucleotide probe. Right panel: The same samples were incubated with [32P]ATP to visualize the LtREL1 and LtREL2 proteins and analyzed on a 10%–20% SDS gel. (B) Association of Ltp26/Ltp28 complex with TUTase and with RNA ligase. IP was performed with preimmune (Pre) or anti-Ltp26 (IP) antibody. Left panel: Total RNA was isolated from mitochondrial extract and the IP material, labeled with [α-32P]GTP. Panel 2, Western analysis of Ltp26 and Ltp28. Panel 3, T1 nuclease protection assay for RNA annealing activity. Panels 4 and 5, co-IP of TUTase and LtREL1 and LtREL2 by anti-Ltp26 antibody.

It was shown previously that the T. brucei gPB21 showed RNA-binding properties and also catalyzed RNA annealing, although only a minority of the annealing activity of the mitochondrial extract could be attributed to this protein (Muller et al. 2001). We confirmed an annealing activity for the gPB21 homolog from L. tarentolae (Ltp28) and showed that Ltp26 and the T. brucei gPB25 homolog of Ltp26 also had RNA annealing activities, although their K<sub>d</sub> values (Table 1) indicate a lower affinity for both single- and double-stranded RNAs. Furthermore, both the purified and reconstituted Ltp26/Ltp28 complex showed higher-affinity RNA binding than either of the component proteins and stimulated RNA annealing approximately 40-fold. The
binding of purified Ltp26/Ltp28 100-kD complex to a synthetic single-stranded RNA was shown to occur stepwise essentially to saturation, with one protein complex for approximately 5–7 nucleotide bases. Although an RNA component was not detected in the purified Ltp26/Ltp28 100-kD complex, the co-IP analysis and the TAP pull-down analysis indicated an interaction of the Ltp26/Ltp28 complex with gRNA and with several 55–60-kD proteins to form a stable high-molecular-weight RNP complex.

T. brucei gPB21 has been shown to coimmunoprecipitate with in vitro RNA editing activity, ATP-labeled RNA ligase proteins, and TUTase activity, and the interaction was sensitive to micrococcal nuclease treatment, suggesting involvement of an RNA component (Allen et al. 1998). The C. fasciculata gPB27 and gPB29 proteins, however, were reported not to interact with any identified editing components or RNA (Blom et al. 2001). We showed by TAP affinity chromatography of tagged Ltp28 targeted to the mi-
tochondrion and incorporated into the 100-kD complex that the 100-kD Ltp26/Ltp28 complex interacts via an RNA component with both the 3' TUTase and the RNA ligase-containing L-complex to form a heterogeneous high-molecular-weight RNP complex sedimenting at approximately 30S in glycerol gradients. We recently found evidence for the existence of an ∼20–25S RNP complex that shows in vitro editing activity and consists of the L-complex interacting via an RNA component with the 3' TUTase and with the 100-kD Ltp26/Ltp28 complex (unpubl. results). We speculate that the differences in sedimentation may be due to variable extents of interaction and/or bound RNA.

We speculate that the Ltp26/Ltp28 RNP complex plays a specific role in the initiation of RNA editing in which a gRNA forms a duplex anchor with the cognate preedited mRNA, thereby creating a substrate for an initial site-directed cleavage. However, as mentioned above, previous evidence for involvement of the T. brucei gBP21 protein in RNA editing is somewhat equivocal. gBP21 was reported to coprecipitate with a 20S in vitro editing activity (Allen et al. 1998), whereas gBP21 null mutants of bloodstream T. brucei showed an overall decrease in the abundance of mitochondrial transcripts but no effect on editing in vivo (Lambert et al. 1999). One explanation of these earlier results could be the observed redundancy of the two proteins of the Ltp26/Ltp28 complex. In this regard, we recently showed that inhibition of expression of T. brucei gBP21 by RNAi had no effect on viability, whereas inhibition of expression of both gBP21 and gBP25 produced a growth defect (A. Simpson, R. Aphasizhev, L. Simpson, unpubl.). Further work is required to determine the precise biological role of the Ltp26 and Ltp28 proteins.

