Mitochondrial tRNA import in *Trypanosoma brucei* is independent of thiolation and the Rieske protein

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ABSTRACT

Due to a complete lack of the tRNA genes in the mitochondrial genome of *Trypanosoma brucei*, all tRNAs needed for mitochondrial translation have to be imported into the organelle from the cytosol. A previous study showed that the modified nucleotide s2U could act as a negative determinant for mitochondrial tRNA import in another kinetoplastid, *Leishmania tarentolae*. We have investigated whether the same type of cytosolic control for tRNA retention exists in *T. brucei*. Based on Northern analysis with subcellular RNA fractions and in vitro import assays, we demonstrate that silencing of the cysteine desulfurase, TbNfs (TbIscS), the key enzyme in tRNA thiolation (s2U) and Fe-S cluster formation in vivo, has no effect on tRNA partitioning. This observation is especially surprising in light of a recent report suggesting that in *L. tropica* the Rieske Fe-S protein is an essential component of the RNA import complex (RIC). In line with the above observation, we also show that down-regulation of the Rieske protein by RNA interference, similar to the TbNfs knockdowns, has no effect on import. The data presented here supports the view that in *T. brucei*: (1) s2U is not a negative determinant for tRNA import; (2) the Rieske protein is not an essential component of the import machinery, and (3) since the Rieske protein is essential for respiration and maintenance of inner mitochondrial membrane potential, neither process plays a critical role in tRNA import. We therefore suggest that the *T. brucei* import machinery differs substantially from what has been described in *Leishmania*.

Keywords: *T. brucei*; tRNA import; 2-thiolation; RIC; Rieske; Fe-S cluster

INTRODUCTION

In a majority of eukaryotes, most of the ATP produced in cells is the result of mitochondrial respiration, which depends on the synthesis and assembly of respiratory complexes. The bulk of the respiratory proteins are encoded in the nucleus, translated in the cytoplasm, and subsequently imported into the mitochondria where together with a few mitochondria-encoded subunits constitute the mature, fully functional complexes. Therefore, nominally, mitochondrial respiration requires both active cytosolic protein import systems and protein synthesizing machinery within the mitochondrial matrix.

Although mitochondrial protein synthesis is invariably essential, many mitochondrial genomes retained only an incomplete set of tRNAs insufficient for translation (Salinas et al. 2008). These organisms have thus evolved mechanisms to import tRNAs across the mitochondrial membranes. To date, tRNA import systems have been described in a number of diverse organisms including the ciliate *Tetrahymena* (Suyama 1986; Rusconi and Cech 1996a,b), the apicomplexan *Toxoplasma* (Esseiva et al. 2004), the kinetoplastid flagellates *Trypanosoma* and *Leishmania* (Simpson et al. 1989; Mottram et al. 1991; Hancock et al. 1992), plants (Marechal-Drouard et al. 1988), yeast (Martin et al. 1979; Rinehart et al. 2005), and most recently mammals (Rubio et al. 2008).

Two main general mechanisms have been described for tRNA import: one mechanism originally described in yeast for the import of tRNA15S is similar to protein import (Tarassov et al. 1995a). This system requires an electrochemical potential across the mitochondrial inner membrane, and with the exception of the mitochondrial outer membrane protein MOM72, all other protein import components play a role in tRNA15S import (Martin et al. 1979; Tarassov and Martin 1996; Tarassov et al. 1995a).
Import also requires the in vivo association of the tRNA with the precursor mitochondrial tRNA synthetase and aminoacylation of the tRNA by the cognate cytosolic synthetase (Schmitz and Lonsdale 1989; Tarassov et al. 1995b), and the metabolic enzyme enolase to help deliver the imported tRNA/synthetase complex to the import machinery on the mitochondrial surface (Entelis et al. 2006).

