Biogenesis of a mitochondrial outer membrane protein in *Trypanosoma brucei*: targeting signal and dependence on an unique biogenesis factor

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Running title: biogenesis of signal-anchored POMP10

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**ABSTRACT**

The mitochondrial outer membrane (OM) contains single and multiple membrane-spanning proteins which need to contain signals that ensure correct targeting and insertion into the OM. The biogenesis of such proteins has so far essentially only been studied in yeast and related organisms. Here we show that POMP10, an OM protein of the early diverging protozoan *Trypanosoma brucei*, is signal-anchored. Transgenic cells expressing variants of POMP10 fused to GFP demonstrate that the N-terminal membrane-spanning domain flanked by a few positively charged or neutral residues is both necessary and sufficient for mitochondrial targeting. Carbonate extraction experiments indicate that, while the presence of neutral instead of positively charged residues did not interfere with POMP10 localization, it weakened its interaction with the OM. Expression of GFP-tagged POMP10 in inducible RNAi cell lines shows that its mitochondrial localization depends on pATOM36 but does neither require Sam50 nor ATOM40, the trypanosomal analogue of the Tom40 import pore. pATOM36 is a kinetoplastid-specific OM protein that has previously been implicated in the assembly of OM proteins and in mitochondrial DNA inheritance. In summary, our results show that while the features of the targeting signal in signal-anchored proteins are widely conserved, the protein machinery that mediates their biogenesis is not.

Mitochondria perform many important functions and are essential for all eukaryotes (1). Whereas a small number of proteins are synthesized in the organelle, more than 95% of the mitochondrial proteome is nuclear-encoded, synthesized in the cytosol and subsequently imported into mitochondria (2-4). Mitochondria consist of four compartments: the outer and the inner membrane that surround the soluble intermembrane space (IMS) and the matrix, respectively. Thus, nuclear-encoded proteins not only need to be targeted to mitochondria but also sorted to their correct intra-mitochondrial destination. The mitochondrial outer membrane (OM) is of special interest. It builds the interface between the organelle and the cytosol and forms a barrier across which all communication between the organelle and its surroundings must occur (5-7). Mitochondrial OM proteins mediate apoptosis, fission and fusion, interaction with other organelles as well as transport of metabolites and precursor proteins. Whereas mitochondrial protein import in general has been extensively studied, we still have large gaps in the understanding
of the biogenesis of mitochondrial OM proteins (8-11).

Integral OM proteins are either anchored by a transmembrane β-barrel or by single or multiple α-helices. The single membrane-spanning proteins can further be categorized into N-terminal (signal)-anchored, internally anchored or C-terminal (tail)-anchored proteins. OM proteins generally contain internal targeting signals. For β-barrel proteins the OM targeting signal appears to be a dedicated β-hairpin motif (12), whereas for single membrane-spanning proteins the targeting signal is confined to the transmembrane domain and positive residues in its flanking regions (13-17).

Biogenesis of α-helically anchored OM proteins involves a number of import pathways, many of which are still poorly characterized. β-barrel proteins are first imported into the IMS by the translocase of the OM (TOM), where they bind to small translocase of the inner membrane (TIM) chaperones that hand them over to the sorting and assembly machinery (SAM) by which they are integrated into the OM (2,18). The insertion of some of the single membrane-spanning OM proteins depends exclusively on TOM subunits. Tom22 on the other hand requires the TOM receptors but subsequently is inserted into the OM by SAM and Mdm10 (19). Finally, insertion of some OM proteins is mediated by a protein complex, consisting of Mim1 and Mim2 (20-22), termed mitochondrial import (MIM) complex. The exact role of the MIM complex and its mode of action is still unknown (23). Moreover, it has recently been shown that a few proteins may be able to insert into the OM independently of protein factors and that at least in one case this is facilitated by specific lipids (24,25).

Mapping the variation of the import systems in different eukaryotes provides insight into the evolution of the mitochondrial protein import systems which is key to understand how the endosymbiotic ancestor of mitochondria converted into an organelle (26-29). However, most of what we know about mitochondrial protein import stems from work done in the yeast Saccharomyces cerevisiae and a few related organisms.

