The J-related Segment of Tim44 Is Essential for Cell Viability: A Mutant Tim44 Remains in the Mitochondrial Import Site, but Inefficiently Recruits mtHsp70 and Impairs Protein Translocation

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Abstract. Tim44 is a protein of the mitochondrial inner membrane and serves as an adaptor protein for mtHsp70 that drives the import of preproteins in an ATP-dependent manner. In this study we have modified the interaction of Tim44 with mtHsp70 and characterized the consequences for protein translocation. By deletion of an 18-residue segment of Tim44 with limited similarity to J-proteins, the binding of Tim44 to mtHsp70 was weakened. We found that in the yeast Saccharomyces cerevisiae the deletion of this segment is lethal. To investigate the role of the 18-residue segment, we expressed Tim44D18 in addition to the endogenous wild-type Tim44. Tim44D18 is correctly targeted to mitochondria and assembles in the inner membrane import site. The coexpression of Tim44D18 together with wild-type Tim44, however, does not stimulate protein import, but reduces its efficiency. In particular, the promotion of unfolding of preproteins during translocation is inhibited. mtHsp70 is still able to bind to Tim44D18 in an ATP-regulated manner, but the efficiency of interaction is reduced. These results suggest that the J-related segment of Tim44 is needed for productive interaction with mtHsp70. The efficient cooperation of mtHsp70 with Tim44 facilitates the translocation of loosely folded preproteins and plays a crucial role in the import of preproteins which contain a tightly folded domain.

Key words: mitochondria • inner membrane • protein translocation • Tim44 • Hsp70

The biogenesis of mitochondria requires the import of nuclear-encoded preproteins from the cytosol. The translocation of preproteins across the mitochondrial membranes is mediated by Tom proteins in the mitochondrial outer membrane and by Tim proteins in the inner membrane (Pon and Schatz, 1991; Ryan and Jensen, 1995; Schatz, 1996; Neupert, 1997; Pfanner et al., 1997; Pfanner and Meijer, 1997). Tim44 was the first Tim protein identified (Maarse et al., 1992; Scherer et al., 1992). It behaves like a peripheral inner membrane protein and is exposed to the matrix (Blom et al., 1993; Rassow et al., 1994). Tim44 was found as a component of complexes together with the integral membrane proteins Tim23 and Tim17 (Berthold et al., 1995; Blom et al., 1995; Bömer et al., 1997) which are putative components of a protein translocation channel (Bauer et al., 1996; Dekker et al., 1997; Jensen and Kinnally, 1997; Lohret et al., 1997; Rassow et al., 1998; Rassow et al., 1999).

Recent efforts to determine the function of Tim44 have concentrated on the finding that Tim44 binds to mitochondrial Hsp70 (mtHsp70) in a 1:1 complex (Kronidou et al., 1994; Rassow et al., 1994; Schneider et al., 1994). Cells combining mutations in the genes encoding these proteins show synthetic growth defects (Rassow et al., 1994). mtHsp70 is essential for driving preproteins across the membranes into the matrix (Kang et al., 1990) and seems to constitute the motor unit of mitochondrial protein import (Pfanner and Meijer, 1995; Rassow and Pfanner, 1995; Schatz, 1996). Nucleotides and cochaperones are involved in the complex formation between both proteins. mtHsp70 initially binds to Tim44 in the ATP-bound state,
but the ATP is rapidly hydrolyzed (von Ahausen et al., 1995; Hörst et al., 1996; Schneider et al., 1996; Ungermann et al., 1996). The interaction with nucleotides is modulated by Mge1, the mitochondrial homologue of the prokaryotic heat shock protein GspE. Mge1 is associated with the Tim44 complex in substoichiometric amounts, acts as a nucleotide release factor, and is required for the efficient function of the import motor (Nakai et al., 1994; Voos et al., 1994; Laloraya et al., 1995; Westermann et al., 1995; Dole and Georgopoulos, 1996; Schneider et al., 1996; Dekker and Pfanner, 1997; Dole et al., 1997a; Miao et al., 1997). Mdj1p and Mdj2p, the mitochondrial homologues of prokaryotic DnaJ, are not associated with the Tim44 complex and there is no indication of an involvement in preprotein translocation (Rowley et al., 1994; Westermann et al., 1996; Dole and Neupert, 1997).

The binding site of Tim44 for mthsp70 is not known. Therefore, it is interesting to note that the sequence of Tim44 contains a short stretch of 18 amino acids (residues 185–202) which shows some similarity to a part of the J-domain which is characteristic of DnaJ and related modulators of Hsp70 proteins (Silver and Way, 1993; Cyr et al., 1994; Rassow et al., 1994; Hartl, 1996; Bukau and Horwich, 1998; Cheetham and Caplan, 1998; Kelley, 1998). This observation is reminiscent of several systems in which J-domains are the structures by which Hsp70s are bound to partner proteins and customized for specific functions (Rassow et al., 1995). It is tempting to speculate that the 18 residues of the J-similarity region of Tim44 may contribute to the interaction with mthsp70. In this case Tim44 could function in analogy to Sec63p, the membrane protein of the ER membrane which binds BiP and thereby plays an important role in the transport of proteins into this organelle (Scidmore et al., 1993; Schlenstedt et al., 1995; Brodsky, 1996; Rapoport et al., 1996; Coris and Schekman, 1997). However, the similarity of Tim44 to J-domains is very limited, raising the question of whether the 18-residue segment in the sequence of Tim44 is of functional relevance for the protein.