Second, and perhaps the most common mechanism for tRNA import, was originally described in *Leishmania* and is now also found in most other organisms ranging from kinetoplastids to humans (Schneider and Marechal-Drouard 2000; Salinas et al. 2008). This mechanism is independent of the protein import pathway and can be efficiently reproduced in vitro in the absence of cytosolic factors (Mahapatra et al. 1994; Yermovsky-Kammerer and Hajduk 1999; Rubio et al. 2000, 2008); however, it may still be influenced by their presence. For example, in *Trypanosoma brucei*, cytosolic translation elongation factor 1a (eEF1a) plays a key role as a specificity determinant for some imported tRNAs (Bouzaidi-Tiali et al. 2007). More recently, the import of tRNA$^{\text{Glu}}$ into yeast mitochondria was shown to occur in vitro in the absence of added cytosolic factors (Rinehart et al. 2005). Therefore, *Saccharomyces cerevisiae* contains two pathways for tRNA import, a feature that is thus far unique to this organism (Rinehart et al. 2005).

What constitutes a conserved tRNA determinant or recognition element for organellar import has also been under debate. In *Leishmania*, the mature tRNA is the in vivo substrate for import (Kapushoc et al. 2000). Although a full-length tRNA is the preferred substrate in vitro (Rubio et al. 2000), smaller substrates representing regions of the full-length tRNA could be efficiently imported and apparently contain useful import information (Mahapatra et al. 1998). Furthermore, in *L. tarentolae*, Suzuki and coworkers showed that tRNA modifications, specifically the presence of 2-thioridine (s$^2$U) in the anticodon of tRNA$^{\text{Glu}}$, could serve as a negative determinant for import. In their system, only the unmodified tRNA (not containing s$^2$U) was the substrate for import in vivo and in vitro, while the modified version was retained in the cytoplasm (Kaneko et al. 2003).

In recent years, numerous protein factors have been implicated as additional requirements for import of tRNAs into mitochondria. In plants, for instance, the voltage-dependent anion channel (VDAC) in the outer membrane was shown to play a crucial role for import in vitro (Salinas et al. 2006). However, the homologous protein is dispensable for import in *T. brucei* (Pusnik et al. 2009). Despite sharing a common basic mechanism, not requiring the protein import pathway, a majority of tRNA import systems may thus possess nuances that set them apart and that may reflect the polyphyletic origin of mitochondrial tRNA import machineries (Schneider and Marechal-Drouard 2000).

To date, one of the most studied import systems is that of *Leishmania*, the causative agent of human leishmaniasis. Adhya and coworkers have identified a putative RNA import complex (RIC) in *L. tropica*, which contains a number of proteins known to serve alternative functions in respiration, introducing the idea of a “moonlighting” function for these proteins (Mukherjee et al. 2007). The core import complex contains six essential factors required for both in vivo and in vitro import: α subunit of the F$_1$-ATP synthase; subunit 6b and the Rieske protein of the respiratory complex III; two proteins with no similarities to any other proteins in the database; and finally, subunit 6 of the respiratory complex IV (Mukherjee et al. 2007).

In this report, we have investigated whether s$^2$U is a negative determinant for import of tRNAs into the single mitochondrion of *T. brucei*, a generally close relative of *Leishmania*, hoping to establish commonalities between the two systems. Using RNA interference (RNAi), we have knocked down two *T. brucei* proteins, the homologs of which play a role in *Leishmania* tRNA import: TbNfs (formerly TbIscS2) and the Rieske protein, both previously shown to be essential for the Fe-S cluster assembly and respiration in *T. brucei*, respectively (Smid et al. 2006). We show that although the inhibition of *T. brucei* TbNfs led to a concomitant decrease of tRNA$^{\text{Glu}}$ thiolation, it had no effect on the distribution of this tRNA species in vivo or in vitro, suggesting that s$^2$U is not a negative determinant for tRNA import in this protist. Furthermore, our findings also question the conservation of the Rieske protein as a general and essential tRNA import factor, since down-regulation of this protein causes dramatic decrease of mitochondrial membrane potential and cytochrome-dependent respiration, yet without causing any effect on the tRNA distribution in vivo or in vitro. Taken together, the data presented here support the view that thiolation, respiration, membrane potential, and most importantly, the Rieske protein are not essential components of the *T. brucei* mitochondrial tRNA import pathway.