Here we have studied the biogenesis of POMP10 (present in the outer mitochondrial membrane proteome 10) (7), a putative signal-anchored mitochondrial OM protein in the parasitic protozoan Trypanosoma brucei, which is essentially unrelated to yeast and one of the most early diverging eukaryotes (30,31). T. brucei has a mitochondrion that in the insect-stage of the parasite’s life cycle is capable of oxidative phosphorylation. In our study we have identified the features of the import signal of POMP10. Moreover, we show that its import is independent of both Sam50 and the trypanosomal functional analog of Tom40, termed archaic translocase of the mitochondrial OM 40 (ATOM40) (32-34). However, using RNAi-mediated ablation it could be demonstrated that the biogenesis of POMP10 does require the trypanosomatid-specific OM protein termed peripheral ATOM36 (pATOM36) that is loosely associated with the ATOM complex (35). pATOM36 has recently been shown to mediate assembly of the ATOM complex as well as mitochondrial DNA inheritance (36).

RESULTS

POMP10 is a signal-anchored mitochondrial OM protein - POMP10 is an abundant mitochondrial OM protein that is conserved in all kinetoplastids and which in T. brucei has a predicted molecular weight of 62.2 kDa. It does neither have conserved domains nor orthologues outside the kinetoplastids. RNAi-mediated ablation of POMP10 neither affects growth nor mitochondrial morphology (Fig. 1), its function therefore remains unknown. Most prediction programs indicate that POMP10 has a single transmembrane domain at its very N-terminus (Table 1) that on both sides is flanked by positively charged amino acids (7). Bioinformatic analysis suggests it to be a signal-anchored protein, which would expose its soluble C-terminal domain to the cytosol. In order to experimentally confirm the predicted topology we produced a transgenic cell line expressing
a full length POMP10 variant whose C-terminus is fused to GFP. Fig. 2A shows that the POMP10-GFP fusion is expressed and that during cell fractionation it behaves like the mitochondrial marker proteins. This was confirmed by immunofluorescence (IF) analysis which yields a staining pattern that is identical to the mitochondrial OM protein ATOM40 (Fig. 4A). Protease protection assays using mitochondria isolated under isotonic conditions show that GFP-tagged POMP10 is protease-sensitive in intact mitochondria like the previously characterized protein import receptor ATOM69 (32). The IMS-localized TbTim9 and the matrix protein mtHsp70 however were only digested after the membranes were dissolved by Triton (Fig. 2B).

This indicates that the C-terminal soluble domain of the protein is exposed to the cytosol and thus establishes POMP10 as a signal-anchored mitochondrial OM protein. The small TIM chaperone, TIM9, which serves as a soluble marker of the IMS, on the other hand is resistant to the treatment illustrating that the purified mitochondria that were used in the assay have an intact OM. Only when the membrane barrier is destroyed by the addition of detergent, TIM9 becomes susceptible to the treatment and gets degraded.

**Transgenic cell lines expressing POMP10-GFP fusions** - In order to identify the signal that directs POMP10 to the OM, we produced a series of transgenic cell lines that allow tetracycline-inducible expression of a number of POMP10 variants all of which were fused to GFP at their C-terminus. The variants include a deletion of the membrane-spanning domain as well as a series of constructs in which the soluble domain was replaced by GFP. The differences between the latter concerns point mutations which replace the positively charged amino acids of either the cytosolic or the IMS-exposed membrane-flanking regions with neutral or negatively charged amino acids, respectively. Moreover, in one variant all positively charged residues were deleted (Fig. 3A). The immunoblot analysis of whole cell extracts from all transgenic cell lines using an anti-GFP antiserum shows that all POMP10-GFP fusions are expressed at comparable levels (Fig. 3B).

The transmembrane domain of POMP10 contains the import signal - The localization of the different POMP10-GFP fusions was analyzed by IF as well as by isolation of crude mitochondrial fractions using digitonin extraction (37). Moreover, in order to test whether the mitochondrially localized fusion proteins are stably inserted into the OM, we performed carbonate extraction at pH 10.8 and 11.5, respectively. This treatment disrupts protein-protein interactions but should not interfere with protein-lipid interactions. Insolubility after carbonate extraction at high pH is used as an operational definition for integral membrane proteins (38).

Fig. 4A shows that the full length POMP10-GFP fusion is correctly localized to mitochondria and behaves as an integral membrane protein (Fig. 5A). However, if the N-terminal transmembrane domain is deleted the POMP10-GFP fusion becomes soluble at low digitonin concentration and it accumulates outside mitochondria in a punctate-like pattern as evidenced by IF (Fig. 4B). The complementary fusion protein in which the transmembrane domain including the positively charged flanking residues was directly fused to the GFP on the other hand was correctly localized and inserted into the OM (Fig. 4C and Fig. 5B).

