

The Taz1p Transacylase Is Imported and Sorted into the Outer Mitochondrial Membrane via a Membrane Anchor Domain

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Mutations in the mitochondrial transacylase tafazzin, Taz1p, in *Saccharomyces cerevisiae* cause Barth syndrome, a disease of defective cardiolipin remodeling. Taz1p is an interfacial membrane protein that localizes to both the outer and inner membranes, lining the intermembrane space. Pathogenic point mutations in Taz1p that alter import and membrane insertion result in accumulation of monolysocardiolipin. In this study, we used yeast as a model to investigate the biogenesis of Taz1p. We show that to achieve this unique topology in mitochondria, Taz1p follows a novel import pathway in which it crosses the outer membrane via the translocase of the outer membrane and then uses the Tim9p-Tim10p complex of the intermembrane space to insert into the mitochondrial outer membrane. Taz1p is then transported to membranes of an intermediate density to reach a location in the inner membrane. Moreover, a pathogenic mutation within the membrane anchor (V224R) alters Taz1p import so that it bypasses the Tim9p-Tim10p complex and interacts with the translocase of the inner membrane, TIM23, to reach the matrix. Critical targeting information for Taz1p resides in the membrane anchor and flanking sequences, which are often mutated in Barth syndrome patients. These studies suggest that altering the mitochondrial import pathway of Taz1p may be important in understanding the molecular basis of Barth syndrome.

Barth syndrome (BTHS) is a recessive X-linked disorder which causes cardiac myopathy, neutropenia, metabolic abnormalities, and atypical mitochondrial function and morphology, with cardiac failure and other potentially fatal complications often, but not always, presenting in infancy and childhood (1–4). The human tafazzin gene (*TAZ*) was first implicated in the disease by linkage analysis (1, 5). Clinical studies and research in several model organisms, ranging from *Saccharomyces cerevisiae* to the fly, zebrafish, and mouse, show that mutations in or deletions of the *TAZ* locus result in abnormal phospholipid species, with the mitochondrial phospholipid cardiolipin (CL) affected the most severely (5–11). Tafazzin functions as a transacylase in the CL maturation pathway by replacing saturated chains with unsaturated chains (7). A hallmark of BTHS is the accumulation of the CL remodeling intermediate monolysocardiolipin (monolysocardiolipin), which lacks a fatty acid (6, 12).

The mitochondrial localization of Taz1p as well as other lipid biogenesis proteins in mitochondria is complex. Cardiolipin synthase, *Crd1p*, synthesizes CL in the matrix-facing leaflets of the inner membrane (13). Localization studies showed that wild-type Taz1p localizes to the inner and outer mitochondrial membranes, facing the intermembrane space (6, 14, 15). When yeast was used as a system to model Taz1p mutations associated with BTHS, a subset of Taz1p mutants correctly localized to the intermembrane space but assembled in complexes that were not functional (6) or were degraded by the intermembrane space *i*-AAA protease *Yme1* (12). Another subset of mutants localized to the mitochondrial matrix, suggesting that mistargeting of tafazzin resulted in BTHS in a subset of patients (6). Given the unexpected localization for tafazzin within mitochondria and that protein mistargeting could be the cause of BTHS in this subset of patients, a detailed study of Taz1p biogenesis is warranted. Taz1p assembly in the membrane is also unusual in that Taz1p is an interfacial membrane protein, anchored by a hydrophobic membrane anchor that protrudes into but not completely through the

lipid bilayer (residues 215 to 232) (6). The membrane anchor of Taz1p is also important for mitochondrial localization because point mutations in the membrane anchor (recapitulating pathogenic point mutations in Taz1p identified in BTHS patients) result in mistargeting to the mitochondrial matrix, where Taz1p either assembles in the leaflet of the inner membrane facing the matrix or forms protein aggregates (6).

The import of mitochondrial proteins is accomplished by a complex translocation system consisting of complexes in the outer membrane, inner membrane, and intermembrane space (16, 17). Whereas many mitochondrial proteins contain a canonical N-terminal targeting sequence, membrane proteins contain internal targeting information that typically resides in the regions near the membrane-spanning domains (18). We have previously shown that Taz1p does not contain a typical N-terminal targeting sequence (6); thus, mitochondrial targeting of Taz1p is directed presumably via an internal targeting sequence.

