Mutations in a tRNA Import Signal Define Distinct Receptors at the Two Membranes of *Leishmania* Mitochondria

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Nucleus-encoded tRNAs are selectively imported into the mitochondrion of *Leishmania*, a kinetoplastid protozoan. An oligoribonucleotide constituting the D stem-loop import signal of tRNA^{Tyr}(GUA) was efficiently transported into the mitochondrial matrix in organello as well as in vivo. Transfer through the inner membrane could be uncoupled from that through the outer membrane and was resistant to antibody against the outer membrane receptor TAB. A number of mutations in the import signal had differential effects on outer and inner membrane transfer. Some mutants which efficiently traversed the outer membrane were unable to enter the matrix. Conversely, restoration of the loop-closing GC pair in reverse resulted in reversion of transfer through the inner, but not the outer, membrane, and binding of the RNA to the inner membrane was restored. These experiments indicate the presence at the two membranes of receptors with distinct specificities which mediate stepwise transfer into the mitochondrial matrix. The combination of oligonucleotide mutagenesis and biochemical fractionation may provide a general tool for the identification of tRNA transport factors.

Mitochondria are genetic parasites presumed to have evolved from endosymbiotic bacteria. The recent sequencing of a large number of mitochondrial genomes has revealed an unexpected diversity in size and gene content (5). The loss of most of the original bacterial protein-coding genes, or their transfer to the nucleus, has been explained in terms of a “big bang” radiation of the different eukaryotic lineages from a single (monophyletic) endosymbiont (5). What is not so easily explained is the loss of a variable number of tRNA genes apparently randomly in different protists, higher plants, and at least one invertebrate lineage (6). At least 24 different tRNA species are required to read the universal genetic code. In kinetoplastid protozoa, including leishmanias and trypanosomes, a complete set of tRNAs are apparently imported in order to compensate for the total lack of mitochondrial tRNA genes (7, 8, 11, 16, 21–23). The mitochondria of tetrhymana import about two-thirds of their tRNAs (2), while in budding yeast only a single tRNA^{Glu} species is imported (15). In higher plants, different species mitochondrially import different sets of tRNAs; for example, mitochondria from wheat but not maize import tRNA^{His} (9). Thus, there appears to be a species-specific selectivity, reflecting perhaps the presence of different mitochondrial receptors recognizing individual or groups of import signals on tRNAs.

To study the basis of the selectivity of tRNA import, we have developed an in organello system using leishmania mitochondria (1, 12, 14). Similar systems from trypanosomes have been recently reported (18, 25). Our experiments showed that tRNA import is selective, e.g., tRNA^{Tyr} is efficiently imported whereas tRNA^{Glu}(CUG) is not (1), and that a conserved sequence motif in the D arm of tRNA^{Tyr} is necessary and sufficient for import (13). The importance of the D loop of tRNA^{His} for import in vivo has also been demonstrated (10). Importable RNA interacts directly with the outer mitochondrial membrane (12), and a 15-kDa RNA binding protein, TAB, associated with the outer membrane (OM) was purified and shown to function as an import receptor (1). Using specific antibody, it was further shown that TAB interacts with tRNA^{Tyr} but not with tRNA^{Glu} (13). Thus, the TAB-tRNA interaction accounts for the selectivity of import in this system.

More recent studies have indicated that, in addition to sequence discrimination at the OM, a second level of selection possibly occurs at the inner membrane (IM). Analysis of the distribution of imported tRNA^{32P} and other transcripts in the different intramitochondrial compartments, i.e., OM, intermembrane space (IMS), IM, and matrix (MX), showed that import occurs in a stepwise fashion, with a distinct kinetic separation of the OM and IM transfer steps (17). While OM transfer requires ATP, IM transfer is driven by both the electrical and chemical components of the electromotive force generated at the IM of energized mitochondria (17). These results lead to the hypothesis that distinct receptors occur at the OM and IM which concertedly determine sequence selectivity for MX entry of tRNAs. However, nothing is known about the specificity of the interaction at the IM or the identity of any of the components of IM transport machinery.