The import signal of POMP10 does not depend on positively charged amino acids - Further GFP variants that were tested either had the three N-terminal positively charged amino acids that flank the transmembrane domain (Fig. 4D), or the four positively charged lysines that flank the region C-terminally (Fig. 4E), or both (Fig. 4F), replaced by serines. All these POMP10-GFP fusions are still mitochondrially localized as evidenced by IF which results in a staining pattern that is congruent with the mitochondrial marker ATOM40. Moreover, like the full length POMP10, all these variants are recovered in a crude mitochondrial fraction obtained by digitonin extraction (Fig. 4D, 4E, 4F, right panel). However,
the carbonate extraction experiments in Fig. 5C, D, E show that replacing the positive charges by neutral ones on either side of the membrane-spanning domain causes a fraction of each of the POMP10-GFP fusions to be released into the supernatant. The soluble fraction becomes larger at higher pH and/or if both sides of the membrane-spanning domain carry neutral residues (Fig. 5E), indicating that in the absence of positively charged flanking amino acids the transmembrane domain, while still targeted to the mitochondrial OM, is less efficiently inserted into or anchored in the lipid bilayer.

Overexpression of most of the truncated POMP10-GFP constructs (Fig. 4D, 4E, 4F) that are targeted to mitochondria altered the morphology of the mitochondrial network as evidenced by enlarged sections that stain positive for mitochondrial marker proteins. All of these cell lines also grew more slowly, albeit to different extents, when expressing GFP fusion proteins (Fig. 6). This suggests that the altered mitochondrial morphology affects the overall fitness of the cells. However, the mechanism by which overexpression of the POMP10 membrane-spanning domain and its mutant derivatives when fused to GFP affect mitochondrial morphology remains to be elucidated.

Negatively charged flanking sequences abolish the POMP10 import signal - While the membrane-spanning domain of POMP10-GFP is still targeted to mitochondria if the flanking positively charged amino acids are replaced by neutral ones (Fig. 4F), targeting is abolished if the transmembrane flanking regions are lost or if the positive residues are replaced by negatively charged glutamates (Fig. 4G, 4H). IF analysis shows that the resulting POMP10-GFP fusions appear in part to be localized to discrete structures that do not overlap with the mitochondrial marker ATOM40. Moreover, the protein is released into the supernatant at low concentration of digitonin.

In vivo biogenesis of POMP10 depends on pATOM36 - Having identified the signal that targets POMP10 to the mitochondrial OM, we wanted to investigate which protein factors might mediate its biogenesis. The mitochondrial OM of *T. brucei* contains three protein complexes involved in mitochondrial protein import. The most important one is the ATOM complex, the pore-forming component of which is the β-barrel protein ATOM40 (32-34). As a functional analogue of the TOM complex it is required for import of essentially all mitochondrial proteins. Insertion of β-barrel protein from the IMS side into the OM requires the SAM complex, the pore-forming component of which is the β-barrel protein SAM50 (39). Finally, there is the trypanosomatid-specific protein pATOM36 which dynamically associates with the ATOM complex (35). Recent work has shown that pATOM36 mediates the biogenesis of a number of OM proteins including subunits of the ATOM complex (36). In order to test the involvement of the three protein complexes in the targeting and import of POMP10, we expressed the full length version of the protein fused to GFP in tetracycline-inducible ATOM40, Sam50 and pATOM36 RNAi cell lines, respectively. Using IF the localization of the GFP-tagged fusion protein was analyzed in the three RNAi cell lines. Cells that had been induced with tetracycline for one day served as a control since the GFP-tagged fusion protein is already expressed but the RNAi effects are not visible yet. At this early time point of induction the staining pattern of the POMP10-GFP fusion protein coincided in all three cell lines with the mitochondrial marker (Fig. 7A). After three days of tetracycline induction all cell lines showed a growth arrest (Fig. 7B) which has been described before (33,35,39). Moreover, as expected if mitochondrial protein import is abolished the morphology of mitochondria is drastically altered (Fig. 7A) (40). However, in the ATOM40 and the Sam50 RNAi cell lines, despite the aberrant morphology of the organelles, the localization pattern of the POMP10-GFP fusion was essentially identical to the mitochondrial markers. This indicates that ablation of neither of the two proteins
affects POMP10 biogenesis. In contrast, ablation of pATOM36 caused accumulation of POMP10-GFP outside mitochondria as evidenced by the diffuse and punctate signal that only partially overlapped with the mitochondrial marker (Fig. 7A). 