To date, four general types of outer membrane proteins have been identified in mitochondria (19–22). The targeting and sorting mechanisms that dictate their correct localization often overlap and, in some cases, are not fully characterized. Signal anchored proteins, such as the Tom20p and Tom70p receptors of the TOM complex, have a moderately hydrophobic alpha-helical transmembrane domain in the N terminus, which, in addition to pos-

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itive charges in the flanking sequences, serves as both an intercellular sorting domain and a membrane anchor. Tail-anchored proteins such as Fis1p, Tom5p, Tom6p, Tom22p, and Gem1p have an alpha-helical C-terminal transmembrane domain and often require a moderately net positive charge located C terminal to the tail-anchor domain (23). Finally, a subset of proteins have multiple transmembrane domains, including those that are alpha-helical (i.e., Fzo1p and Ugo1p) and the β -sheets of β -barrel proteins (i.e., Tom40p and porin). Mim1 mediates the insertion of outer membrane proteins with alpha-helical membrane domains (21, 22). In contrast, the β -barrel proteins have targeting information at the very C terminus and are imported via the TOM complex and then inserted from the intermembrane space side via the sorting and assembly (SAM) complex (24). The small Tim proteins (Tim9p-Tim10p and Tim8p-Tim13p) chaperone the β -barrel proteins through the intermembrane space (25).

Membrane proteins of the inner membrane can use the TIM22 or TIM23 import pathways. The TIM22 pathway (16, 17, 26) mediates the insertion of polytopic carrier proteins such as the ADP/ATP carrier (AAC) and the phosphate carrier. In this pathway, the 70-kDa small Tim chaperone complexes of the intermembrane space, Tim8p-Tim13p and Tim9p-Tim10p, bind to hydrophobic regions of the precursor and mediate translocation to the insertion complex that consists of Tim12p, Tim22p, Tim54p, Tim18p, and a fraction of the small Tim proteins Tim9p and Tim10p. Both matrix targeted and inner membrane proteins that contain an N-terminal targeting sequence are translocated through the TIM23 pathway and may be inserted into the inner membrane by the TIM23 complex or Oxa1p, depending on sorting information (27, 28).

Even though the Tim9-Tim10p complex has been previously implicated in Taz1p biogenesis (15, 29), in this study, we investigated Taz1p import in detail. Taz1p is first imported into the outer membrane using the TOM complex in a Tim9p-Tim10p dependent manner. The membrane anchor and N-terminal flanking sequences are important for targeting. Taz1p then is sorted to the inner membrane via intermediate density membranes to be distributed in both the outer and inner membranes, where it may modify cardiolipin in the membrane leaflets that line the intermembrane space.

MATERIALS AND METHODS

Strains and growth conditions. Standard conditions were used for the growth, manipulation and transformation of yeast, with the parental strain GA74-1A used except where indicated. Characterization of the following temperature-sensitive or deletion mutant strains with the following relevant genotypes was detailed previously: *tim22-4* (30), *tim10-1* (31), $\Delta tom20$ (YTJB10) and $\Delta tom70$ (YVH1) (32), *tim23-6* (33), and $\Delta taz1$ (6). All strains were grown in YPEG to mid-log phase and mitochondria were harvested from yeast as previously described (34). $\Delta taz1$ and $\Delta taz1$ [pTaz1] strains were described previously (6). The $\Delta taz1$ [pTaz1 Δ MA²¹⁵⁻²³²] strain was generated by transformation of the $\Delta taz1$ parental strain.

Cloning and *in vitro* transcription/translation. The *TAZ1* open reading frame was subcloned into an SP6 polymerase *in vitro* transcription/translation vector with an N-terminal Kozac sequence to enhance transcription (GCC GCC ATG TCT). The V224R mutant was subcloned from pRS425Taz1^{V224R} using NdeI and AvrII restriction enzyme sites. The pRS425Taz1 Δ MA²¹⁵⁻²³² plasmid was made with overlap extension PCR to remove the membrane anchor; briefly, the DNA products from the primers sets A and B (GAAGCCATATCCCAGAAG and ATGCTATTTGATCATTCCTGGTAATACCCC, respectively) and C and D (CA GAATGATCAAAATAGCATCCGAAGCAGTC and GAACCGCTTCCA

CCATGAGG, respectively) were diluted and used as a template with primer set A and D, and the product of this reaction was cut with NdeI and AvrII and subcloned into the pSP64Taz1 plasmid. The dihydrofolate reductase (DHFR) fusion constructs were subcloned into the pSP65DHFR-Tim23-DHFR vector in which Tim23 was replaced with Taz1 sequences; the inserts were made by amplifying the appropriate Taz1 regions with primers that would add an AvaI restriction enzyme site and a NAAIRS linker region (AATGCTGCTATACGATCG) N terminally and a BamHI I site and Gly linker region (GGTGGCGGAGGGGGTGGC) C terminally. To construct the pSP65DHFR-Taz1^{V224R}-DHFR plasmid, the same primers were used to amplify a fragment from the pSP64Taz1^{V224R} template. All ³⁵S-radiolabeled precursors were generated with a Promega TNT kit, but in early experiments (with the *tim22-4* and *tim10-1* strains), a rabbit reticulocyte lysate was used as previously described (33).