Here we show that a short oligoribonucleotide containing the D arm of tRNA^{Tyr}(GUA) rapidly and efficiently transits to the mitochondrial MX in vitro. Site-specific mutagenesis of the D arm of tRNA^{Tyr}(GUA) rapidly and efficiently transits to the mitochondrial MX in vitro. Site-specific mutagenesis of the D arm was employed to detect similarities and differences in the sequence and/or structural requirements for OM and IM transfer.

**MATERIALS AND METHODS**

**Cell culture and preparation of mitochondria.** Promastigotes of *Leishmania tropica* strain UR6 were cultured at 22°C on solid blood agar medium (3) supplemented with 150 µg of biotin per ml and 50 µg of adenine per ml. Mitochondria were purified from DNase I-treated lysates by Percoll gradient centrifugation and stored in a 50% glycerol storage buffer, as described previously (14). Before use, mitochondria were diluted with cold isotonic sucrose-Tris-EDTA (STE) buffer (14), washed by centrifugation, and resuspended in STE at a final protein concentration of 8 to 10 mg/ml.

**Submitochondrial fractionation.** Separation of submitochondrial compartments was performed as previously described (17). Briefly, mitochondria were treated with 320 µM digitonin in STE buffer for 15 min on ice to selectively solubilize the OM (19). Mitoplasts were separated from the soluble fraction (OM plus IMS) by centrifugation. To further fractionate the mitoplasts, they were suspended in a solution of 0.6 M sucrose, 10 mM Tris-HCl (pH 7.5), and 1 mM EDTA and subjected to three freeze-thaw cycles. Soluble MX and particulate IM fractions were subsequently separated by centrifugation.
Preparation of import substrates. 32P-labeled tRNA\textsubscript{GUA} transcripts were synthesized by runoff transcription of the plasmid pSKB1 (1), which contains a cloned genomic copy of the corresponding Leishmania gene, using T7 RNA polymerase and [\textsuperscript{32}P]UTP, as described previously (4). Wild-type and mutant D arm minihelix RNAs were synthesized by runoff transcription of the corresponding double-stranded oligonucleotides containing a T7 RNA polymerase promoter. To prepare the templates, the promoter primer GG\textsubscript{T}GCG\textsubscript{T}CTG\textsubscript{T}CTA\textsubscript{G}CTC\textsubscript{TAC}A\textsubscript{TAC}TAC\textsubscript{T}A\textsubscript{G}G\textsubscript{G}G\textsubscript{G}A\textsubscript{G}C\textsubscript{T}A\textsubscript{G} (containing an EcoRI linker, a T7 RNA polymerase promoter, and nucleotides 5 to 13 of tRNA\textsubscript{GUA}) (11) (see Fig. 1), was annealed to the following oligonucleotides, each containing sequences complementary to positions 5 to 27 of tRNA\textsubscript{GUA}: wild-type, ATGC\textsubscript{T}GCT\textsubscript{C}AC\textsubscript{C}AAC\textsubscript{T}G\textsubscript{A}G\textsubscript{T}AC\textsubscript{G}\textsubscript{A}GT\textsubscript{C} (C\textsubscript{17}–A\textsubscript{23}–A, ATGC\textsubscript{T}GT\textsubscript{C}A\textsubscript{T}A\textsubscript{C}T\textsubscript{A}T\textsubscript{G}G\textsubscript{A}G\textsubscript{T}AC\textsubscript{G}AC\textsubscript{T}G\textsubscript{C}G\textsubscript{A}G\textsubscript{C}GT\textsubscript{C}T\textsubscript{A}C\textsubscript{A}T\textsubscript{C}AC\textsubscript{G}T\textsubscript{G}C\textsubscript{T} (C\textsubscript{21}–A\textsubscript{27}–A, ATGC\textsubscript{T}GT\textsubscript{C}A\textsubscript{T}A\textsubscript{C}T\textsubscript{A}T\textsubscript{G}G\textsubscript{A}G\textsubscript{T}AC\textsubscript{G}AC\textsubscript{T}G\textsubscript{C}G\textsubscript{A}G\textsubscript{C}GT\textsubscript{C}T\textsubscript{A}C\textsubscript{A}T\textsubscript{C}AC\textsubscript{G}T\textsubscript{G}C\textsubscript{T} (C\textsubscript{22}–A\textsubscript{28}–A, ATGC\textsubscript{T}GT\textsubscript{C}A\textsubscript{T}A\textsubscript{C}T\textsubscript{A}T\textsubscript{G}G\textsubscript{A}G\textsubscript{T}AC\textsubscript{G}AC\textsubscript{T}G\textsubscript{C}G\textsubscript{A}G\textsubscript{C}GT\textsubscript{C}T\textsubscript{A}C\textsubscript{A}T\textsubscript{C}AC\textsubscript{G}T\textsubscript{G}C\textsubscript{T})). The resulting partially double-stranded molecule was end-filled with 50 m\textsuperscript{M} carbonyl cyanide m-chlorophenylhydrazone (CCCP), containing 0.2 m\textsuperscript{M} of ATP and 2.5 m\textsuperscript{M} Mg\textsuperscript{2+}, 10 mM magnesium acetate, 1 mM dithiothreitol, 0.1 M KCl) for 30 min on ice. After pelletting the cells, the centrifugation, the cells were suspended in 0.1 ml of STE buffer by centrifugation. To measure total uptake, RNase-treated mitochondria were incubated with 32P-labeled RNA, excess RNA was removed by isopropanol precipitation, as described previously (13). Alternatively, intramitochondrial RNAs were assayed by subjecting postimport RNase-treated mitochondria to digestion treatment, followed by freeze-thaw lysis to separate the submembranous mitochondrial compartments (as described above) and recovery of 32P-labeled RNA from each fraction. Antibody inhibition experiments. This observation raises the attractive possibility of using small synthetic oligonucleotides to probe the transport systems on mitochondrial membranes. Accordingly, a 27-nucleotide RNA hairpin (minihelix) (Fig. 1A) containing the wild-type D arm sequence of the tRNA\textsubscript{GUA} was synthesized by T7 RNA polymerase-mediated transcription of the appropriate oligonucleotide template. The energy-minimized secondary structure of this molecule (Fig. 1A) contains the 4-bp stem and 8-base loop present in intact tRNA. Neither the 5' terminal extension, containing the GGA initiation sequence, nor the 3' tail forms any detectable base pair with the remainder of the molecule, and this is true for the wild type as well as for all the mutant sequences used in this study (see below). Uptake of the minihelix by isolated leishmania mitochondria was dependent on ATP; entry into the membrane-bound organelle was evident from the susceptibility of the internalized RNA to RNase in the presence of detergent (Fig. 1B). Moreover, import of the minihelix was specifically competed out by the parental molecule tRNA\textsubscript{GUA} (see Fig. 1B), indicating the utilization of a common import pathway (Fig. 1B).