**RESULTS**

**TbNfs is essential for growth of *T. brucei***

TbNfs is the master desulfurase responsible for both Fe-S cluster formation and tRNA thiolation. The putative function of the s$^2$U modification as a negative determinant for tRNA import into the *T. brucei* mitochondria can be tested in cells depleted for TbNfs, as this protein is known to be essential for thiolation at U34 in tRNA$^{\text{Glu}}$ (Kambampati and Lauhon 2000, 2003; Mihara et al. 2002; Nilsson et al. 2002; Nakai et al. 2004; Ikeuchi et al. 2006). We therefore took advantage of the previously described *T. brucei* procyclic cells, in which TbNfs (=TbIscS2) can be inducibly down-regulated (Smid et al. 2006). With these cells, inhibition of growth is apparent 3 d after the induction.
of RNAi, and the cells virtually stopped growing by day 5 (Fig. 1A). The RNAi-induced cells showed no recovery even after 8 d of cultivation (Fig. 1; data not shown). The extent of TbNfs silencing was evaluated using specific polyclonal anti-TbNfs antibodies, which revealed almost complete elimination of the targeted protein by day 3 of RNAi induction (Fig. 1B). This observation leads to the conclusion that TbNfs is essential for growth of procyclic T. brucei.

$s^2$U is not a determinant for tRNA import into the T. brucei mitochondrion

To determine whether TbNfs has a role in tRNA import, levels of several tRNAs were measured in mitochondria isolated from the noninduced and RNAi-induced TbNfs knockdown cells. Day 3, after the addition of tetracycline to induce RNAi, was selected for subsequent experiments, as the growth of the induced cells decreases after this time point (Fig. 1A), while the TbNfs protein is virtually eliminated (Fig. 1B). After day 3 the culture still remains in proper physiological condition as determined by light microscopy (data not shown), which effectively excludes the influence of possible secondary phenotypic effects.

Two different approaches were used to investigate the role of the 2-thiouridine modifications in tRNA import. RNA isolated from organelles purified from the noninduced and induced TbNfs cells were analyzed by Northern blotting using a panel of oligonucleotide probes specific for either thiolated or nonthiolated tRNAs (Fig. 1C). The former class was represented by tRNA$^{Glu}$ and tRNA$^{Gln}$, while four nonthiolated tRNAs were assayed (tRNA$^{Gly}$, tRNA$^{Ile}$, tRNA$^{Cys}$, and tRNA$^{Val}$). The transcript levels of 12S mitochondrial rRNA should be unaltered by the interference against TbNfs and thus served as a reference for calculation of relative abundances of the examined tRNA species. Prior to Northern analysis for tRNA distribution, subcellular RNA fractions were routinely tested for compartment cross-contamination by hybridization with a probe against the spliced leader (SL) RNA, which is confined to the cytosol. Indeed, no SL RNA signal was detected in the organellar RNA fractions while a clear signal was seen in the cytosolic fractions, indicating that the mitochondrial fractions had negligible levels of cytosolic contamination (Fig. 1C).

Surprisingly, quantification of the import of all tRNAs revealed no major differences between the nonthiolated and thiolated tRNA species (Fig. 1C). For individual tRNAs, differences between their presence in mitochondria isolated from the noninduced and RNA-induced cells ranged from 81% to 112% (or a ratio of 0.81 to 1.21 when the noninduced signal is divided by the induced for each tRNA). The mitochondrial localization did not correlate with the thiolation status of the molecules. A slight increase in the levels of tRNA$^{Glu}$ was observed upon RNAi induction of TbNfs; however, this increase was deemed insignificant in that a slight decrease was documented for the other thiolated tRNA (tRNA$^{Gln}$) examined (Fig. 1C). These experiments lead to the conclusion that $s^2$U formation in tRNA is not a negative determinant for in vivo mitochondrial import in T. brucei.