Similarly unclear is the role of Tim44 in the mechanism of mthsp70-driven protein import. A complete inactivation of Tim44 retains the mitochondrial protein import channels intact but causes defects in the translocation of the mature parts of matrix-targeted preproteins (Bömer et al., 1998). Therefore, it is possible that the only function of Tim44 is the cooperation with mthsp70. Since up to now no method was available to specifically modulate the interaction between Tim44 and mthsp70, suggestions on the involvement of Tim44 in the mechanism of protein transport were often based on circumstantial evidence or referred to studies using temperature-sensitive mutants of mtHsp70. In these mutants, the interactions between Tim44 and mthsp70 are impaired, but additional defects and thus indirect effects could not be ruled out. In fact, there is evidence that the mitochondrial Tim machinery contains two binding sites for mthsp70, one site directly at Tim44 and a second binding site at the Tim23/Tim17 complex (Rassow et al., 1995; Voos et al., 1996; Bömer et al., 1997). Therefore, it was unclear whether the defects in mitochondrial protein import which are observed with the sslc1-mutant strains are due to the impaired interaction of mthsp70 with Tim44 or due to other functions of mtHsp70.

In this study, we asked whether residues 185–202 of Tim44 are required for the function of Tim44. We found that Tim44 lacking this segment (Tim44185–202) is unable to substitute for authentic Tim44 in a yeast strain lacking the wild-type Tim44 gene although Tim44185–202 is correctly imported into mitochondria and assembled into the inner membrane. Our results indicate that the 18-residue segment of Tim44 is required for the efficient interaction of Tim44 with mthsp70. Moreover, we find that this interaction is necessary for the efficient action of mthsp70 on translocating folded preproteins and thus for the full activity of the mitochondrial protein import motor.

Materials and Methods

Construction of Strains and Plasmids

For expression of Tim44 in vivo, a 2.7-kb HindIII fragment containing the Tim44 gene (M aarse et al., 1992) was cloned into YEpplac33 and YEpplac181 (Gietz and Sugino, 1988). Tim44-ura3 was constructed using the Promega A Itered Sites System. The 2.7-kb HindIII fragment was cloned into pSELECT1 (= pALTER-1) and mutagenized with the mutagenic oligonucleotide 5′CAGAGAGACTTACAACTGCAAGAAAGCAAACGTG3′. The S4-lp deletion was verified by DNA sequence analysis. The mutagenized HindIII fragment was subsequently cloned into YEpplac181 (as a multi-copy vector) or YCplac111 (as a single-copy vector), respectively. The following Saccharomyces cerevisiae strains were used: PK82 (MATa his4-722 lys2-53 ttp1 leu2-3,112; Gambill and Egelman, 1993), PK81 (MATa ade2-101 lys2-53 leu2-3,112 ttp1 scp1-2 U E2; Gambill et al., 1993), MB3 (MATa ade2-101 his3-J200 leu2-11 lys2-801 ura3-1; M aarse et al., 1992), and MB20 (MATa ade2- trp1-1 ura3-1 leu2-44 Tim44: Y ES3 + YEplac33:Tim44(URA3); this study). For plasmid shuffling, double transformed cells were grown in rich liquid broth (YPD) and then plated on solid medium containing 5-fluoro-orotic acid according to Boeke et al. (1987).

Import of Preproteins into Isolated Mitochondria, Cross-linking by EGS, and Blue Native Electrophoresis

The cDNA constructs encoding mitochondrial preproteins were cloned into pGEM4 (Promega), the transcription was performed using SP6 RNA-polymerase (Stratagene), and the precursor proteins were subcloned into pGEM4 (Promega). For expression of Tim44 in vivo, a 2.7-kb HindIII fragment containing the Tim44 gene (M aarse et al., 1992) was cloned into YEpplac33 and YEpplac181 (Gietz and Sugino, 1988). Tim44-ura3 was constructed using the Promega A Itered Sites System. The 2.7-kb HindIII fragment was cloned into pSELECT1 (= pALTER-1) and mutagenized with the mutagenic oligonucleotide 5′CAGAGAGACTTACAACTGCAAGAAAGCAAACGTG3′. The S4-lp deletion was verified by DNA sequence analysis. The mutagenized HindIII fragment was subsequently cloned into YEpplac181 (as a multi-copy vector) or YCplac111 (as a single-copy vector), respectively. The following Saccharomyces cerevisiae strains were used: PK82 (MATa his4-722 lys2-53 ttp1 leu2-3,112; Gambill and Egelman, 1993), PK81 (MATa ade2-101 lys2-53 leu2-3,112 ttp1 scp1-2 U E2; Gambill et al., 1993), MB3 (MATa ade2-101 his3-J200 leu2-11 lys2-801 ura3-1; M aarse et al., 1992), and MB20 (MATa ade2- trp1-1 ura3-1 leu2-44 Tim44: Y ES3 + YEplac33:Tim44(URA3); this study). For plasmid shuffling, double transformed cells were grown in rich liquid broth (YPD) and then plated on solid medium containing 5-fluoro-orotic acid according to Boeke et al. (1987).
Fractionation of Mitochondria

The procedures of protease treatment, swelling of mitochondria, and carbonate extraction were published previously (Blom et al., 1993; Rassow et al., 1994). The tendency of mitochondrial proteins to form aggregates was tested by lysis of the organelles and subsequent centrifugation. Mitochondria from the wild-type and from the strain expressing Tim44 was tested by lysis of the organelles and subsequent centrifugation. The procedures of protease treatment, swelling of mitochondria, and carbonate extraction were published previously (Blom et al., 1993; Rassow et al., 1994). The tendency of mitochondrial proteins to form aggregates was tested by lysis of the organelles and subsequent centrifugation. Mitochondria from the wild-type and from the strain expressing Tim44 was tested by lysis of the organelles and subsequent centrifugation.