The distribution of the minihelix in various intramitochondrial compartments following entry through the OM was then studied by biochemical fractionation (17). Briefly, mitochondria were incubated with 32P-labeled RNA, excess RNA was digested with RNase, and then the washed vesicles were treated with digitonin to selectively permeabilize the OM (19). After centrifugal separation of the soluble fraction (OM plus IMS) from the insoluble mitochondria, the latter was subjected to freeze-thaw cycles to liberate the soluble MX contents with the remaining insoluble fraction representing the IM. At shorter incubation times (5 min), more than 50% of the RNA was found to be associated with the IM, with the remainder being in the MX (Fig. 1C). By 10 min, most of the RNA was in the MX fraction (Fig. 1C). This kinetic pattern indicates that OM transfer is faster than IM transfer and is similar to what was previously observed with the parental molecule (17), except that the rate of IM transfer of the minihelix is noticeably higher, probably due to the absence of extra sequences which hinder import.

In the above system using intact mitochondria, IM transfer is dependent on OM transfer, making it difficult to independently assess the requirements of the former step. Therefore, mitochondrial vesicles were incubated with 32P-labeled D arm minihelix, treated with RNase, and analyzed for their RNA content. Transfer of the minihelix into the MX was dependent on ATP and inhibited by protonophore uncouplers such as CCCP (Fig. 1D), demonstrating the requirement of a proton motive force across the IM (17). Under otherwise identical conditions, the amount of wild-type oligonucleotide entering the MX was the same (2 to 2.5 fmol/100 \mu g of mitochondrial protein), irrespective of whether intact mitochondria or mitoplasts were used, indicating that the essential components of the IM machinery are not lost during mitoplast isolation.

To further assess the validity of the oligonucleotide import system, the transfer of the D arm minihelix across the OM and IM was directly compared with that of intact tRNA\textsubscript{GUA} under a variety of conditions. ATP, temperature, high concentrations of monovalent cations, dissipation of IM electromotive force by protonophores (CCCP and nigericin), and disruption of membrane potential by K\textsuperscript{+} in the presence of valinomycin all had similar or identical quantitative effects on the transfer of either substrate (Table 1). Thus, except for the kinetics of transfer (see above), there is little or no biochemical distinction between full-length tRNA and the isolated D arm.
The loop (G18) may be summarized as follows. (i) Single mutations in mutants were directly assayed for their import into mitoplasts, since the amount internalized is very low. Therefore, these to accurately assess IM transfer using intact mitochondria, those mutants which are deficient in OM transfer, it is difficult

nalized RNA present in the MX represents IM transfer. For measurement of OM transfer, while the fraction of the total internalized RNA contained negligible amounts of RNA [data not shown], is a IM-plus-MX fractions, whereas the OM-plus-IMS fractions were RNase treated and subfractionated, and the contents of RNA in the IM and MX fractions were analyzed. (C) Intramitochondrial distribution of D arm minihelix. Intact mitochondria were incubated with wild-type RNA and ATP at 37°C for the time intervals shown. After each incubation, the mitochondria were RNase treated and subfractionated, and the contents of RNA in the IM and MX fractions were analyzed. (D) Import of D arm minihelix into mitoplasts. Mitoplasts (100 μg of protein) were incubated with the wild-type minihelix in the absence (lane 1) or presence (lanes 2 through 6) of 4 mM ATP. In lane 3, Triton X-100 (0.5%) was added after the incubation. Reactions 4 and 5 contained 1 pmol of yeast tRNA or tRNA Tyr, respectively, as the competitor. In reaction 6, mitoplasts were preincubated with 50 μM CCCP before the import incubation. RNase-resistant RNA was recovered for analysis. The region of the major minihelix band in each lane was excised and counted; after background subtraction, femtomole values were computed from the specific activity.

Effect of point mutations in the D arm on transfer through OM and IM in organello. The D arm of tRNA^{35S}(GUA) contains the motif UGGUAGAGC (Fig. 1A), which is conserved in the corresponding region of tRNAs imported in leishmania and trypanosome mitochondria as well as in a synthetic transcript imported through the same pathway (13). To determine the role of the primary sequence and secondary structure in the import signal, point mutations were introduced within this region. Mutant RNAs synthesized by in vitro transcription of the corresponding oligonucleotide templates were assayed for intramitochondrial distribution after uptake in the presence of ATP. In this assay, total uptake (i.e., the amount of RNA in the IM-plus-MX fractions, whereas the OM-plus-IMS fractions contained negligible amounts of RNA [data not shown]), is a measure of OM transfer, while the fraction of the total internalized RNA present in the MX represents IM transfer. For those mutants which are deficient in OM transfer, it is difficult to accurately assess IM transfer using intact mitochondria, since the amount internalized is very low. Therefore, these mutants were directly assayed for their import into mitoplasts.