To further substantiate the results described above, an in vitro import assay was performed following a previously described protocol (Rubio et al. 2000). First, we tested the import of native tRNA$^{Glu}$ and compared its import behavior with a similar sample that was pretreated with hydrogen peroxide to remove the $s^2$U group. To ensure dethiolation, the H$_2$O$_2$-treated tRNA was separated in an (N-acryloylamino) phenyl-mercuric chloride (APM) gel (Crain et al. 2002). Both the H$_2$O$_2$-treated and untreated native tRNA$^{Glu}$ species were then $5'$-labeled. Increasing concentrations of radioactively labeled dethiolated or thiolated tRNA$^{Glu}$ were then used in import assays by incubating increasing concentrations of either substrate with
Percoll gradient-purified mitochondria in the presence of ATP (Fig. 2). Reactions in which tRNAs were incubated in the import buffer but in the absence of mitochondria or micrococcocal nuclease served as markers during electrophoresis, as well as quantification standards. Reactions in which tRNAs were incubated in the presence of nuclease but in the absence of mitochondria served as a negative control for import. These experiments showed that regardless of whether or not a given tRNA is thiolated, it is efficiently imported into the mitochondrion in vitro (Fig. 2).

We also performed similar reactions with in vitro radioactively labeled transcripts representing either tRNA\textsuperscript{Glu} or tRNA\textsuperscript{Ile} (a nonthiolated tRNA) comparing mitochondrial tRNA import purified from Percoll gradient-purified mitochondria from the noninduced TbNfs cells (Fig. 3A,B). We found that, analogous to the in vivo situation, import is not affected by the elimination of the cysteine desulfurase TbNfs (Fig. 3C). Both tRNAs are imported in vitro with comparable efficiencies. Taken together, we can conclude that neither TbNfs nor the presence of s\textsuperscript{2}U in the tRNA play significant roles as tRNA import determinants in vivo or in vitro.

**Rieske is essential for growth but not for mitochondrial tRNA import in *T. brucei***

It was recently proposed that the Rieske protein is an essential component of the RIC complex in *L. tropica* (Mukherjee et al. 2007). This Fe-S cluster containing subunit of the respiratory complex III is essential for respiration and also for maintenance of the membrane potential across the mitochondrial inner membrane. In light of our observation that down-regulation of TbNfs had no effect on tRNA import into the *T. brucei* mitochondrion, and the known requirement of TbNfs for overall Fe-S cluster assembly, it is difficult to reconcile our findings with the idea that the Rieske protein could also be essential for tRNA import in *T. brucei*. To test this proposal further, we have generated clonal cell lines for the inducible down-regulation of the Rieske protein expression by RNAi (Horváth et al. 2005). Since the cell growth decreases at day 4 following RNAi induction (Fig. 4A), this time point was selected for further experiments. As in the case of TbNfs knockdowns, no resistance against RNAi was observed even in cells cultivated for a prolonged period of time. Western analysis using specific antibodies showed a dramatic decrease of the Rieske protein levels upon the addition of tetracycline (Fig. 4B).