Assessment of the Mitochondrial Membrane Potential

The membrane potential (ΔΨ) of isolated yeast mitochondria was determined by recording the fluorescence decrease of the voltage-sensitive dye 3,3′-dipropylthiacyclobacarboxylic iodine [DiSC3(5); Molecular Probes; Sims et al., 1974]. The assays were performed using a Perkin-Elmer 640-40 fluorescence spectrometer at 25°C (excitation at 622 nm, slit width 5 nm). The mitochondria (100 μg protein) were incubated in 1 ml of 0.6 M sorbitol, 0.1% (vol/vol) Triton X-100, 1 mM EDTA, 0.5 mM PMSF, pH 7.4. The final concentration of the fluorescent dye DiSC3(5) was 3.6 μM. The membrane potential was dissipated by the addition of 3 mM (final concentration) KCN. The difference between the fluorescences before and after the addition of KCN represents a rough assessment to the membrane potential.

For coimmunoprecipitations, mitochondria (25 μg/sample) were lysed in 200 μl 250 mM sucrose, 80 mM KCl, 20 mM MOPS-KOH, pH 7.2, 0.1% (vol/vol) Triton X-100, 5 mM EDTA, 0.5 mM PMSF. In the experiment shown in Fig. 6 F, the mitochondria were lysed in 150 mM NaCl, 0.1% Triton X-100, 0.5 mM PMSE, 10 mM Tris- HCl, pH 7.4. After a spin of 16,000 g for 10 min, the lysates were incubated with antibodies directed against mtHsp70. The antibodies were covalently coupled to protein A-Sepharose as described previously (Vooos et al., 1994).

Figure 1. Deletion of residues 185–202 in Tim44 is lethal for S. cerevisiae. (A) The J-similar segment of Tim44. Shown is a comparison of the amino acid sequence of S. cerevisiae Tim44 residues 185–202 with sequences in E. coli DnaJ, S. cerevisiae Sis1p, and Sec63p. Dark boxes are identical residues; white boxes are similar amino acids. Secondary structure predictions were performed according to Garnier and Robson using the program Protean. Indicated is the number of the residues in the corresponding amino acid sequences of the compared proteins. Serpentine lines are the predicted α helix; dark dots are predicted unstructured amino acids (turn or coil). Helix I and helix III of DnaJ were defined according to the NMR structure. (B) The segment of residues 185–202 in Tim44 is essential for viability of yeast. Strain M B20 contains a deletion of chromosomal TIM44 sequences and harbors the multicopy plasmid YEp plac33 (withURA3 as a selective marker) encoding authentic Tim44. (Left plate) M B20 was transformed with the LEU2 marked 2 μm plasmid YEp plac181 carrying the wild-type TIM44 gene (upper half of the plate) or tim44Δ18 (lower part of the plate). (Right plate) Double transformants were grown on nonselective rich broth (YPD) and then plated on solid medium containing glucose as a carbon source and 5-fluoro-otic acid (FOA) for selection of cells which have lost the URA3 marked expression plasmids. The cells retaining YEp plac181 containing the authentic TIM44 coding sequence grew normally (upper half of the plate). Cells retaining only the Tim44Δ18 encoding construct were not viable (lower half of the plate).
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Results

A Segment of Tim44 Essential for Viability of S. cerevisiae

The region of sequence similarity between Tim44 and characteristic J-domains is shown in Fig. 1 A. The similarity extends from residue 185 to residue 202 and is depicted for the two yeast proteins Sec63p (Sadler et al., 1989; Feldheim et al., 1992) and Sis1p (Luke et al., 1991; Zhong and Arndt, 1993), and for the Escherichia coli protein DnaJ (Ohki et al., 1986; Liberek et al., 1991). The structure of the J-domain of DnaJ has been determined by NMR spectroscopy (Szyperski et al., 1994; Hill et al., 1995). The J-similarity segment of Tim44 corresponds to α helix II of DnaJ (residues 18–30) and the first half of the following turn region. In DnaJ, the turn is followed by helix III comprising residues 41–55. The secondary structures which are predicted for the corresponding part of Tim44 similarly show two hydrophilic α helices which are connected by a putative turn element. However, it should be emphasized that the similarity of Tim44 to J-domains is still very limited. Tim44 obviously does not belong to the family of homologous J-proteins.

To determine the relevance of the J-similar segment for the function of Tim44 in vivo, we constructed a plasmid encoding a Tim44 with a deletion of residues 185–202 (Tim44_{Δ18}). We then tested whether Tim44_{Δ18} can substitute for authentic Tim44 in a genetic assay (Fig. 1 B). In a strain of S. cerevisiae expressing both forms of Tim44 from different plasmids, we used the URA 3/FOA technique to deplete the authentic Tim44. It turned out that the cells lacking the authentic Tim44 were not viable. Tim44_{Δ18} could not substitute for wild-type Tim44, and neither on glycerol nor on glucose was any growth observed. Thus, this result demonstrates that the 18-residue segment of Tim44 is essential for viability of yeast.