The observed effects on OM and IM transfer (Fig. 2 and Table 2) may be summarized as follows. (i) Single mutations in the loop (G18→C, U17→A, A23→C) resulted in a decline of OM transfer by four- to eightfold compared to the wild-type sequence. The effect of these mutations on IM transfer was somewhat less, being reduced to 30 to 40% of the wild-type level. (ii) The mutation G18→C at the loop-closing pair of the D arm also reduced OM and IM transfer by about 10-fold. (iii) However, if the loop-closing pair was restored by a second mutation, C13→G, OM transfer remained low (15% of wild type), but at least half of the RNA internalized into intact

TABLE 1. Comparison of import characteristics of tRNA^{35S} and D arm minihelix

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Membrane</th>
<th>Value with substrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATP</td>
<td>K_m</td>
<td>OM 1.52 mM 2.15 mM</td>
</tr>
<tr>
<td>KCl</td>
<td>IC_{50}</td>
<td>IM 0.74 M 0.80 M</td>
</tr>
<tr>
<td>CCCP</td>
<td>IC_{50}</td>
<td>IM 3.0 μM 3.0 μM</td>
</tr>
<tr>
<td>Nigericin</td>
<td>IC_{50}</td>
<td>IM 2.6 μM 3.5 μM</td>
</tr>
<tr>
<td>KCl + 5 μM</td>
<td>IC_{50}</td>
<td>IM 12.5 M 13.5 M</td>
</tr>
<tr>
<td>Valinomycin</td>
<td>IC_{50}</td>
<td>IM 0.0066 0.0053</td>
</tr>
</tbody>
</table>

a Import reactions were carried out with either mitochondria or mitoplasts (100 μg of protein) in the presence of 100 fmol of 32P-labeled RNA substrate for 15 min at 25°C (except for the temperature-dependence experiment, in which temperature was varied). The amount of RNA transferred was quantified by RNase protection, PAGE, autoradiography, and denisotmetric band scanning. Various concentrations of the indicated reagents or various temperatures were used. b K_m, the concentration of ATP for half-maximal (50%) rate of import (determined from titration curves); IC_{50}, the concentration of reagent for 50% inhibition of transfer; Δν/Δt, increase in rate of transfer (Δν) per degree Celsius rise in temperature (Δt) before saturation (between 25 and 37°C in both cases).
mitochondria was found in the MX, indicating efficient IM transfer. This was confirmed with mitoplasts; the \( G^{22}\rightarrow C \), \( C^{13}\rightarrow G \) mutant was transferred across the IM with about 80% of the efficiency of the wild-type sequence. (iv) The mutation \( A^{23}\rightarrow U \) reduced both OM and IM transfer to 20 to 30% of that of the wild type. When the AU pair was restored by a second mutation, \( U^{12}\rightarrow A \), OM transfer increased from 29 to 64% (compared to wild type), but IM transfer was not restored. (v) A double mutation in the stem, \( C^{13}\rightarrow U \), \( U^{12}\rightarrow C \), which destabilizes the stem without altering the primary sequence of the conserved motif (Fig. 1), resulted in a significant amount of OM transfer (59% of wild type) but had a more severe effect on IM transfer (reduced to 30% of wild type). These results indicate that certain mutations in the import signal result in different efficiencies of transfer across the IM and OM.

**Targeting of the D arm minihelix to the mitochondrial MX in vivo.** From the preceding in organello analysis, it is evident that mutations in the conserved D arm motif result in deficiency of transport into the mitochondrial MX, due to a defect in either OM or IM transfer. To examine whether this is also true in vivo, \( ^{32}\text{P} \)-labeled minihelices were introduced into leishmanial promastigotes by electroporation. Transfected cells were lysed by syringe passage in hypotonic medium, the mitochondrial fraction was recovered and treated with DNase and RNase, and the nuclease-resistant RNA was analyzed. Hypotonic treatment results in leakage of the contents of the IMS; this assay therefore scores exclusively for MX-localized RNA and does not distinguish between OM and IM transfer.

Approximately 20 to 40 molecules of the wild-type minihelix per transfected cell were recovered from the mitochondrial fraction under optimal conditions (Fig. 2C). Entrapment of the RNA within a membrane vesicle was evident from an enhanced RNase sensitivity in the presence of Triton X-100, but RNase resistance was unaffected by treatment with digitonin at a concentration which selectively permeabilizes the OM (17), indicating MX localization. Transfection in the presence of the mitochondrial inhibitors CCCP and oligomycin resulted in reduction of MX transfer in vivo (Fig. 2C), consistent with the sensitivity of in organello IM transfer to these agents (17) (Fig. 1). About 20% of the RNA remained RNase resistant in the presence of Triton X-100 or the inhibitors; this could be due to sequestration into an unknown subcellular component present in the crude mitochondrial fractions.