Next, the mitochondrial RNAs isolated from the noninduced cells, as well as from cells 4 d after RNAi induction, were subjected to Northern analysis with the same set of oligonucleotide probes used to analyze both nonthiolated and thiolated tRNAs (Fig. 4C). The values obtained from A were used to calculate the fmoles of tRNA protected in the assay, which was plotted versus input tRNA in fmoles.
and thiolated tRNAs. As before, probes against the SL RNA and 12S tRNA were used to assess sample purity and served as loading controls, respectively (Fig. 4C). These results showed no differences between the import behavior of the tRNAs tested, including the nonthiolated and/or thiolated species (tRNA Glu and tRNA Gln) were used as loading controls. (C) The effect of RNAi on the Rieske protein levels was analyzed by Western blots with extracts from the noninduced (Tet−) and RNAi-induced (Tet+) cells. Cells were collected 4 d after RNAi induction. Each lane was loaded with total protein from ~5 × 10⁶ cells, and polyclonal anti-Rieske protein antibodies were used for detection. Similar Western blots with anti-TbNfs antibodies were used as loading controls. (C) Northern blot analysis of mitochondrial tRNAs purified from the noninduced (−) and RNAi-induced (+) Rieske protein cells. tRNAs were visualized with radioactively labeled oligonucleotide probes specific for tRNA^{Glu}, tRNA^{Gln}, tRNA^{Gly}, tRNA^{Ile}, tRNA^{Cys}, and tRNA^{Val}. Amounts of the individual tRNAs, after RNAi induction, were calculated as the ratio of noninduced (Tet−) to induced (Tet+) signals, after normalization against the 12S mitosomal RNA. The numbers indicate differences between the noninduced and induced cells, where a ratio of 1 indicates no differences between the two samples. (Arrows) Thiolated tRNAs. Cytosolic spliced leader (SL) RNA was used as a control for mitochondrial RNA purity.

**DISCUSSION**

Most eukaryotic cells contain an extensive degree of intracellular compartmentalization that allows subcellular organelles to keep distinct metabolic processes separated, often as a means to prevent unwanted reactions. In kinetoplastid flagellates, for example, recent studies have shown that compartmentalization of the glycolytic enzymes prevents accumulation of toxic glycolytic intermediates in the cytoplasm (Haanstra et al. 2008). What leads to intracellular compartmentalization is an extensive network of membrane systems, which are the foundations of intracellular architecture. Mitochondria and various types of plastids are unique membrane-bound intracellular compartments that usually contain their own genome, which encodes a variable number of protein-coding genes essential for organellar functions. However, these genomes are dynamic and throughout evolution have relegated a number of functions to the nucleus via gene transfer and gene loss. Indeed, a wide majority of mitochondrial proteins are encoded in the nucleus, synthesized in the cytoplasm and subsequently imported into the organelle. Remarkably, compartmentalization and the loss of mitochondrial genetic material frequently led to the disappearance of tRNA genes from mitochondrial genomes. This has in turn driven the evolution of mechanisms for the import of tRNAs from the cytoplasm to enable translation of the protein coding mRNAs still encoded in mitochondria (Salinas et al. 2008). Currently, however, the factors and/or mechanisms that control tRNA import are not fully understood.

Despite differences in the number and types of factors shown to be associated with tRNA import in various organisms, tRNAs can mechanistically be imported in two different ways: either via the protein import pathway (e.g., coimport with cytosolic aminoacyl-tRNA synthetase) (Tarassov et al. 1995b) or via a pathway independent from protein import that does not require cytosolic factors (Rubio et al. 2000). Future studies may reveal additional pathways. However, the disparate nature of the import mechanisms suggests that tRNA import evolved independently in different systems (Schneider and Marelchal-Drouard 2000). Differences among various systems may