***In vitro* import assays.** The precursor from the *in vitro* transcription/translation reaction mixture was incubated at 25°C with isolated mitochondria in import buffer (1 mg/ml of bovine serum albumin, 0.6 M sorbitol, 0.6 M KCl, 10 mM HEPES-KOH [pH 7.1], 10 mM MgCl₂, 2.5 mM EDTA, 5 mM L-methionine, 2 mM ATP, and 2 mM NADH). All import reactions were performed at 25°C, and the reactions were quenched at specific time intervals. When an energy-regenerating system was used, the reaction buffer was supplemented with 100 μ g/ml of creatine phosphate kinase (CPK), 16 mM creatine P_i, and 10 mM succinate. Where indicated, the potential across the mitochondrial inner membrane was dissipated using 1 μ M valinomycin and 25 μ M *p*-trifluoromethoxy carbonyl cyanide phenylhydrazone (FCCP). Nonimported radiolabeled precursor was removed by treatment with 20 μ g/ml of trypsin or 5 μ g/ml of proteinase K for 30 min on ice. Trypsin was inhibited with 100 μ g/ml of soybean trypsin inhibitor and proteinase K with 1 mM phenylmethylsulfonyl (PMSF). For alkali extraction, the samples were first pelleted, resuspended in 100 mM Na₂CO₃ (pH 10.5), incubated on ice for 30 min, and then centrifuged at high speed (100,000 \times g) in an Airfuge. Unless indicated otherwise, all samples were resuspended in SDS sample buffer supplemented with 2-mercaptoethanol (BME), loaded onto 12% polyacrylamide gels, and dried for quantitative autoradiography using a Bio-Rad FX Pro plus fluorimager/phosphorimager. Bio-Rad Quantity 1 software was used to quantitate the gels, and the last time point of import into wild-type mitochondria was set to 100%.

Cross-linking and immunoprecipitation. The cross-linking and immunoprecipitation protocol was as previously described (30), with the following modifications: the import reaction mixture contained 1 μ M valinomycin and 25 μ M FCCP to trap the import intermediate. The mitochondria were incubated for 30 min on ice with 1 mM dithiobis[succinimidyl propionate] (DSP), and the reaction was quenched for 15 min with 10 mM Tris-HCl (pH 8.0). For immunoprecipitation, the mitochondria were first solubilized with TNET buffer (50 mM Tris-HCl [pH 7.5], 150 mM NaCl, 5 mM EDTA, and 1% Triton X-100) and incubated with the indicated polyclonal rabbit antibodies coupled to protein A-Sepharose beads overnight. Before loading to the 10% polyacrylamide gel, the cross-links of the immunoprecipitation samples were treated with 1 mM 2-mercaptoethanol to reduce the cross-linked products.

Subcellular fractionation, mitoplast manipulation, and digitonin solubilization assays. Subcellular fractionation was performed as previously described (6). Osmotic shock was performed by incubating the mitochondria after the import assay in 0.03 M sorbitol and 20 mM HEPES-KOH (pH 7.4) for 30 min on ice, with gentle agitation. When necessary, 10 μ g/ml of proteinase K with or without 0.1% Triton X-100 was added to the solution. The mitoplasts (mitochondria that have had the outer membrane disrupted) were pelleted in a microcentrifuge and resuspended in sample buffer, and the released soluble proteins were precipitated with 20% trichloroacetic acid (TCA). For the digitonin solubilization assay (35), aliquots were taken from either resuspended mitochondria or the import assay, pelleted, and resuspended in SEMK (250 mM sucrose, 1 mM EDTA, 10 mM morpholinepropanesulfonic acid [MOPS]-KOH [pH 7.2], 100 mM KCl) buffer that contained increasing amounts of digitonin