Tim44 Lacking Residues 185–202 Is Imported into Mitochondria and Reaches Its Functional Location

The lethality of the 18-residue deletion could be caused by two different reasons. Either the 18-residue segment is crucial for the biogenesis of Tim44, or this segment is required for the function of Tim44 within the mitochondria. To address the first possibility, we expressed Tim44_{Δ18} in reticulocyte lysate and tested whether the protein could be imported into isolated mitochondria. In parallel we imported the authentic Tim44 (Fig. 2 A). Both preproteins were processed to a mature form (Fig. 2 A, lanes 1 and 5) and translocated into a location protected against externally added proteinase K (lanes 3 and 7). The import was dependent on the mitochondrial membrane potential, confirming the specificity of the reaction. The result demonstrates that the 18-residue segment of Tim44 is not required for efficient transport into mitochondria in vitro.

To test whether Tim44_{Δ18} is similarly imported into vivo,
Tim44 was expressed in a S. cerevisiae wild-type strain from a multi-copy vector, and mitochondria were isolated to determine the amount of imported protein (Fig. 2 A, lanes 9 and 10). We found that the mitochondria had imported both the authentic Tim44 and Tim44\textsubscript{18}. The amount of authentic Tim44 was not significantly reduced as compared with the original wild-type strain; Tim44\textsubscript{18} was overexpressed about threefold. The location of the imported Tim44 proteins was confirmed by a fractionation experiment. Both proteins were resistant against externally added trypsin in intact mitochondria (Fig. 2 B, lane 2) and after opening of the intermembrane space by swelling (Fig. 2 B, lane 5). Only after disruption of the inner membrane by sonication did the proteins become accessible for the protease (Fig. 2 B, lanes 3 and 6), in agreement with results which were obtained previously for the wild-type Tim44 (Blom et al., 1993; Rassow et al., 1994). A faster lysis of the mitochondria by detergent, both proteins were similarly soluble (Fig. 2 C). Even after prolonged incubation at 0 or 25°C, no formation of aggregates was observed (Fig. 2 C, lanes 2 and 4). We then tested whether the deletion of the 18-residue segment may affect the association of Tim44 with the inner membrane. The extraction of Tim44 and Tim44\textsubscript{18} was monitored by sodium carbonate and again both proteins showed the same behavior (Fig. 2 D). In contrast to the ADP/ATP carrier (AAC) which was resistant against this treatment, both Tim44 proteins were extracted. The association of Tim44 with the inner membrane was also determined in the presence of different salt concentrations (Fig. 2 E). Mitocondria were sonicated in

Figure 2. Tim44\textsubscript{18} is efficiently imported into mitochondria. (A) Import of Tim44\textsubscript{18} in vitro and in vivo. Lanes 1–8 show the import of 35S-labeled proteins into isolated mitochondria. Wild-type Tim44 and Tim44\textsubscript{18} (containing a deletion of residues 185–202) were synthesized in reticulocyte lysate and incubated with mitochondria isolated from a S. cerevisiae wild-type strain. Samples 2, 4, 6, and 8 contained valinomycin to dissipate the membrane potential (Δψ). The mitochondria were reisolated and treated with proteinase K (Prot. K) as indicated. The imported proteins were analyzed by SDS-PAGE and fluorography. p, precursor protein; m, mature protein. Lanes 9 and 10 show the import of Tim44 and Tim44\textsubscript{18} in vivo. The multi-copy plasmid YEpplac181 encoding Tim44\textsubscript{18} was transformed into the yeast wild-type strain MB3. Mitocondria were isolated to determine the expression and import of Tim44\textsubscript{18} and of the authentic Tim44. The mitochondrial proteins were separated by SDS-PAGE and analyzed by Western blotting. Tim44 and mThsp70 were visualized by specific antibodies. WT, wild-type mitochondria; WT + Tim44\textsubscript{18}, mitochondria from the strain MB3 expressing Tim44\textsubscript{18} in addition to the authentic Tim44. The same mitochondria as in lane 10 were analyzed in the experiments shown in B–E. (B) Fractionation of mitochondria. Mitocondria isolated from the yeast strain expressing both authentic Tim44 and Tim44\textsubscript{18} were swollen to open the intermembrane space (lanes 4–6) or sonified to open the matrix (lanes 3 and 6) and tested for the accessibility of Tim44 by trypsin. Both forms of Tim44 became accessible for the protease only after sonication. (C) Solubility of Tim44 after lysis of the mitochondria. Mitocondria containing authentic Tim44 together with Tim44\textsubscript{18} were lysed in the presence of 0.1% Triton X-100 and 80 mM KCl, and after a spin of 5 min at 16,000 g the supernatants were incubated for 1 h at 0 or 25°C, respectively. The samples were subsequently subjected to a centrifugation of 1 h at 100,000 g. Supernatants and pellets were separated and analyzed by SDS-PAGE and Western blotting. The blots were decorated with antibodies directed against Tim44 and mThsp70.
the presence of up to 500 mM KCl and the membranes were subsequently pelleted by centrifugation. Tim44 and Tim44$_{118}$ were found stably associated with the membranes; only minor amounts of both proteins were released at higher ionic strength. As a control, we followed the distribution of Mge1p which was soluble in all samples. The AAC was completely resistant against extraction.

Eventually, we investigated the involvement of both forms of Tim44 in the formation of high molecular weight complexes using the method of blue native electrophoresis (BN-PAGE; Schägger and von Jagow, 1991; Dekker et al., 1997). As shown in Fig. 2 F, Tim44 and Tim44$_{118}$ both showed the same distribution, suggesting that both proteins participate in the same interactions with the components of the Tim machinery. In a previous study we found that Tim44 is mainly associated with Tim23 (Bömer et al., 1997). Therefore, we also determined the complex formation of Tim23 and found that it showed the same running behavior in the BN-PAGE, irrespective if only the authentic Tim44 was present or if Tim44$_{118}$ was overexpressed in addition.