MATERIALS AND METHODS

Cell growth and isolation of mitochondria

L. tarentolae cells were grown in 15-L batches in brain-heart infusion medium with a supplement of hemin to late log phase (∼180 × 10^6 cells/mL) in a BioFlow IV Fermentor (New Brunswick). Mitochondria were isolated by the hypotonic lysis Reno-
Gene cloning and protein expression

N-terminal sequences were obtained from the 26-kD and 28-kD bands and two minor bands of 20 kD and 22 kD. Two internal peptide sequences from the 26-kD band and three peptides from the 28-kD band were used to construct degenerate oligonucleotides with the Codehop program. All possible pairwise combinations of oligonucleotides were tested for PCR using genomic DNA. A 262-bp product was obtained for the 26-kD protein and a 378-bp product for the 28-kD protein. The PCR products were used as probes to clone 4-kb KpnI and 3.5-kb PstI genomic fragments that contained the entire genes for Ltp26 and Ltp28, respectively. The Ltp26 ortholog from T. brucei was cloned using a PCR-derived probe based on the EST sequence AI881040 (Parasite Genomes Database) from a KpnI subgenomic library. The 5.3-kb genomic clone contained the full-length gene. For expression in E. coli, cloned genes were PCR-amplified, with NdeI and BamHI sites introduced at the 5′ and 3′ ends, respectively, and cloned into the pET 15b vector, which provides an N-terminal 6 His affinity tag. The experimentally determined mitochondrial localization signal (position −22) for Ltp26 from T. tarentolae, and the predicted signals (position −22) for Ltp28 and (position −30) for the Ltp26 ortholog from T. brucei were removed. Both Ltp26 proteins were partially soluble (30%–50%) when expressed at 25°C and were purified by Talon metal affinity chromatography as suggested by the manufacturer, followed by chromatography on a 1-mL heparin HiTrap column as described above which was down-scaled fivefold. The Ltp28 formed inclusion bodies and was purified under denaturing conditions. The protein was refolded while still attached to the Talon resin at 20°C by gradually decreasing the urea concentration from 8 to 0 M in 20 mM sodium phosphate buffer pH 7.0, 300 mM NaCl. The gradient was run for 15 h at 1 column volume per h. The protein was eluted with 200 mM of imidazole and further purified on a heparin column. All proteins were at least 95% pure as determined by Sypro Ruby (Molecular Dyes) staining of SDS gels.

Protein–protein interactions

DNA, DMP, and DMS are homobifunctional imidoester crosslinking agents (Pierce) that react with primary amines and differ in spacer arm length (8.6, 9.2, and 11 A, respectively). Purified Ltp26/28 complex (0.1 mg/mL) in 50 mM sodium phosphate buffer, pH 8.0, 150 mM KCl, was incubated for 40 min at 5°C in the presence of 10 mM of the crosslinking reagent. The reaction was stopped by adding 1 M Tris-HCl, pH 6.8 to 1/10 of the volume, and the products were analyzed on a 10%–20% SDS gel followed by Sypro Ruby staining.

The full-length coding regions for the Ltp26 and Ltp28 L. tarentolae proteins, and the L. tarentolae 3′ TUTase were cloned in the proper reading frames into both the ‘bait’ (pGBK7T7) and ‘prey’ (pGADT7) vectors used in the Matchmaker 3 Yeast Two-Hybrid Screening Kit™ (Clontech). Each construct alone or in various pairwise combinations with other constructs were used to transform yeast strain AH109 (MATα, trp1–109, leu2–3, 112, ura3–5, his3–200, ade2–101, gal4Δ, gal80Δ, LYS2-GAL1-UAS-GAL1-TATA-HIS3, GAL1-UAS-GAL2-TATA-AD2, URA3-MEL1-UAS-MEL1-TATA-lacZ) using appropriate nutritional selection; the resulting yeast transformants were checked for fusion protein expression by Western analysis. In this manner, each of the three L. tarentolae proteins were shown to be expressed fused at their amino-terminal end to either the DNA-binding domain of the Gal4 protein epitope-tagged with Myc or the activation domain of the Gal4 protein epitope-tagged with HA. Each of the plasmid constructs was also used for coupled transcription/translation (Tnl7 T7 System, Promega). Because of the location of the T7 promoter in the parent pGBK7T7 and pGADT7 vectors, in vitro translation products retain the amino-terminal epitope tags, Myc and HA, respectively, but do not contain any portion of the Gal 4 protein. Pairwise mixtures of these were analyzed for association in vitro using a co-IP protocol (Matchmaker Co-IP Kit) that included clarification of reactions to remove any nonspecific precipitation and/or nonspecific adsorption.