**FIGURE 4.** Effects of Rieske protein RNAi on parasite growth and tRNA localization in vivo. (A) Growth curves of the noninduced (TET−) (triangles) and RNAi-induced (TET+) (squares) knockdowns for the Rieske protein. Cell densities were measured using the Burkner cell chamber. The y-axis is labeled by a log scale and represents the products of measured cell densities and the total dilutions. The arrow indicates sampling time point for latter experiments. (B) The effect of RNAi on the Rieske protein levels was analyzed by Western blots with extracts from the noninduced (−) and RNAi-induced (+) cells. Cells were collected 4 d after RNAi induction. Each lane was loaded with total protein from ~5 × 10⁶ cells, and polyclonal anti-Rieske protein antibodies were used for detection. Similar Western blots with anti-TbNfs antibodies were used as loading controls. (C) Northern blot analysis of mitochondrial tRNAs purified from the noninduced (−) and RNAi-induced (+) Rieske protein cells. tRNAs were visualized with radioactively labeled oligonucleotide probes specific for tRNA^{Glu}, tRNA^{Gln}, tRNA^{Gly}, tRNA^{Ile}, tRNA^{Cys}, and tRNA^{Val}. Amounts of the individual tRNAs, after RNAi induction, were calculated as the ratio of noninduced (Tet−) to induced (Tet+) signals, after normalization against the 12S mitosomal RNA. The numbers indicate differences between the noninduced and induced cells, where a ratio of 1 indicates no differences between the two samples. (Arrows) Thiolated tRNAs. Cytosolic spliced leader (SL) RNA was used as a control for mitochondrial RNA purity.
also suggest import as a dynamic process that has adapted to intracellular environmental changes provided by a particular organism.

Among the most studied tRNA import pathways are those of kinetoplastids, where a number of in vitro and/or in vivo determinants for import have been described (Salinas et al. 2008). It was reported that thiolation of tRNAs in the cytoplasm of *L. tarentolae* (Suyama et al. 1998). Among the closely related genera *Trypanosoma* and *Leishmania*, TbNfs also serves as a key enzyme in Fe-S clusters synthesis pathway (Lill and Muehlenhoff 2008). Since the Rieske protein contains several Fe-S clusters, the lack of effect of the TbNfs knockdown on tRNA import also brings into question the essentiality of the Rieske protein for this process. Thus, in line with results obtained with the TbNfs knockdown cells, down-regulation of the Rieske protein levels by RNAi had no effect on tRNA import in vivo and/or in vitro in *T. brucei*.

In an attempt to find commonalities among the various kinetoplastid systems, and prompted by the availability of RNAi in *T. brucei*, we explored the use of thiolation as a conserved negative determinant for import in vivo in these ancestral flagellates. To this end, we have generated conditional RNAi knockdown cell lines of TbNfs and showed that like in all other organisms described so far, TbNfs is essential for tRNA thiolation in *T. brucei* (J Lukës and JD Alfonzo, in prep.). Indeed, reduction in the expression levels of TbNfs leads to the expected reduction in thiolation of tRNA^{Glu} and tRNA^{Gln} but, significantly, causes no defect in tRNA distribution. Based on this result, we postulate that, rather unexpectedly, the use of thiolation as a negative import determinant is not conserved between the closely related genera *Trypanosoma* and *Leishmania*. TbNfs also serves as a key enzyme in Fe-S clusters synthesis pathway (Lill and Muehlenhoff 2008). Since the Rieske protein contains several Fe-S clusters, the lack of effect of the TbNfs knockdown on tRNA import also brings into question the essentiality of the Rieske protein for this process. Thus, in line with results obtained with the TbNfs knockdown cells, down-regulation of the Rieske protein levels by RNAi had no effect on tRNA import in vivo and/or in vitro in *T. brucei*.

It was shown earlier in *T. brucei* that the Rieske subunit of respiratory complex III is essential for both maintenance of membrane potential and respiration (Horváth et al. 2005). Our current findings that the Rieske protein is not involved in tRNA import also firmly established that neither process plays a major role in mitochondrial tRNA localization and corroborates previous observations.
as needed (50 μg/mL hygromycin, 15 μg/mL neomycin, and 2.5 μg/mL phleomycin). Synthesis of double-stranded (ds) RNA was induced by the addition of 1 μg/mL tetracycline. Growth rates of the noninduced and RNAi-induced clonal cell lines were determined every 24 h using the Beckman Z2 Coulter counter over a period of 8 d. Mitochondrial vesicles were isolated from the noninduced and RNAi-induced cells by hypotonic lysis following protocols described elsewhere (Kapushoc et al. 2000).