We conclude from these experiments that Tim44$_{118}$ is efficiently imported into mitochondria and acquires the correct topology at the inner side of the inner membrane. The results of the BN-PAGE indicate that Tim44$_{118}$ adopts the native folding state and engages in the same interactions with the Tim machinery as the authentic Tim44.

These conclusions are corroborated by the interactions of Tim44$_{118}$ with different forms of mtHsp70 (see below, Fig. 6 F).

**Tim44$_{118}$ Interacts with Mitochondrial Preproteins**

If Tim44$_{118}$ is recruited by the Tim machinery, is it also present at import sites during translocation of preproteins? We addressed this question by chemical cross-linking (Fig. 3). As a substrate we synthesized the hybrid protein Su9-DHFR, containing the presequence of subunit 9 of the mitochondrial ATP synthase fused to the complete DHFR. The import of this protein is dependent on the membrane potential $\Delta \psi$ (Fig. 3 A, lanes 1-4) and ATP (Fig. 3 A, lanes 5-8). A fter depletion of ATP, Su9-DHFR is accumulated in import sites as a membrane-spanning translocation intermediate. Following this protocol, Su9-DHFR was accumulated in mitochondria of both the Tim44$_{118}$ overproducing strain and the wild-type (Fig. 3 B, lanes 1 and 2). The translocation intermediates were cross-linked to the proteins in the vicinity by addition of the agent EGS. In wild-type mitochondria, two products were formed, corresponding to the precursor form and the processing intermediate of Su9-DHFR (Fig. 3 B, lane 4, bands labeled Tim44* and Tim44**; Blom et al., 1993). In mitochondria, which in addition contained Tim44$_{118}$, a third cross-linking product was formed which could be precipi-
tated by antibodies against Tim44 (Fig. 3 B, lane 7, band labeled Tim44***). No reaction product was precipitated by the preimmune serum. According to its size in the SDSPAGE, the additional product corresponds to cross-linking of Tim44<sub>18</sub> to the processing intermediate of Su9-DHFR. The cross-linking of a protein in transit across the mitochondrial membranes confirms that Tim44<sub>18</sub> is present at protein import sites.

**Coexpression of Tim44<sub>18</sub> Causes Membrane Potential-independent Defects in Protein Import**

To determine the possible role of the 18-residue segment in Tim44, additional preproteins were imported and tested for defects in distinct steps of translocation across the mitochondrial membranes. In a first series of experiments, the β subunit of the mitochondrial ATP synthase (F1β) was synthesized in reticulocyte lysate in the presence of [35S]methionine/[35S]cysteine and imported into mitochondria which were isolated from the Tim44<sub>18</sub>-overexpressing strain and the corresponding wild-type (Fig. 4 A). It is known from previous studies that the import of F1β is very sensitive against defects in the import machinery and requires the mtHsp70-dependent unfolding machinery of the mitochondria (Kang et al., 1990; Rassow and Pfanner, 1991; Gambill et al., 1993). We now observed only a slight reduction in processing of F1β, suggesting that the deletion of the 18-residue segment of Tim44 does not cause major changes in the insertion of the presequence into the Tim machinery of the inner membrane (Fig. 4 A, top). However, a protease-protection assay revealed a delay in the translocation of the mature part of the preprotein (Fig. 4 A, bottom), indicating that the deletion affected the completion of translocation. After an import time of 20 min the efficiency of translocation was reduced by 75–80% (Fig. 4 B, column 2 vs. column 3). Some reduction was also observed after overexpression of the authentic Tim44, but the effect was much less pronounced (Fig. 4 B, column 3, and see below, Fig. 5 C).

Modifications of mitochondrial inner membrane proteins can easily lead to a reduction of the mitochondrial membrane potential and thereby indirectly cause reduced efficiencies of protein transport. To address this possibility, we compared the membrane potential of the mitochondria which had been used in the previous experiments. A sensitive assay we determined the membrane potential-dependent uptake of the dye DiSC<sub>3</sub>(5) (Sims et al., 1974). The uptake is reversible and can be quantified by following the change in the fluorescence of the dye. With mitochondria from the Tim44<sub>18</sub> mutant strain no reduction in the membrane potential was observed (Fig. 4 C). An indirect effect of the mutation on mitochondrial protein import mediated by a weakened membrane potential can thus be excluded.