The effect of mutations on targeting in vivo was analyzed. All the mutants examined were deficient in MX localization (Fig. 2C). The \( C^{11}\rightarrow A \), \( U^{12}\rightarrow C \) mutant was targeted at about 15% of the wild-type level. These data confirm that the D arm signal is necessary and sufficient for transport into the mitochondrial MX in vivo as well as in vitro.

**Sequence-specific binding of D arm variants to OM and IM.** To determine whether the above effects on OM and IM transfer reflect altered binding to membrane-bound receptors, \( ^{32}\text{P} \)-
labeled minihelices were allowed to interact with intact mitochondria or mitoplasts under conditions favoring specific binding but discouraging translocation (i.e., in the presence of 0.1 M KCl at 0°C, in the absence of ATP) (12). The vesicles were then washed, and bound RNA was analyzed by gel electrophoresis. Control experiments (data not shown) demonstrated that the bound RNA was completely sensitive to RNase, i.e., it was associated with the membrane surface.

Both the G22→C and G22→C, C13→G mutants were bound inefficiently to the OM, but the second mutation restored IM binding to about 60% of that of the wild type (Fig. 3A). In contrast, restoration of the second base pair in the stem (i.e., A23→U, U12→A) increased binding to the OM but not to the IM (Fig. 3A). The wild-type sequence bound to the OM and IM with apparent Kd values of 0.93 and 3.86 nM, respectively (Table 3). The mutation G22→C increased Kd(OM) nearly 8-fold, while Kd(IM) was increased 12-fold. But whereas Kd(IM) remained high for the double mutant G22→C, C13→G, Kd(IM) was restored to near-wild-type levels (Table 3). The converse effect of base pair restoration at the next position (A23) was also evident: reformation of the AU pair reduced Kd(IM) by a factor of 2 but increased Kd(IM) by a factor of 3.5 (Table 3). The double mutation C11→A, U12→C had a much more severe effect on Kd(IM) (8-fold) than on Kd(OM) (2-fold). Thus, the binding efficiencies parallel the transfer efficiencies and indicate the presence of distinct receptors on the two membranes which recognize different structural features of the D arm.

Although the net yields of the complexes formed with the wild-type sequence and the G22→C, C13→G mutant were comparable, there was a notable difference in their rates of dissociation (Fig. 3B). Thus, while the wild-type complex remained stable for at least 30 min at 0°C, more than 90% of the mutant complex was dissociated within this time period.

Competition between D arm variants at OM and IM. From the above experiments, two mutants were identified which are transferred through the OM, but not IM, with good efficiency: the A23→U, U12→A mutant and the C11→A, U12→C mutant. Conversely, the G22→C, C13→G mutant is transferred efficiently through the IM but not the OM. Only the wild-type sequence is transferred through both the membranes efficiently. To examine whether the different structures use the same or distinct receptors at either membrane, competition assays were performed in which one labeled oligonucleotide was challenged with an excess of another oligonucleotide (either unlabeled or labeled to low specific activity). OM transfer of both the A23→U, U12→A and C11→A, U12→C mutants was efficiently competed out by the wild-type sequence, although in the latter case competition was somewhat less effective than in the self-self situation (Fig. 4A). At the IM, the wild-type