Reconstruction of Ltp26/Ltp28 complex

Purified recombinant proteins were mixed at a 1:2 (Ltp28:Ltp26) molar ratio and a final protein concentration of 2 mg/mL, and dialyzed against 20 mM sodium phosphate, 300 mM NaCl, for 16 h at 25°C. We found that an excess of the Ltp26 protein was necessary for in vitro complex formation. After 10 min centrifugation at 200,000 g, the samples were loaded on a Superose 12 column and chromatographed in the same buffer at 0.3 mL/min. Fractions of 0.4 mL were collected and analyzed on a 10%–20% acrylamide-SDS gel and stained with Sypro Ruby. Molecular weight markers were bovine gamma globulin, ovalbumin, myoglobin, and Vitamin B12 (Biorad). For gRNA-protein reconstruction, purified Ltp26/28 complex and gRNA were incubated with in vitro transcribed RPS12-I gRNA (see Results) and separated on Superose 6 columns in 20 mM HEPES, pH 7.5, 5 mM MgCl2, 150 mM KCl buffer. Molecular weight markers were thyroglobulin, ferritin, catalase, and BSA. Native precast 8%–16% gels (Invitrogen) were run for 12 h at 40 V, fixed, dried, and exposed to X-ray film. Premixed high-molecular-weight markers for native gel electrophoresis were from AP Biotech.

Catalysis of RNA annealing and RNA binding

RNA molecules (RNA-III and RNA-IV in Fig. 7) were synthesized by run-off transcription from a linear DNA template (Kapushch et al. 1999) using T7 RNA polymerase. RNA-IV was labeled with [α-32P]UTP during transcription and purified on an 8M urea- 5% polyacrylamide gel. The RNA annealing reaction contained 0.5 nM of mRNA and 0.5-50 nM of gRNA, and the protein concentration varied from 0 to 5 μM. All reactions were performed in 20 μL of 20 mM HEPES, pH 7.6, 100 mM NaCl, 1 mM MgCl2, 1 mM DTT for 0.5, 1, 2, 4, and 10 min at 27°C. The reaction was stopped by the addition of 1 μL T1 RNase (18u/μL) and incubated for 10 min at 27°C, followed by phenol/chloroform extraction and ethanol precipitation. Products were analyzed in 15% polyacrylamide/urea gels, exposed to PhosphoImager plates, and quantified with ImageQuant software.

The first- and second-order association constants were calculated as described (Muller et al. 2001). The affinity of RNA binding was measured as the equilibrium dissociation constant, Kd, by fitting nitrocellulose filter binding data from three independent experiments into an equilibrium binding model. RNA-III (Fig. 7) was used as the single-stranded RNA substrate. The double-stranded RNA substrate was obtained by annealing of RNA-III to RNA-V, a complementary RNA of the same size: 3′-GGCTCTTC TAGTATACAGAAAAATATAATAGCACTACGCGGA-5′.
For annealing, the RNAs at 5 μM were denatured by incubating at 95°C for 2 min in 20 mM HEPES, pH 7.1, and cooled to 20°C over 30 min followed by addition of MgCl₂ to 10 mM. RNA-III was denatured and quick-cooled to melt out a possible hairpin structure produced by self-folding. For Kd determinations, the RNA concentration was kept constant at 5 nM, and increasing amounts of protein were used. The percentage of active protein was determined in reciprocal experiments with 2 nM of protein and increasing concentrations of RNA and used to adjust concentration values in the Kd calculations. The purified complex was determined to be 60% active, and the recombinant Ltp28 and Ltp26 were 25%–35% active, depending on the preparation. The Prizm3 (GraphPad) software package was used for all nonlinear regression curve fitting and statistical analyses.