**Coexpression of Tim44<sub>18</sub> Leads to Reduced Import Efficiencies of Folded Protein Domains**

We then asked if the effect of the deletion of the 18-residue segment on protein import is dependent on the folding state of the preprotein. Previous studies have shown that the heme-binding domain of cytochrome b<sub>2</sub> is tightly folded and requires an intact mtHsp70 system to drive the unfolding of this domain (Glick et al., 1993; Voos et al., 1994). Figure 4. Translocation of a preprotein across the mitochondrial membranes is impaired in the presence of Tim44<sub>18</sub>. (A) Import of the β subunit of the mitochondrial ATP synthase (F1β). Mitochondria were isolated from the strain overexpressing Tim44<sub>18</sub> in addition to the authentic Tim44 (WT + Tim44<sub>18</sub>) and the corresponding wild-type strain MB3 containing only the authentic Tim44 (WT). 35S-labeled F1β was synthesized in reticulocyte lysate and incubated with isolated mitochondria at 25°C for the times indicated. In samples 4 and 8 the membrane potential (Δψ) was dissipated by addition of valinomycin. After incubation at 25°C, the samples were cooled to 0°C and divided into halves. One-half was treated with 50 μg/ml proteinase K for 20 min (+ Prot. K), the other half was left without protease (− Prot. K). A fter incubation with PM SF, the samples were analyzed by SDSPAGE and digital autoradiography using a PhosphorImager system. (B) Import of F1β into mitochondria isolated from the wild-type (column 1), in mitochondria isolated from the strain expressing Tim44<sub>18</sub> in addition to authentic Tim44 (column 2), and in mitochondria from the strain overexpressing the authentic Tim44 (column 3). F1β was imported for 20 min at 25°C following the protocol described in A. The amounts of processed and protease-protected F1β were quantified using the PhosphorImager. The value obtained for import into wild-type mitochondria was set to 100% (control). (C) A assessment of the mitochondrial membrane potential. The membrane potential (Δψ) was determined at 25°C, using the fluorescence dye 3,3′-dipropylthiadi-carbocyanine iodide, DiSC<sub>3</sub>(5). The Δψ is indicated by the difference in fluorescence before and after addition of potassium cyanide (KCN).
Proteins were synthesized in reticulocyte lysate and incubated with residues of cytochrome b$_2$. The preproteins of Tim44 were expressed in the strain MB3 using the vector pBluescript II SK(+) carrying the heme-binding domain (HB; position 81–179), fused to DHFR. In both constructs, a part of the sorting signal (residues 47–65) is deleted within the presequence. (A) The hybrid protein pb220-DHFR contains the first 220 amino acid residues of cytochrome b$_2$, including the complete heme-binding domain (HB; position 81–179), fused to DHFR. In pb167-DHFR, the first 167 amino acid residues of cytochrome b$_2$ (lacking the carboxy-terminal part of the heme-binding domain) are fused to DHFR. In both constructs, a part of the sorting signal (residues 47–65) is deleted within the presequence. (B) Import of pb220-DHFR and pb167-DHFR. The radiolabeled preproteins were synthesized in reticulocyte lysate and incubated with mitochondria isolated from the Tim44-overproducing strain (WT + Tim44$_{18}$) subjected to the same strain as applied in the experiments shown in Figs. 2–4) or the corresponding wild-type strain M B3 (WT). Lanes 4 and 8 contained valinomycin to dissipate the membrane potential. Subsequently, all samples were treated with proteinase K. The imported proteins were analyzed by SDS-PAGE and fluorography. (C) Ratio of the import efficiencies of pb220-DHFR and pb167-DHFR after overexpression of Tim44$_{18}$ and after overexpression of authentic Tim44. Both versions of Tim44 were expressed in the strain M B3 using the vector YEplac181 (LEU2). The preproteins pb220-DHFR and pb167-DHFR were imported as described in the legend to Fig. 4 B and quantified for calculation of the import efficiencies (imported protein in percentage of total preprotein added). Shown are the ratios of the efficiencies of both proteins after 5 or 10 min of import at 25°C. The experiments shown in Figs. 2 and 3 have indicated that the interactions of Tim44$_{18}$ with the Tim machinery are not disturbed by the deletion of the 18-residue segment. We now ask if Tim44$_{18}$ shows an altered interaction with mtHsp70. We lysed mitochondria from the wild-type in the presence of detergent and performed coimmunoprecipitations using specific antibodies raised against mtHsp70. The precipitates were analyzed by immunoblotting and demonstrated the association of mtHsp70 with Tim44$_{18}$. The precipitates were analyzed by immunoblotting and demonstrated the association of mtHsp70 with Tim44$_{18}$.

**Deletion of the 18-Residue Segment Causes Reduced Complex Formation of Tim44 with mtHsp70**

The experiments shown in Figs. 2 and 3 have indicated that the interactions of Tim44$_{18}$ with the Tim machinery are not disturbed by the deletion of the 18-residue segment. We now ask if Tim44$_{18}$ shows an altered interaction with mtHsp70. We lysed mitochondria from the wild-type in the presence of detergent and performed coimmunoprecipitations using specific antibodies raised against mtHsp70. The precipitates were analyzed by immunoblotting and demonstrated the association of mtHsp70 with Tim44$_{18}$. The precipitates were analyzed by immunoblotting and demonstrated the association of mtHsp70 with Tim44$_{18}$.