Fig. 7C shows immunoblots for the three RNAi cell lines. RNAi directed against pATOM36 leads to a rapid ablation of the protein. There is a decrease of ATOM14 and ATOM46 whose biogenesis has previously been shown to depend on pATOM36 (36). The levels of ATOM40, ATOM69 and VDAC however remain constant, except for a slight drop that is observed after 4 days of induction (Fig. 7C, left panel). As expected ATOM40 is early and efficiently downregulated in the ATOM40-RNAi cell line. The same is seen for the ATOM complex subunits ATOM14 and ATOM46. Moreover, we see downregulation of ATOM69 and VDAC. The level of pATOM36 however remains constant throughout induction and only slightly decreases at the latest time point (Fig. 7C, middle panel). RNAi of Sam50 finally leads to ablation of the two β-barrel proteins VDAC and ATOM40. As a consequence of the latter a decline of ATOM14 and ATOM46, and to a lesser extent of ATOM69, is also observed. The level of pATOM36 on the other hand remains essentially constant (Fig. 7C, right panel). Finally, elongation factor 1α (EF1α) which serves as a cytosolic control is not affected in any of the three cell lines.

In summary, these results illustrate that this is not the case since in the absence of pATOM36 only very little of the staining corresponding to the POMP10-GFP fusion protein overlaps with the ER.

All above mentioned RNAi cell lines expressing full length POMP10-GFP protein was also analyzed by extraction with low concentrations of digitonin. In Fig. 7D the resulting total, mitochondria-enriched pellet and supernatant fractions are analyzed on immunoblots probed for the POMP10-GFP fusion protein as well as for EF1α and lipoamid dehydrogenase (LipDH) that serve as cytosolic and mitochondrial markers, respectively. Whereas in cell lines ablated for ATOM40 and SAM50 the POMP10-GFP fusion protein is quantitatively recovered in the mitochondria-enriched pellet, a fraction of the protein is released into the supernatant in the absence of pATOM36. This indicates that the mitochondrial localization of POMP10 depends on pATOM36. The same is observed not only for the full length POMP10 fusion protein, but also if only the membrane-spanning domain is fused to GFP (ΔGFP(33-560)) (Fig. 7E).

In vitro biogenesis of POMP10 depends on pATOM36 - In order to confirm that pATOM36 plays an important role in the biogenesis of POMP10 we performed in vitro insertion experiments. To that end we used the POMP10 variant, in which the membrane-spanning domain was directly fused to GFP (GFP(Δ33-560)) and whose in vivo biogenesis was shown to depend on pATOM36 (Fig. 7D). Thus, the substrate was in vitro translated in the presence of [35S]methionine using reticulocyte lysate and subsequently incubated with mitochondria isolated from either uninduced cells or from cells ablated for pATOM36. To measure insertion of the protein into the mitochondrial OM, the reisolated organelles were extracted with sodium carbonate at pH 11.5 and the resulting pellets were analyzed by SDS/PAGE analysis (36). Fig. 9A shows a time-dependent increase of the radioactive substrate protein in the carbonate insoluble pellet fractions of organelles that stem
from uninduced cells, indicating the substrate is inserted into the OM. However, in mitochondria isolated from pATOM36-ablated cells much less insertion is observed. Thus, as observed in vivo, pATOM36 facilitates the in vitro insertion of POMP10-GFP(Δ33-560) into the mitochondrial OM. Interestingly, another pATOM36 substrate, ATOM46, behaved differently in the same assay (Fig. 9B, middle panel). As described previously in the case of ATOM46 pATOM36 mediates the integration of the protein into the ATOM complex rather than its insertion into the membrane (36). Fig. 9B (right panel) finally shows that membrane insertion of POMP10-GFP(Δ33-560) is not affected in mitochondria ablated for ATOM40, even though in these organelles the levels of all ATOM subunits are lower than in the induced pATOM36 cell line (Fig. 7C) (32,36).

DISCUSSION

In a recent study we have purified the OM of T. brucei and shown by a combination of protein abundance profiling and label-free quantitative mass spectrometry that its proteome consists of 82 different proteins of which 42 contain one or more putative transmembrane domains (7). Among them seven proteins are bioinformatically predicted to be signal-anchored. In the present study we selected one of these proteins, termed POMP10, to investigate its biogenesis pathway.

Using a protease protection assay we confirmed that POMP10 is indeed a signal-anchored protein. In a first series of experiments we showed that the targeting signal of POMP10 for the OM is confined to the predicted membrane-spanning domain and a few flanking residues. We then demonstrated that the positively charged flanking residues can be replaced by neutral ones without affecting mitochondrial localization. However, if the positively charged flanking residues are replaced by negatively charged ones or deleted, targeting of the protein is abolished.