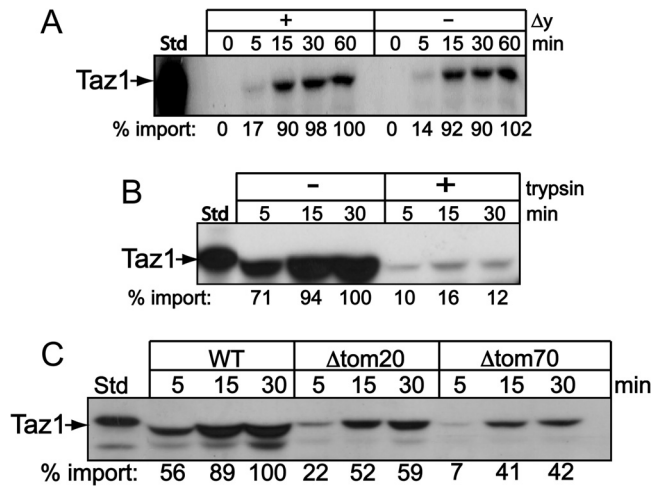


FIG 1 Taz1p import is not dependent on the $\Delta\psi$ and is first imported through the TOM complex. (A) Radiolabeled Taz1p was incubated with wild-type mitochondria, and aliquots were removed at the indicated time points and incubated with 20 $\mu\text{g}/\text{ml}$ of trypsin. Valinomycin at 1 μM and FCCP at 50 μM were added to the reaction buffer to dissipate the $\Delta\psi$. Samples were separated by SDS-PAGE, and imported Taz1p was detected by autoradiography. Imported Taz1p was quantitated using Bio-Rad Quantity One software, with the last time point in the energized mitochondria set to 100%. (B) Taz1p was imported into isolated yeast mitochondria that were pretreated with 20 $\mu\text{g}/\mu\text{l}$ of trypsin. p, precursor; m, mature form of Su9-DHFR. (C) Import of Taz1p into isolated $\Delta\text{tom}20$ and $\Delta\text{tom}70$ mitochondria.

(0% to 0.5% in 0.1% increments). The samples were quickly pelleted in the Airfuge at $100,000 \times g$ at 4°C. The pellet (P) was resuspended in SDS sample buffer supplemented with β -mercaptoethanol, and the supernatants (S) were TCA precipitated.

Blue native gel electrophoresis. After the import assay, the mitochondria were pelleted and then solubilized in 20 mM HEPES-KOH (pH 7.4), 50 mM NaCl, 10% glycerol, 2.5 mM MgCl_2 , 1 mM EDTA, 1 mM PMSF, and 1% digitonin for 30 min on ice. Insoluble material was removed by spinning the samples in the microcentrifuge at high speed for 15 min. The solubilized complexes were supplemented with 50 mM 6-aminocaproic acid and 0.5% Coomassie brilliant blue G-250 and then loaded to a 6 to 16% gradient gel at 4°C. The gel was either dried and exposed to film for autoradiography or blotted to polyvinylidene difluoride (PVDF) membranes for Western blot analysis.

RESULTS

Taz1p import is not dependent on a membrane potential. The atypical localization of Taz1p to both the inner and outer mitochondrial membranes lining the intermembrane space warrants detailed characterization of the biogenesis of Taz1p (6). We first investigated whether a membrane potential ($\Delta\psi$) was required for Taz1p import. Whereas proteins that are imported across the inner membrane and use the TIM23 or TIM22 translocons require a $\Delta\psi$ for translocation, proteins that are imported directly to the outer membrane do not require a $\Delta\psi$ (16). Radiolabeled Taz1p was imported into isolated mitochondria in which the electrochemical potential was established with NADH or dissipated with valinomycin and *p*-trifluoromethoxy carbonyl cyanide phenylhydrazide (FCCP). An import time course was performed, nonimported precursor was removed by exogenous treatment with protease, and insertion into the membrane was verified by carbonate extraction (Fig. 1A). Samples were separated by SDS-PAGE and data quantitated following autoradiography. The rates of Taz1p

import were similar in the absence or presence of a $\Delta\psi$, consistent with a previous study (15). Taz1p translocation thus seems to bypass the TIM23 and TIM22 translocons of the inner membrane.

To determine the role of the TOM translocon, we performed “trypsin shaving” in which mitochondria were pretreated with trypsin to remove the TOM receptors prior to import. Pretreatment with protease decreased Taz1p import by 80% compared to import into untreated mitochondria (Fig. 1B), suggesting a requirement for the cytosolic portion of outer membrane receptors in the early steps of Taz1p translocation (36). To dissect transport of Taz1p through the TOM translocon in greater detail, purified mitochondria from both a $\Delta\text{tom}20$ and a $\Delta\text{tom}70$ strain were used in the *in vitro* protein import assay. Tom20p facilitates the import of precursors with a typical N-terminal targeting sequence as well as β -barrel proteins (37), while Tom70p mediates the import of the carrier proteins that have internal targeting sequences (38). As controls, import of the carrier AAC and Su9-DHFR were compromised in $\Delta\text{tom}70$ and $\Delta\text{tom}20$ mitochondria, respectively (see Fig. S1 in the supplemental material). In contrast, Taz1p import was deficient in both mutant mitochondria, indicating that both Tom70p and Tom20p facilitate Taz1p targeting to the TOM translocation channel (Fig. 1C).