Western blotting and immunoprecipitation

Rabbit and mouse polyclonal antibodies were raised against the Ltp26/28 complex from *L. tarentolae*, which was purified by heparin fraction and preparative 10%–20% SDS gel electrophoresis, and also against recombinant proteins (Covance Research Products). Antibodies were affinity-purified on immobilized antigen columns by standard techniques. IPs were carried out in 300–400 μL of extract (2 mg/mL) or glycerol gradient fractions (supplemented with BSA to 1 mg/mL) with 10–15 μL of GammaBind protein G beads (AP Biotech) coated with 1–5 μg of affinity-purified antibodies or 100 μg of total IgG fraction for 2 h at 4°C with constant mixing. Beads were pelleted by brief centrifugation, washed 3 times with 1 mL of 20 mM Tris HCl, pH 7.5, 500 mM NaCl, 0.5% TritonX100 and resuspended in 50 μL of 1×SDS loading buffer. Alternatively, total RNA was isolated from mitochondrial extract or IP material by adding SDS to 1%, followed by phenol/chloroform extraction and ethanol precipitation. Electrophoretic transfer on nitrocellulose membrane from SDS gels was performed for 2 h at 80 V in Mini Trans-Blot cells (Bio-Rad) in 25 mM Tris, 190 mM glycine, 10% methanol. Immunodetection was carried out with SuperSignal West Pico chemiluminescent substrate (Pierce) by standard techniques.

Glycerol gradient sedimentation

Mitochondrial pellets were resuspended in 10 mM HEPES, pH 7.6, 10 mM MgCl₂, 60 mM KCl, 125 mM sucrose at 25 mg total protein per mL, and lysed on ice by adding TritonX100 to 0.3%. The extract was clarified by centrifugation at 200,000 g for 15 min and 200 μL of supernatant loaded on 10%–30% glycerol gradient formed in a centrifuge tube for the SW 41 Beckman rotor. After centrifugation for 20 h at 30,000 rpm, 750-μL fractions were collected from the top of the gradient. Adenylation reactions were performed on each fraction as described (Peris et al. 1997). RNA was isolated from each fraction and the gRNA specifically labeled with [α-32P]GTP as described (Blum et al. 1990).

Tandem affinity purification (TAP) of Ltp28-associated proteins

The entire Ltp28 gene was fused at the C-terminus with the TAP cassette and inserted into the pX vector (LeBowitz et al. 1990) at the BamHI site. *L. tarentolae* cells were transfected with this plasmid and selected for resistance to G418 by agar plating. The expression of the Ltp28-TAP and targeting of the protein to the mitochondrion was confirmed by Western analysis of subcellular fractions using the PAP reagent which recognizes the protein A moiety of the fusion protein. Cells were grown to late log phase, and the mitochondrial fraction was isolated.

The TAP protocol was adapted from the published method (Puig et al. 2001). Mitochondria (200 mg of total protein) were lysed on ice for 15 min in 12 mL of TMK buffer (20 mM Tris-HCl, pH 7.6, 60 mM KCl, 10 mM MgCl₂) plus 0.5% NP40. One tablet of Complete protease inhibitors (Roche) was added directly to the extract. After centrifugation at 200,000 g for 10 min, the pellet was sonicated 3 times for 10 sec in 12 mL of the same buffer without detergent and the centrifugation was repeated. Supernatants from both extraction were pooled and incubated with 0.6 mL of IgG Sepharose FF (AP Biotech) for 2 h in batch, transferred to three disposable 5-mL columns (Pierce) and washed sequentially with 20 mL of TMK buffer plus 0.1% of NP40 and 10 mL of TEV protease cleavage buffer (20 mM Tris-HCl, pH 7.6, 60 mM KCl, 10 mM MgCl₂, 0.1% NP40, 1 mM DTT). One mL of the same buffer with 100 U of TEV protease was added to each column, and the columns were incubated overnight with constant mixing. The IgG resin was drained and washed with 2 mL of CB buffer (20 mM Tris-HCl, pH 7.6, 60 mM KCl, 10 mM MgCl₂, 1 mM imidazole, 10 mM β-mercaptoethanol, 2 mM CaCl₂, 0.1% NP40). Eluted material was incubated with 0.6 mL of calmodulin agarose (Stratagene) for 1 h, transferred to two disposable 5-mL columns (Pierce), washed with 20 mL of CB buffer, and eluted with 3 mL of 20 mM Tris-HCl, pH 7.6, 60 mM KCl, 10 mM MgCl₂, 2 mM EGTA, 2 mM CHAPS. Eluted proteins were concentrated with Slide-A-Lyzer solution (Pierce) to 300 μL and loaded on a 10%–30% glycerol gradient. Sedimentation was performed in the SW41 rotor (Beckmann) for 20 h at 30,000 rpm. Fractions of 750 μL were collected from the top of the gradient.

**REFERENCES**