The targeting sequence of signal-anchored proteins has previously been analyzed in yeast, humans and plants. In all three systems the signal was confined to a membrane-spanning domain of "moderate" hydrophobicity which however did not show sequence similarity between different signal-anchored proteins of the same or different species (13,14). Moreover, whereas for human and plant signal-anchored proteins a C-terminal positively charged flanking region was an additional requirement for OM targeting (13,41,42), this was not the case for the corresponding proteins of yeast since their positively charged residues could be replaced by neutral ones without affecting the targeting (14,15). While in the latter case the protein was still correctly targeted, its integration into the OM membrane was weakened.

The targeting signal of trypanosomal POMP10 conforms to this picture. As in the other systems it consists of a transmembrane domain with moderate hydrophobicity that includes N-terminal and C-terminal flanking regions containing positively charged residues. However, as in yeast but unlike in plants and humans, these residues are not essential for mitochondrial targeting but do influence the strength of the interaction with the OM.

Yeast and humans belong to the eukaryotic supergroup of the Opisthokonts and plants to the supergroup of the Archeaplantidae, which represent two of the six main branches of the eukaryotic phylogenetic tree (43). T. brucei is a member of the supergroup of the Excavates which are essentially unrelated to Archeaplantidae and Opisthokonts. The fact that the general features of the targeting signal for signal-anchored OM proteins are similar in all three systems suggests that they may be conserved in all eukaryotes.

While the targeting signal of signal-anchored mitochondrial OM proteins has been investigated in fungi, mammals and plants our knowledge of the components required for the biogenesis of these proteins is restricted to the two fungal species S. cerevisiae and Neurospora crassa. It has been shown in
yeast that neither the SAM complex nor the import receptors, Tom20 and Tom70, both of which are themselves signal-anchored proteins, are required for the biogenesis of signal-anchored OM proteins. Moreover, it has been demonstrated that the pore protein Tom40 can be blocked without affecting the import of signal-anchored proteins, although Tom20 still requires Tom40 for its biogenesis (9-11). Finally, a protein complex consisting of Mim1 and Mim2 has been characterized that is involved in import and assembly of at least some signal-anchored OM proteins (22,23,44,45). Mim1 and Mim2 were also shown to be involved in the insertion and/or assembly of some tail-anchored as well as multi membrane-spanning proteins. However, the mechanism by which Mim1 and Mim2 exert their function remains to be investigated.

In trypanosomes we showed that membrane insertion in vivo and in vitro of the signal-anchored POMP10 did neither require ATOM40 nor Sam50, the pore forming components of the ATOM and the SAM complexes. However, biogenesis of POMP10 did depend on pATOM36 a trypanosomatid-specific OM protein. A recent study has shown that pATOM36 has a dual localization. Similar to ATOM40 it is localized all over the OM, however, in contrast to ATOM40 it is also enriched at the tripartite attachment complex (TAC), the structure that links the mitochondrial DNA to the basal body of the flagellum. The dual localization of pATOM36 reflects its dual function in: i) the assembly of OM proteins, including most subunits of the ATOM complex (32) and ii) the segregation of the replicated mitochondrial genome (35,36).

Ablation of pATOM36 resulted in the accumulation of a fraction of POMP10 outside mitochondria in vivo and much reduced membrane insertion in vitro, suggesting that the protein is required for targeting and/or insertion of the protein into the OM. This is different to the previously analyzed substrates of pATOM36, such as ATOM complex subunits, which in the absence of the pATOM36 were still inserted into the membrane but not assembled into high molecular weight complexes (36).

It is tempting to speculate that the kinetoplastid-specific pATOM36 might be a functional analogue of fungal Mim1 and 2 which were shown to be involved in the insertion and/or assembly of some signal-anchored OM proteins in yeast (22,23,44,45). Thus, while the targeting signals for signal-anchored proteins appear to be conserved in all eukaryotes, this is not the case for the factors mediating their biogenesis. There is no sequence similarity between pATOM36 and Mim1/Mim2 and the molecular weights of the two sets of proteins are also very different: pATOM36, 36 kDa; Mim1, 13 kDa; Mim2, 11 kDa. The only recognizable motif that is shared between the two groups of proteins are two GxxxG(A) motifs in their predicted transmembrane domains. However this motif is quite frequent in transmembrane domains.