The TIM23 and TIM22 membrane translocation complexes are not required for Taz1p import. We took advantage of temperature-sensitive mutants in the TIM22 and TIM23 translocation pathways to assess the requirement of inner membrane translocons. Taz1p was imported into mitochondria isolated from the *tim23-6* and *tim22-4* mutants (Fig. 2A and B) (30, 33). The impairment of each translocation pathway was demonstrated using AAC and Su9-DHFR precursors for the *tim22-4* and *tim23-6* mutant strains, respectively (see Fig. S1C and D in the supplemental material). Taz1p import was similar in the mutant and wild-type (WT) mitochondria. Thus, Taz1p import is independent of the TIM22 and TIM23 translocons, suggesting that the inner membrane translocons do not play a direct role in the initial import of Taz1p. In contrast, the import of Taz1p was decreased in the *tim10-1* mutant mitochondria (Fig. 2C). This supports our previous result in which Taz1p import was blocked by the Tim10p-specific inhibitor MitoBloCK-1 (29). To confirm that Tim10p played a role in Taz1p import, we used mitochondria that contained a suppressor mutation in *TIM9* that restored import to the *tim10-1* mutant (designated *tim10-1tim9^S*) (29, 31). Indeed, import was restored to levels near that of WT mitochondria (Fig. 2C).

To confirm that Tim10p played a direct role in Taz1p import, we used cross-linking and coimmunoprecipitation to characterize import components that were bound to Taz1p. Radiolabeled Taz1p was imported into mitochondria, followed by cross-linking with 1 mM dithiobissuccinimidylpropionate (DSP) and quenching with excess Tris buffer. Mitochondria were solubilized and immunoprecipitated with antibodies directed against Tim10p, Tom40p, cytochrome (CYT) *b*₂, cytochrome *c* peroxidase (CCP1), Tim22p, and Tim23p (Fig. 2D). Taz1p was bound directly to Tim10p and Tom40p but not Tim22p or Tim23p. Taz1p also was not bound to the negative controls CCP1 and CYT *b*₂, verifying a specific interaction of Taz1p with Tom40p and Tim10p. This series of import experiments supports a direct task for Tom40p and Tim10p in the biogenesis of Taz1p.

The Taz1p precursor is initially imported into the outer membrane. We took advantage of the differing solubilities of the mitochondrial outer and inner membranes to digitonin (35) to substantiate Taz1p import via the outer membrane. Mitochondria

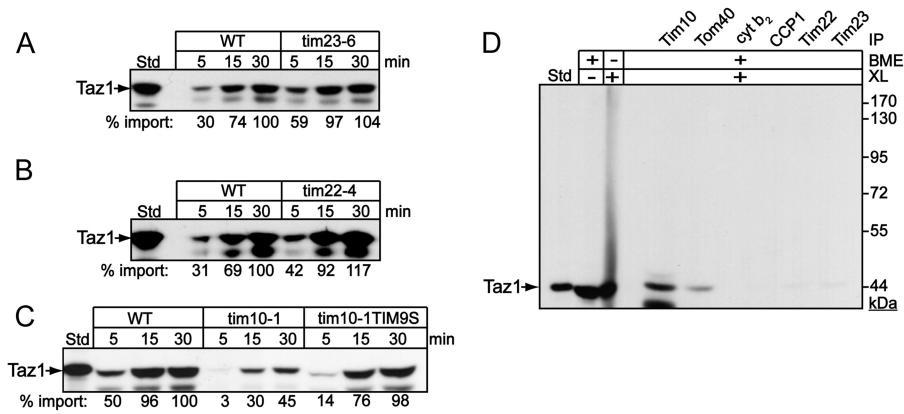


FIG 2 Taz1p import is independent of the TIM22 and TIM23 translocons. (A to C) Taz1p was imported into *tim23-6*, *tim22-4*, *tim10-1*, and *tim10-1tim9^S* mitochondria as described for Fig. 1A. (D) Taz1p was imported into WT mitochondria followed by cross-linking (XL) with 1 mM DSP and then coimmunoprecipitated (IP) with antibodies directed against Tim10p, Tom40p, cytochrome *b₂*, CCP1, Tim22p, and Tim23p. Cross-links were broken with BME prior to loading the samples to the SDS-PAGE gel to confirm that the conjugates were Taz1p.