DHFR is about two times more reduced as compared with the import of pb167-DHFR, demonstrating that the deletion of the 18-residue segment mainly affects the import of the preprotein containing the tightly folded heme-binding domain.
Figure 6. Deletion of the 18-residue segment of Tim44 reduces complex formation with mtHsp70. (A) Coimmunoprecipitation of Tim44 and Tim44_{18} with antibodies directed against mtHsp70. Mitochondria were lysed in the presence of detergent and protein complexes were precipitated during an incubation of 1 h by antibodies raised against mtHsp70. The precipitates were analyzed by SDSPAGE, Western blotting, and immunostaining, using polyclonal antisera raised against Ssc1p (Hsp70) or Tim44. Sample 1 is total protein of wild-type mitochondria (10 μg protein). Sample 2 is Tim44-mtHsp70 complex precipitated from wild-type mitochondria (100 μg protein). (B) Sample 1 is total protein (10 μg protein); sample 2 is complex precipitated from mitochondria (100 μg protein) of the strain coexpressing Tim44_{18} together with authentic Tim44. (C) Samples 1 and 2 are total protein (10 μg) of mitochondria from the same strain as in B. To compare the stability of Tim44_{18} and Tim44 in the lysis buffer, the proteins of sample 1 were precipitated by trichloroacetic acid immediately after lysis of the mitochondria; the proteins of sample 2 were precipitated after the lysed mitochondria had been incubated for 1 h at the same temperature as the coimmunoprecipitations. Samples 1 and 3 are similar to the samples of B, except that the lysis buffer contained 80 mM KCl instead of 10 mM KCl. The different stability of the complexes formed between mtHsp70 and Tim44 versus mtHsp70 and Tim44_{18} is indicated by the ratio of the bands corresponding to both forms of Tim44 in the immunoprecipitation (lane 3) in comparison to the mitochondria (lanes 1 and 2). (D) Ratio of the amounts of Tim44_{18} and Tim44 as determined by quantifications of coimmunoprecipitations by antibodies against mtHsp70 in comparison to the ratio of both proteins in total mitochondria (as shown in B and C). Columns 1 and 2 are coimmunoprecipitation in the presence of 10 mM KCl; columns 3 and 4 are coimmunoprecipitation in the presence of 80 mM KCl. (E) Release of Tim44 and Tim44_{18} from mtHsp70 in the presence of ATP. The complex of Tim44 and Tim44_{18} with mtHsp70 was precipitated by protein A-Sepharose-bound antibodies against mtHsp70 (−ATP). Parallel samples were incubated in the presence of 2 mM ATP and 5 mM MgCl₂ (+ATP). The protein A-Sepharose was separated from the supernatant by centrifugation and the bound complex was eluted at pH 2.5. The proteins were precipitated by trichloroacetic acid and analyzed by SDS-PAGE and immunoblotting. The relative amounts of coprecipitated Tim44 and Tim44_{18} were quantified; the combined amounts of bound and free protein were set to 100% (total). Pellet is the coimmunoprecipitated protein (released from the protein A-Sepharose at pH 2.5); Sup. is protein in the supernatant of the coimmunoprecipitation. (F) Interaction of Tim44 and Tim44_{18} with wild-type mtHsp70 and with mutant mtHsp70 encoded by the allele ssc1-2. Tim44_{18} was expressed from a single-copy vector in the strain ssc1-2; mitochondria were isolated (containing both the authentic Tim44 and Tim44_{18}), lysed, and analyzed by coimmunoprecipitation using the antibodies raised against mtHsp70. Mitochondria from a strain containing wild-type mtHsp70 were analyzed in parallel (WT). Shown are the relative amounts of Tim44 and Tim44_{18}, respectively. The relative amounts of coprecipitated protein obtained in the strain expressing wild-type mtHsp70 were set to 100%. To determine the affinity of both forms of mTsp70 to a preproteins, Su9-DHFR was imported into mitochondria of the strain ssc1-2 and the corresponding wild-type strain. The imported Su9-DHFR was coimmunoprecipitated by the antibodies raised against mTsp70. A fiter quantification, the amount of coprecipitated Su9-DHFR were calculated relative to the amounts of imported Su9-DHFR. The ratio found for binding of Su9-DHFR to Ssc1-2p was set to 100%.

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contented about three- to fourfold more Tim44_{18} than Tim44 (Fig. 6B, lane 1, and Fig. 6C, lane 1). By coimmunoprecipitates from these lysates we compared the association of mtHsp70 with Tim44_{18} and the authentic Tim44. The ratio of Tim44_{18} to Tim44 in the precipitates was close to 1:1, demonstrating that complex formation of Tim44_{18} to mtHsp70 was reduced about three- to fourfold by the deletion of the 18-residue segment (Fig. 6B, lane 2, and Fig. 6C, lane 3). Both forms of Tim44 were stable upon prolonged incubation after lysis, confirming that the reduced amount of Tim44_{18} in the immunoprecipitates was due to a reduced complex formation with mtHsp70 (Fig. 6C, lanes 1 and 2).

The reduced affinity of mTsp70 to Tim44_{18} as compared with the authentic Tim44 was confirmed by systematic quantifications (Fig. 6D). Complex formation of mTsp70 with Tim44_{18} was reduced by ~70%. In the presence of ATP, Tim44_{18} and Tim44 were both effi-
Josephson et al. (1998) have shown that su8-DHFR is more than fourfold higher than that of mtHsp70 with both Tim44 proteins and a substrate protein by coimmunoprecipitations (Fig. 6 F). While the efficiency of binding of the mutant mtHsp70 to the substrate protein Su9-DHFR was more than fourfold higher than that of wild-type mtHsp70 (Fig. 6 F, columns 5 and 6), the association of the mutant mtHsp70 with both Tim44 and Tim44,18 was blocked (Fig. 6 F, columns 2 and 4). This result implies that not only the authentic Tim44 but also the truncated Tim44,18 is recognized by mtHsp70 as a partner protein of special properties, and not as a substrate protein.

Discussion
In this study we have characterized the role of complex formation of mtHsp70 with Tim44 in mitochondrial protein import. A system of reduced binding between both proteins was created by the deletion of an 18-residue segment in Tim44 (residues 185–202) which shows a limited similarity to one of the two α helices of J-domains. The intracellular localization of Tim44,18 is not altered by the deletion. In all fractionation experiments Tim44,18 showed the same behavior as the authentic Tim44, and by chemical cross-linking we found that Tim44,18 is localized at the inner membrane protein import sites.

The correct topology of Tim44,18 within the mitochondria allowed us to test whether the 18-residue segment of Tim44 is required to recruit mtHsp70 to the Tim machinery of the inner membrane. We found that binding of mtHsp70 to Tim44,18 was reduced by ~70% as compared with the authentic Tim44. The 18-residue segment of Tim44 is obviously not the only site for binding to mtHsp70. In contrast to the integral membrane protein Sec63p which is exposed to the ER lumen only by short segments of its sequence (Feldheim et al., 1992), Tim44 is a peripheral protein and in larger parts exposed to the matrix (Blom et al., 1993; Rassow et al., 1994). This topology may allow the formation of multiple binding sites.