<table>
<thead>
<tr>
<th>Mutation(s)</th>
<th>OM transfer (% of w.t.)</th>
<th>Intramitochondrial distribution (fraction of total)</th>
<th>IM transfer (% of w.t.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None (w.t.)</td>
<td>100</td>
<td>0.30</td>
<td>100</td>
</tr>
<tr>
<td>U17→A</td>
<td>12.2</td>
<td>0.67</td>
<td>38.5</td>
</tr>
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<td>G19→C</td>
<td>23.9</td>
<td>0.73</td>
<td>30.3</td>
</tr>
<tr>
<td>A11→C</td>
<td>11.9</td>
<td>0.80</td>
<td>38.5</td>
</tr>
<tr>
<td>G22→C</td>
<td>9.0</td>
<td>0.82</td>
<td>12.7</td>
</tr>
<tr>
<td>G22→C, C13→G</td>
<td>12.8</td>
<td>0.73</td>
<td>88.3</td>
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<tr>
<td>A23→U</td>
<td>29.3</td>
<td>0.84</td>
<td>19.2</td>
</tr>
<tr>
<td>A23→U, U12→A</td>
<td>64.3</td>
<td>0.82</td>
<td>10.5</td>
</tr>
<tr>
<td>C11→A, U12→C</td>
<td>58.7</td>
<td>0.81</td>
<td>29.7</td>
</tr>
</tbody>
</table>

a Measured by total uptake into intact mitochondria and/or the sum of the distribution in intramitochondrial compartments. Quantitations were performed by gel band counting (Fig. 2). The amount of wild-type (w.t.) sequence imported (2.0 to 2.8 fmol) was taken as 100%.

b Ratio of the amount of RNA in a particular compartment to the total amount internalized (i.e., IM + MX).

c Measured by uptake into mitoplasts; quantitation performed as described for OM transfer.
The use of synthetic oligonucleotides as import substrates is based on our observation that the D arm of tRNA_Y is necessary and sufficient for import in organello (13) as well as in vivo (Fig. 2), and this approach also greatly facilitates the generation of mutations for the study of structure-function relationships. Furthermore, fractionation of the sub mitochondrial compartments allows an assessment of the effect of an individual mutation on the precise location of the corresponding structure within the mitochondrion. By all the criteria examined, i.e., competition by intact tRNA (Fig. 1), effect of anti-TAB antibody on import into mitochondria or mitoplasts (Fig. 5), dependence on ATP and temperature (Table 1), and inhibition of transfer by inhibitors and uncouplers (Fig. 1 and Table 1), the D arm minihelix appears to be using the same pathway for import as the parental molecule, although the rate and extent of import of the minihelix are noticeably higher (Fig. 1), presumably due to its smaller size and/or the lack of inhibitory sequences.

Site-specific mutagenesis of the D arm minihelix revealed a number of distinct phenotypes (Fig. 2 and Table 1). In the first group of mutants (U17→A, G→G; A23→C; A25→C; C11→A, A12→U RNA (100 fmol), respectively. Wild-type competitor (1 pmol) was included in reactions 2, 4, and 6. In panel B, high-specific-activity wild-type (lanes 1 to 5) or G22→C, C13→G (lanes 6 to 10) RNA (100 fmol) was incubated without competitor (lanes 1 and 6), with 0.5 pmol (lanes 2 and 9) or 5 pmol (lanes 3 and 10) of wild-type competitor, or with 0.5 pmol (lanes 4 and 7) or 5 pmol (lanes 5 and 8) of G22→C, C13→G competitor. Quantitations were performed by densitometric scanning of the major band in each lane.

In marked contrast, transfer of tRNA_Y as well as transfer of the wild-type D arm minihelix and the G22→C, C13→G mutant across the IM of isolated mitoplasts was resistant to anti-TAB antibody (Fig. 5). These results imply that the RNA sequence and/or structural specificities at the two membranes are nonidentical.