drial markers from different compartments can be differentially released to the supernatant upon treatment with increasing concentrations of digitonin, because digitonin selectively solubilizes the outer membrane with its high lipid content, followed by the inner membrane with its increased protein concentration. Thus, soluble intermembrane space proteins, such as cytochrome *b₂*, are released to the supernatant following a high-speed spin after solubilization with a low digitonin concentration. Outer membrane proteins that are fully integrated into the membrane, such as porin and OM45, are released at intermediate digitonin concentrations, and finally, inner membrane proteins, such as AAC, are released at

a higher detergent concentration (Fig. 3A; see also Fig. S2A in the supplemental material). Endogenous tafazzin had a release profile that is intermediate between the outer and inner membrane markers in both WT and *tim23-6* mitochondria (Fig. 3A and C; see also Fig. S2A and C). This profile could be indicative of contact sites or reflect the dual residence of Taz1p on both the outer and inner membranes. However, the Taz1p release profile shifted to that of an outer membrane protein in *tim10-1* mitochondria (Fig. 3B; see also Fig. S2B), suggesting that Taz1p distribution is aberrant. We also investigated the distribution of radiolabeled Taz1p following import; similarly, radiolabeled porin and Tim54 were imported as

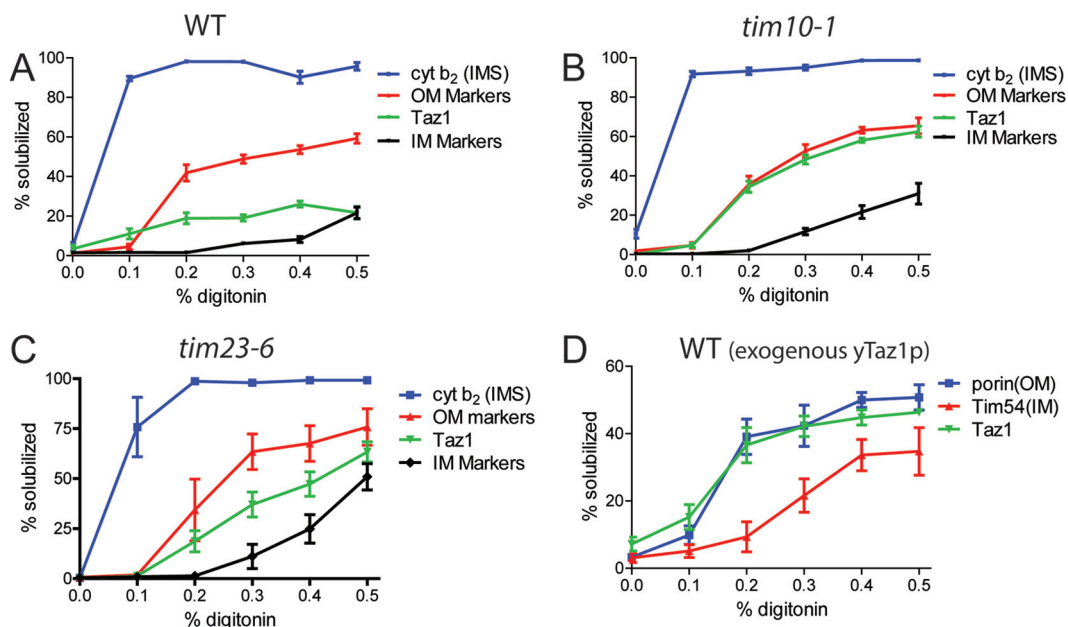


FIG 3 Taz1p is first imported into the outer mitochondrial membrane. (A) WT mitochondria (15 mg/ml) were solubilized with the indicated concentration of digitonin, and the soluble proteins were separated from the mitochondrial pellet by centrifugation. Outer (porin and OM45) and inner (Tim45p and AAC) membrane markers were averaged, and *cyt b₂* marked the intermembrane space. Equal amounts were analyzed by immunoblotting and quantitated with Bio-Rad Quantity 1 software. The percentage solubilized was calculated as $[(S)/(S + P) \times 100]$ for each detergent concentration (mean \pm SD, $n = 3$). Immunoblots are provided in Fig. S2A to D in the supplemental material. (B) Same as panel A, with *tim10-1* mitochondria. (C) Same as panel A, with *tim23-6* mitochondria. (D) Taz1p was imported into WT mitochondria for 15 min, and then fractionation was investigated for the imported Taz1p as in panel A. Markers included imported porin and Tim54p.