In summary our results show the existence of evolutionary distinct biogenesis factors for signal-anchored proteins in the different eukaryotic supergroups. Moreover, the fact that Mim1 and Mim2 are absent from mammals, which as the fungi belong to the Opisthokonts, indicates that there might be different biogenesis factors even within the same supergroup. Identification of biogenesis factors for signal-anchored OM proteins in different phylogenetic groups is important, as it may allow the identification of shared traits between them which will help to define the fundamental biochemical features mandatory for their function.

**EXPERIMENTAL PROCEDURE**

**Production of transgenic cell lines**

- Transgenic procyclic cell lines are based on *T. brucei* 29-13 (46). Cells were grown at 27°C in SDM79 supplemented with 10% (v/v) FCS. All plasmids are based on the pLew100 expression vector in which the phleomycine resistance gene has been replaced by a puromycine resistance gene (47) and in which the ORF coding for enhanced GFP was inserted. PCR amplicons corresponding to the different variants of POMP10 (Tb927.11.13180) were cloned into this enhanced GFP...
expression vector to allow expression of C-terminally tagged variants. Cells were grown to mid exponential phase and transfected with the corresponding linearized plasmid (48), followed by selection in medium containing puromycine. DNA constructs were verified by sequencing. RNAi constructs targeting ATOM40 (33), SAM50 (49) and pATOM36 (35) have been described elsewhere.

**Immunoblotting** - Protein samples were separated by SDS-PAGE and transferred onto nitrocellulose membranes (GE Healthcare). Membranes were blocked in 5% (w/v) milk in PBS-Tween (137 mM CaCl₂, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.8 mM KH₂PO₄, 0.1% (v/v) Tween-20, pH 7.4) and incubated overnight, with the appropriate antibody at 4°C. Monoclonal primary antibodies used were: mouse anti-GFP (Roche, dilution 1:1’000) and mouse anti-EF1α (Merck Millipore, dilution 1:10’000). Polyclonal primary antibodies produced in rabbits were: anti-VDAC (dilution 1:1’000) (50), anti-ATOM40 (dilution 1:1’000) (32), anti-Cyt c (dilution 1:1’000) (7) and anti-TIM9 (1:20) (7) and rabbit anti-LipDH (1:10’000) (provided by L. Krauth-Siegel, Heidelberg University). Detection of the antibodies was done with an Odyssey Infrared imaging system using IRDye secondary antibodies (Li-COR Biosciences, dilution 1:20’000).

**Immunofluorescence** - The cells were collected, washed in PBS, and subsequently resuspended in ice-cold buffer. All further steps were performed in a wet chamber. Cells were fixed for 10 minutes in freshly made 4% paraformaldehyde in PBS (pH 7.2) and rinsed with ice-cold PBS. Cells were incubated in PBS containing 0.2% Triton-X100 for 5 minutes. Primary antibodies were rabbit anti-VDAC (1:100), rabbit anti-ATOM40 (1:1’000), rabbit anti-ATOM69 (1:3) (32) and rabbit anti-BIP (1:5’000). Cy3-conjugated goat anti-rabbit (dilution 1:500) was used as a secondary antibody. Cells were rinsed in ice-cold buffer, post-fixed in cold methanol and mounted with VectaShield containing DAPI (Vector Laboratories). Fluorescence images were taken with Leica fluorescence microscope (Leica Microsystems). Images stacks were recorded and deconvoluted using LAS X software from Leica (Leica Microsystems).

**Digitonin extractions** - Cell membranes were lysed by resuspension of 10⁶ cells in SoTe buffer (20 mM Tris-HCl pH 7.5, 0.6 M sorbitol and 2 mM EDTA) containing 0.015% (w/v) digitonin followed by differential centrifugation. This yielded a mitochondria-enriched pellet and a fraction enriched for cytosolic proteins (37). All samples were analyzed by immunoblot experiments.

**Carbonate extractions** - Mitochondria-enriched fractions were resuspended in 160 µl of 100 mM Na₂CO₃, pH 11.5 or 10.8. Thereof, 80 µl were removed and mixed with 20 µl 5x SDS loading buffer and boiled to serve as the ‘total’ sample. The remaining 80 µl were incubated on ice for 10 minutes and centrifuged (1000’000g, 4°C, 10 min). The supernatant was transferred to a new tube and 20 µl 5x SDS loading buffer was added prior to boiling. The pellet was resuspended in 80 µl of 100mM Na₂CO₃ and 20 µl 5x SDS loading buffer was added prior to boiling. All samples were analysed by SDS-PAGE.