**DISCUSSION**

In this report, a simple method for probing the structural basis of the selectivity of mitochondrial tRNA import is described.
The differences are particularly pronounced when bases in the stem of the D arm (Fig. 1A) are altered. Each of the single mutations (G12-C and A23-U), or the double mutation on the same side of the helix (C11-A, U12-C), results in a drastic reduction of secondary structure stability (Table 3); the FOLDRNA program (26) predicts relatively unstable energy-minimized structures with the canonical 4-bp stem replaced by loops, bulges, and non-Watson-Crick pairs (data not shown). Restoration of the loop-closing pair by the second C13-G mutation, or of the neighboring pair by the U12-A mutation, results in restoration of the wild-type conformation. It would seem from this limited structure-function study that the loop-closing pair (G22:C13) is critical at both membranes, but for different reasons: while the G base is specifically recognized at the OM, base pairing is more important at the IM, since restoration of the pair in reverse results in reappearance of IM transfer (Fig. 2) and an increase in binding affinity for the IM receptor (Table 3). Conversely, the A23 residue critically interacts with the IM receptor(s) but may be replaced by U at the OM. This would explain the lack of reversion for IM transfer in the A23-U, U12-A mutant (Fig. 2); in fact, the binding affinity of this mutant for the IM is further lowered (Table 3). The importance of base pairing for IM transfer is further illustrated by the effects of the double mutation of C11 and U12 in the stem, which results in considerable weakening of the secondary structure (Table 3), while leaving the critical A23 contact and other residues in the conserved region intact. This mutation, however, only marginally affects OM transfer (Fig. 2) or OM binding (Table 3), suggesting recognition of primary sequence, rather than secondary structure, by the OM receptor TAB. In fact, synthetic transcripts containing the nonanucleotide conserved motif but lacking significant secondary structure are transferred efficiently through the OM by the TAB-dependent pathway (13). Regardless of the precise nature of the interactions, it is evident from these experiments that distinct receptors exist for OM and IM transfer. This conclusion was reinforced by the resistance of IM transfer to antibody against TAB (Fig. 5).

Almost nothing is currently known about the nature of tRNA import factors. A major problem in systems such as leishmania, ciliates, and higher plants is the nonavailability of mutants with defective mitochondrial function. The use of oligonucleotide probes, coupled with submitochondrial fractionation, constitutes an alternative and general approach for the identification of the components of the membrane transport machinery. Previous experiments using intact tRNA molecules indicated the occurrence of nonproductive binding to mitochondrial membranes, with only about 10% of RNA bound to the OM being internalized (12); moreover, tRNA^{Gln} binds efficiently to the OM in a TAB-independent manner but is poorly imported (13). This occurrence of nonproductive binding makes it difficult to decide whether a particular tRNA binding membrane protein is an import factor or is unrelated to import. In contrast, binding of D arm oligonucleotides to the IM shows a strict correlation with importability. The most striking example of this correlation is the loss of binding as well as import in the G12-C mutant, and their simultaneous restoration by the second mutation, C13-G (Fig. 2 and 3 and Table 3). This should allow the identification of putative IM receptors by photochemical cross-linking and other methods.

The observed differential effects of the same mutation on OM and IM transfer support and refine the stepwise transport model (17) of mitochondrial tRNA import (Fig. 6). In this model, a tRNA species containing the conserved D arm import signal is proposed to bind to TAB at the OM and to be transferred in an ATP-dependent manner into the IMS, where it interacts with a different receptor or receptor complex on the IM. Subsequent translocation into the MX is driven by both the electrical (ΔΨ) and chemical (ΔpH) components of the proton motive force across the IM. In the in vitro system, the proton motive force is generated by F_{1}F_{0} ATPase-catalyzed ATP hydrolysis, with concomitant translocation of protons across the IM (Fig. 6).

Mitochondrial translation is dependent on the presence of an adequately balanced pool of different tRNA species in the MX. Independent evolution of OM and IM receptor specificities may be a means of ensuring this balance. Since the MX concentration of an individual species is dependent on the efficiencies of OM and IM transfer, a low efficiency of OM transfer would be compensated for by a high efficiency of IM transfer, and vice versa. There is also evidence that other domains of the D arm contain import signals. In tetrahymena, the anticodon of tRNAGln (UUG) functions as an import signal (20). Moreover, a tRNA^{Ile} derivative containing a nonfunctional D arm sequence from tRNA^{Gln} is nonetheless imported into leishmania mitochondria in vivo (10), indicating the presence of a signal elsewhere in the molecule, possibly in the anticodon arm, in addition to the one in the D arm. The presence of distinct OM and IM transport machineries would allow the sequential use of different signals on the same molecule for MX entry.

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