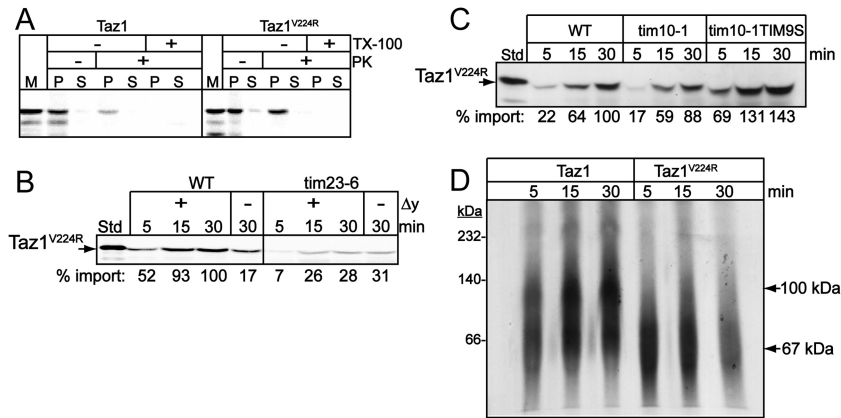


FIG 4 Taz1p^{V224R} is imported into the matrix via the Tim23 pathway. (A) Taz1p and Taz1p^{V224R} were synthesized *in vitro* and imported into WT mitochondria, which were subjected to osmotic shock, followed by centrifugation to separate the supernatant (S) from the mitoplast pellet (P). Mitochondria were also treated with 20 μ g/ml of proteinase K or 20 μ g/ml of proteinase K plus 1% Triton X-100. As a control, M was included. (B) Radiolabeled Taz1p^{V224R} was imported into both WT and *tim23-6* mitochondria in the presence and absence of a membrane potential, followed by protease treatment to remove nonimported precursor. (C) Taz1p^{V224R} was imported as in panel B in WT, *tim10-1*, and *tim10-1tim9^S* mitochondria. (D) Taz1p and Taz1p^{V224R} were imported into WT mitochondria, and the samples were solubilized in 1% digitonin and separated on a 6 to 16% blue native gel.

controls for outer and inner membrane markers, respectively (Fig. 3D; see also Fig. S2D). Imported Taz1p was localized to the outer membrane after import into WT mitochondria because Taz1p shared a digitonin solubilization profile with that of imported porin. The Taz1p profile was also similar to another imported outer membrane marker, Tom40p (data not shown). The same analysis was not performed with *tim10-1* mitochondria because the import signal was too weak for analysis. We chased the import for up to 2 h, but the imported Taz1p remained in the outer membrane and a distribution profile similar to Taz1p at steady state in WT mitochondria (Fig. 3A; see also Fig. S2A) was not observed. These results suggest that Taz1p is first imported into the outer membrane of the mitochondrion and subsequently traffics into the inner membrane or contact sites.

Taz1pV224R is imported into the matrix via the Tim23 pathway. Using yeast as a model, BTHS patients with mutations (V223D, V224R, I226P—equivalent mutations in yeast Taz1p) in the membrane anchor (residues 215 to 232) of Taz1p display an altered protein import pathway in which the mutant protein is imported to the matrix (6). This altered topology suggests that the membrane anchor is important for trafficking in mitochondria. We therefore investigated the role of the membrane anchor in Taz1p assembly by characterizing the import pathway of one representative mutant, Taz1p^{V224R} (Fig. 4). The mutant Taz1p^{V224R} was imported into isolated mitochondria, followed by osmotic shock to disrupt the mitochondrial outer membrane and generate mitoplasts. Centrifugation was used to separate the soluble intermembrane space from the mitoplasts (inner membrane compartment with matrix and intermembrane space proteins that bind to the inner membrane or outer membrane). In WT mitochondria, Taz1p localized to the mitoplast (P) fraction after osmotic shock but was sensitive to protease, confirming a localization facing the intermembrane space (Fig. 4A). In contrast, Taz1p^{V224R} was protected from protease after osmotic shock, indicating a localization facing the matrix (Fig. 4A). As a control, the treatment with Triton X-100 confirmed that Taz1p^{V224R} was sensitive to protease.