While this manuscript was in preparation, a publication appeared by Greene et al. (1998) showing that the binding site of DnaJ for DnaK is the helix II of the J-domain, which corresponds exactly to the segment of similarity to Tim44. Since Tim44 seems not to belong to the family of J-proteins (Wada and Kanwar, 1998) we assume that the J-related segment of Tim44 does not represent a J-homology in the strict sense but rather a J-analogous development to facilitate the interaction with mtHsp70. According to Greene et al. (1998), the helix II of DnaJ interacts with the ATPase domain of DnaK. Following the analogy between Tim44 and DnaJ, Tim44 should similarly bind to the ATPase domain of mtHsp70. However, other J-proteins were found to interact with the carboxy-terminal domain of Hsp70s (Freeman et al., 1995; Demand et al., 1998) or to require both domains for binding (Ungewickell et al., 1997). Therefore, it may be speculated that the interaction between mtHsp70 and Tim44 is mediated by multiple attachment sites, as was shown recently by the x-ray structure for the complex of DnaK with GrpE (Harrison et al., 1997). We cannot completely rule out allosteric effects of the deletion of the 18-residue segment. However, the only difference to the wild-type protein we observed was restricted to the interaction with mtHsp70. The very sensitive assays of chemical cross-linking (Fig. 3 B) and blue native electrophoresis (Fig. 2 F) demonstrate that the oligomeric state of Tim44, and the direct interactions with preproteins and other components of the Tim machinery were retained. The comparison to the DnaJ-DnaK complex as analyzed by Greene et al. (1998) suggests that the 185–202 segment of Tim44 provides the major binding site for mtHsp70.

Several data indicate that Tim44 binds to Tim23 and provides a dynamic link between the Tim proteins which form the protein import channel and the soluble mtHsp70 system of the matrix (Bömer et al., 1997; Dekker et al., 1997). With Tim44,18 the function of this link is specifically impaired in the interactions of Tim44,18 with mtHsp70. Our import experiments demonstrate that the presence of Tim44,18 causes a significant reduction in the import efficiencies of different preproteins, including proteins which are regarded as loosely folded. The import of all of these preproteins is strictly dependent on mtHsp70 as demonstrated by previous studies using temperature-sensitive strains of SSC1 (encoding mtHsp70) (Gambill et al., 1993; Voo s et al., 1993). The strongest inhibition of import was observed with preproteins which contain a tightly folded domain. Such domains cause restrictions in the translocation across the mitochondrial membranes which are due to the requirement of unfolding within the import channel (Gambill et al., 1993; Glück et al., 1993; Voo s et al., 1993, 1996; Matouschek et al., 1997). To overcome these restrictions, the mtHsp70 system of the matrix has to exert a force on the translocating protein which is sufficient to pull the protein across the membranes.

Studies to elucidate the mechanism by which this force is generated made use of the ssc1-2 mutant of mtHsp70 (Kang et al., 1990; Schneider et al., 1994; von Ahsen et al., 1995; Voos et al., 1996). The mtHsp70 of this mutant binds efficiently to translocating preproteins but is impaired in binding to Tim44. This defect correlates with an inhibition in the import of tightly folded protein domains. However, conclusions could only be drawn with reservation. The Tim machinery seems to contain at least two binding sites for mtHsp70, one at Tim44 and a second site at the Tim23/Tim17 import channel, and both interactions are inhibited by the ssc1-2 mutation (Bömer et al., 1997). The principle which governs the mechanism of mtHsp70-dependent protein import is still unknown. A Brownian ratchet mechanism (Simon et al., 1992; Ungermann et al., 1994; Gaume

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et al., 1998) and a mechanism of mtHsp70/Tim44-mediated pulling (Glick, 1995; Pfanner and Meijer, 1995) have been suggested. In this context it is remarkable that the effect of Tim44ΔΔ on the import of different preproteins and on the viability of yeast resembles the effects of ssc1-2. This similarity in the phenotype thus corroborates and specifies the concept that the cooperation of Tim44 with mtHsp70 is of particular importance in the import of tightly folded protein domains. In a previous study on a complete inactivation of functional Tim44 in isolated mitochondria we showed that Tim44 acts at the inner side of the inner membrane (Bömer et al., 1998). The results obtained with the Tim44ΔΔ construct suggest that in this location the functions of Tim44 in protein import may be confined to specific interactions with mtHsp70.

In summary, the results of this study demonstrate that the J-related segment of Tim44 (residues 185-202) is required for the essential functions of Tim44 in mitochondria. This segment is not the only element involved in the interaction of Tim44 with mtHsp70, but it is required for productive cooperation of both proteins and the optimal efficiency of mitochondrial protein import. mtHsp70 is an essential motor protein in the translocation of all proteins which are imported into the mitochondrial matrix, irrespective of whether or not they contain tightly folded domains (Schatz, 1996; Neupert, 1997; Pfanner et al., 1997). In contrast, the requirement for an interaction of mtHsp70 with Tim44 seems to be less strict and appears to play an important role primarily in situations which require the full activity of the import motor, for example in overcoming stronger restrictions in the translocation of preproteins. The import of loosely folded preproteins is facilitated by Tim44, but the effect is much more pronounced in the case of tightly folded domains.

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