**Protease protection assay** - Isotonically isolated mitochondria (25µg) from cells (51,52) expressing POMP10-GFP were resuspended in 20 mM Tris-HCl pH 7.2, 15 mM KH₂PO₄, 20 mM MgSO₄, 0.6 M sorbitol in a total volume of 50 µl in presence of proteinase K (10 mg/ml) containing or not 0.5% (v/v) Triton-X100 followed by incubation on ice for 15 min. Reactions were stopped by adding PMSF (phenylmethylsulfonylfluorid) at 5mM and mitochondria were centrifuged (6’800 g, 4°C), resuspended in SDS-loading buffer and boiled.

**In vitro insertion of proteins into the mitochondrial OM** - The membrane insertion assay was done exactly as described in (36). ³⁵S-Met-labelled POMP10-GFP(A33-560) and ATOM46 were synthesized using the TNT T7 Quick for PCR in vitro translation kit (Promega). For the coupled transcription and translation, gel-eluted PCR-fragments
were used that consisted of the T7 RNA polymerase promoter fused to the complete ORF of the corresponding substrates.

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Author contributions: JB designed and performed research (results shown in Fig.1-8), SK designed and performed research (results shown in Fig. 7C and 9). JB and SK analyzed the results and prepared the Figures. JM planed and analyzed the experiments. AS analyzed the data and wrote the paper.

REFERENCES


FIGURE LEGENDS

FIGURE 1. RNAi-mediated ablation of POMP10 does neither affect growth nor mitochondrial morphology. (A) Growth curve of uninduced (-Tet) and induced (+Tet) POMP10-RNAi cell line. Inset, Northern blot of the POMP10 mRNA after two days of RNAi induction. Ethidiumbromide-stained gel showing the rRNAs served as a loading control. (B) Immunofluorescence analysis of induced (2 d and 7 d) and uninduced RNAi cells. Cells were stained with ATOM40 antiserum as a mitochondrial marker (red) and 4′,6-diamidin-2-phenylindol (DAPI) to highlight nuclear and mitochondrial DNA (blue).

FIGURE 2. POMP10 is a signal-anchored mitochondrial OM protein. (A) Upper panel, outline of the mitochondrial purification scheme. Lower panel, immunoblot analysis of the indicated fractions of a mitochondrial purification from the cell line expressing full length POMP10 that is C-terminally tagged with GFP (POMP10-GFP). The fractions (5 µg each) depicted are: whole cells (wc), cytosol (cyto), crude mitochondria (crude) and pure mitochondria (pure). The samples were separated by SDS-PAGE, blotted and probed for: GFP; the mitochondrial markers ATOM40, cytochrome C (Cyt c) and lipoamid dehydrogenase (LipDH) and the cytosolic marker elongation factor 1α (EF1α). (B) Immunoblots of a protease protection assay using gradient-purified structurally intact mitochondria isolated from cell lines expressing POMP10-GFP analyzed by anti-GFP antiserum. Additions of proteinase K and of Triton X-100 are indicated. The OM protein ATOM69 serves as a control OM protein. The IMS protein Tim9 and the matrix protein mtHsp70 serve as controls for OM integrity. In the presence of Triton X-100 mtHsp70 is degraded to a protease resistant fragment.

FIGURE 3. Transgenic cell lines expressing POMP10-GFP fusions. (A) FL, full length GFP fusion. For the other fusion proteins the extent of the deletions (Δ) as well as the positions that were replaced with either serine (S) or glutamate (E) are indicated on the left. Amino acid sequences of the signal-anchor domains in the indicated POMP10-GFP fusion proteins are shown in the middle. The predicted membrane-spanning domain is underlined. The serine and glutamate residues that were used to replace the positively charged amino acids are indicated in bold. The symbols used for the different GFP constructs are shown on the right: black line, cytosolic domain; box, signal-anchor domain; large white box, predicted membrane-spanning domain; small black boxes, positively charged flanking sequence(s); small white boxes, uncharged flanking sequence(s); small grey boxes, negatively charged
flanking sequences. (B) Immunoblot analysis of total cellular extracts from equal cell numbers of the parental strain *T. brucei* 29-13 and the indicated transgenic cell lines expressing the various POMP10-GFP fusion proteins were probed for GFP and EF1α, which serves as a loading control.

FIGURE 4. Characterization of the targeting signal in POMP10. IF analysis of the indicated POMP10-GFP fusion proteins (A–H, left columns) in procyclic forms of transgenic *T. brucei* (A–H, middle column). Samples were probed for GFP and the mitochondrial marker ATOM40, respectively. An overlap of both signals is also shown (Merge). Bar, 10µm. Crude digitonin-based cell fractionation analysis (A–H, right column) of cell lines expressing the indicated POMP10-GFP fusion proteins. Total cellular extract (T), crude mitochondrial fraction (P) and digitonin-extracted cytosolic fraction (S). LipDH and EF1α serve as mitochondrial and cytosolic markers, respectively.