**Immunoblot analysis**

Cell lysates of *T. brucei* procyclics corresponding to 5 × 10⁶ cells/lane were resolved on 12% SDS–polyacrylamide gel. The polyclonal rabbit antibodies against the *T. brucei* TbNfs and Rieske proteins were used as described previously (Horváth et al. 2005; Smid et al. 2006). Secondary anti-rabbit IgG antibodies (1:2000) coupled to horseradish peroxidase (Sevapharma) were visualized according to the manufacturer’s protocol using the ECL kit (Pierce).

**tRNA localization using Northern blot analysis**

RNA was isolated using the guanidinium thiocyanate/phenol/chloroform extraction method (Chomczynski and Sacchi 1987) either from whole cells or from the mitochondrial fractions. Samples (2.5 μg each) were separated on denaturing 8% polyacrylamide gel with 8 M urea and electroblotted to Zeta-probe (Bio-Rad) membranes, which were subsequently probed with [³²P]-5′ end-labeled oligonucleotides specific for each RNA as follows:

<table>
<thead>
<tr>
<th>tRNA</th>
<th>Oligonucleotide Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cys-GCA</td>
<td>5′-GGGACCATTCGGAACCCT-3′</td>
</tr>
<tr>
<td>Gin-UUG</td>
<td>5′-CAGGATTGGAACCTGGGTT-3′</td>
</tr>
<tr>
<td>TTCG-3′</td>
<td></td>
</tr>
<tr>
<td>Glu-UUC</td>
<td>5′-TCCGGATACCCGGAATACACG-3′</td>
</tr>
<tr>
<td>Gly-UCC</td>
<td>5′-TCTCCGGGCTTGGAAAGGGCGAAAATC-3′</td>
</tr>
<tr>
<td>Ile-UAU</td>
<td>5′-CGGTTTCGACCCCGATATTCGGT-3′</td>
</tr>
<tr>
<td>Val-UAC</td>
<td>5′-ACGTTCCTGCTGAAGACAGACATC-3′</td>
</tr>
<tr>
<td>125 rRNA</td>
<td>5′-GCTGCTACTGAGGAGCCTTTCATAC-3′</td>
</tr>
<tr>
<td>SLRNA</td>
<td>5′-AGGAGAATGAGCATTGGCCT-3′</td>
</tr>
</tbody>
</table>

Hybridization procedures were carried out according to the manufacturer’s instructions (Bio-Rad). Images were taken with a Storm PhosphorImager (Molecular Dynamics). Abundance of individual tRNA species, isolated from cells prior to or after RNAi induction, was calculated in percentages using the ImageQuant program after normalization to the 12S RNA signal.

**In vitro import assays**

For import assays, 1 mg of purified *T. brucei* mitochondria was incubated with 50–100,000 cpm of radioactively labeled tRNA,
Reactions were then stopped by the addition of 10 mM EGTA (pH 8.0). To purify the protected tRNAs, the mitochondria were washed with 0.25 M sucrose/20 mM Tris–HCl (pH 8.0), pelleted, suspended in 90 μl of 10 mM Tris–HCl (pH 8.0), 1 mM EDTA, and phenol extracted, followed by ethanol precipitation. The radioactively labeled tRNAs were separated by electrophoresis through a 7 M urea/6% polyacrylamide gels. After electrophoresis, the gels were dried onto Whatman 3MM chromatography paper and visualized with the Storm PhosphorImager imaging system. Import assays were performed with either radioactive in vitro transcript representing the various tRNAs or with native tRNAs. Whenever native substrates were used, they were hybridized with an antisense biotinylated oligonucleotide specific for each species and purified by affinity chromatography through streptavidin beads as described elsewhere (Crain et al. 2002). Native tRNAs were 5’-end-labeled as described (Kaneko et al. 2003).

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