FIGURE 5. Carboxylate extractions at high pH of membrane-associated POMP10-GFP fusion proteins. Immunoblots of equal cell equivalents of total (T), pellet (P) and supernatant (S) fractions of a carboxylate-extracted crude mitochondrial pellet obtained by digitonin extraction of cells expressing the indicated POMP10-GFP variants (A–E). The extraction was performed both pH 10.8 and pH 11.5 and immunobots were analyzed by anti-GFP. ATOM40 and Cyt c serve as markers representing an integral and peripheral membrane protein, respectively.

FIGURE 6. Expression of some of the truncated POMP10-GFP fusions interferes with growth. Growth curves of uninduced (black line) and induced (red line) cell lines expressing the indicated recombinant POMP10-GFP fusion proteins. All analyzed cell lines show an altered mitochondrial morphology (see Fig. 4D, 4E, 4F). Insets, immunoblots confirming expression of the GFP fusion proteins. VDAC serves as loading control.

FIGURE 7. In vivo biogenesis of POMP10 depends on pATOM36. (A) IF analysis of full length POMP10 that is C-terminally tagged with GFP (GFP(FL): green) and mitochondrial marker proteins (VDAC and ATOM69: red) in tetracycline-inducible pATOM36 (left panels), ATOM40 (middle panels) and SAM50 (right panels) RNAi cell lines. Time of induction in days (d) is indicated at the top. Bar, 10µm. (B) Growth curves of tetracycline-inducible pATOM36 (left panels), ATOM40 (middle panels) and SAM50 (right panels) RNAi cell lines that co-express full length POMP10-GFP and that were used in (A). (C) Equal cell equivalents of the same RNAi cell lines shown in (A) (left panel, pATOM36; middle panel, ATOM40; right panel, SAM50) were analyzed for the presence or absence of pATOM36, ATOM40, ATOM14, ATOM46, ATOM69, VDAC and cytosolic EF1α using immunobots. Time of RNAi induction is indicated at the top of each panel. (D) Crude digitonin-based cell fractionation analysis of the same cell lines shown in (A) and (B). Total cellular extract (top panels), crude mitochondrial fraction (pellet, middle panels) and digitonin-extracted cytosolic fraction (SN, bottom panels) were probed for POMP10-GFP (GFP(FL)). LipDH and EF1α serve as mitochondrial and cytosolic markers, respectively. (E) Same as (D) but a tetracycline-inducible pATOM36 RNAi cell line that co-expresses the POMP10 fusion protein in which the transmembrane domain including the positively charged flanking residues was directly fused to the GFP was analyzed.

FIGURE 8. Ablation of pATOM36 does not cause mislocalization to the ER. IF analysis of full length POMP10 that is C-terminally tagged with GFP (POMP(FL): green) and the ER marker protein BiP (red) in the tetracycline-inducible pATOM36 RNAi cell line. Time of induction in days (d) is indicated at the top. Bar, 10µm.

FIGURE 9. In vitro biogenesis of POMP10 depends on pATOM36 (A) Left panel, in vitro insertion assay of the 35S-Met-labeled POMP10-GFP(D33-560) using mitochondria isolated
from the uninduced (-Tet) and induced (+Tet, 2 days) procyclic pATOM36-RNAi cell line. Incubation times are indicated at the top. The pellet fraction of an alkaline carbonate extraction were separated by SDS-PAGE and analyzed by autoradiography. Lower panel, section of the Coomassie-stained gel serve as loading controls. Graph on the right, quantification of triplicate experiments shown on the left. Standard errors are indicated. (B) Graph of triplicate in vitro insertion assays (9 min time point) using isolated mitochondria of uninduced and induced pATOM36 and ATOM40 RNAi cell lines, respectively. The tested substrates are indicated at the top. Standard errors are indicated. The SDS-gels representing a typical experiments including their corresponding loading controls are indicated below the graphs. Bottom panels, immunoblot confirming the specific downregulation of pATOM36 and ATOM40 in the respective RNAi cell lines.

**TABLES**

<table>
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<tr>
<th>Name of server</th>
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<td>TOPCONS</td>
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**Table 1.** Prediction of transmembrane domains using the indicated prediction tools.
Figure 1
Figure 2
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<tr>
<th>Name</th>
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Figure 3
Figure 4
Figure 5
Figure 6
Figure 7
Figure 8
Figure 9