The altered topology of Taz1p^{V224R} suggests that it may be imported independently of Tim10p and rely on the TIM23 import

pathway. Indeed, Taz1p^{V224R} import was dependent on a membrane potential and strongly decreased in *tim23-6* mutant mitochondria (Fig. 4B). In addition, Taz1p^{V224R} import bypassed the requirement for Tim10p because import was restored in the *tim10-1* mitochondria and an additional mutant that contains a suppressing mutation in Tim9p, designated *tim10-1tim9^S* (Fig. 4C) (31). Therefore, the imported mutant is likely found on the inner leaflet of the inner mitochondrial membrane, facing the matrix.

Finally, we investigated assembly of imported Taz1p using blue native gels in WT mitochondria. Taz1p and Taz1p^{V224R} were imported and solubilized in 1% digitonin, followed by centrifugation to remove insoluble material. As reported previously (39), Taz1p assembles in several complexes, including three complexes at approximately 70, 120, and 240 kDa in WT mitochondria (Fig. 4D). However, Taz1p^{V224R} only assembled in the smallest complex (70 kDa) and failed to assemble in larger complexes. Taken together, the import pathway of mutant Taz1p^{V224R} was altered compared to that of wild-type Taz1p, which supports the notion that sequences in the membrane anchor are important for interacting with Tim10p.

The membrane anchor of Taz1p is required for import. We constructed a version of Taz1p with a deletion in the membrane anchor, designated Taz1p Δ MA²¹⁵⁻²³², and investigated import *in vitro* and *in vivo*. When Taz1p Δ MA²¹⁵⁻²³² was expressed from a multicopy plasmid in a strain lacking Taz1p (Δ *taz1*), the mutant protein was detected at a lower molecular mass faintly in a yeast lysate (Fig. 5A); in contrast, wild-type Taz1p was readily detected when expressed from the same plasmid in the Δ *taz1* strain. The yeast lysate was fractionated into a mitochondrial enriched pellet (M) and the postmitochondrial supernatant (PMS) (Fig. 5B). Whereas Taz1p was detected in the mitochondrial fraction, Taz1p Δ MA²¹⁵⁻²³² was not detected after fractionation. However, an RNA transcript for Taz1p Δ MA²¹⁵⁻²³² was detected (see Fig. S3 in the supplemental material). Thus, Taz1p Δ MA²¹⁵⁻²³² is not a stable protein.

From the aforementioned evaluation, it was not possible to determine if Taz1p Δ MA²¹⁵⁻²³² was targeted to mitochondria.

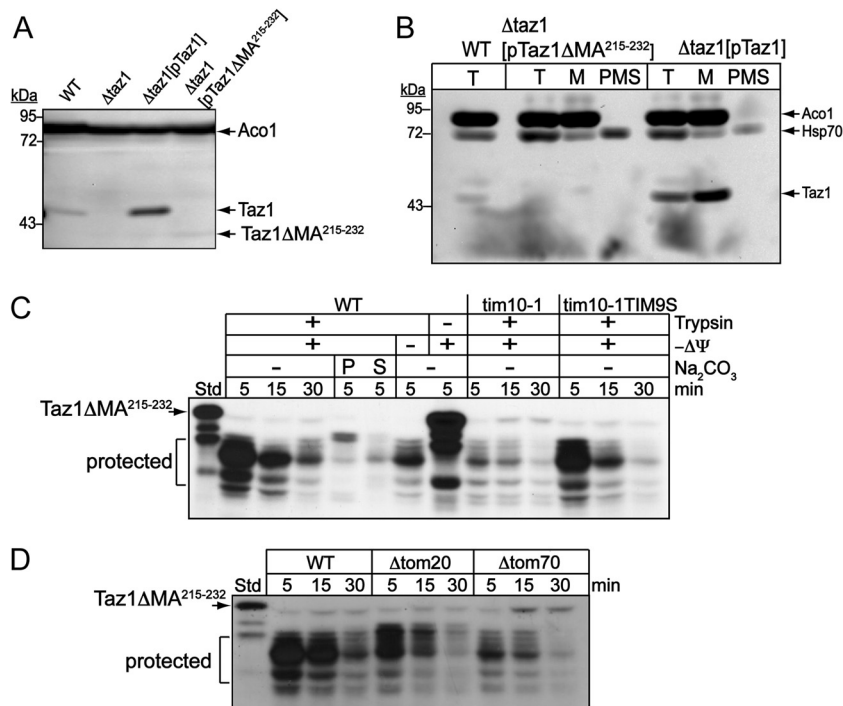


FIG 5 The membrane anchor of Taz1p is required for mitochondrial localization. (A) Taz1p Δ MA^{215–232}, which lacks the membrane anchor, was expressed in a Δ taz1 strain and the truncated protein was detected by immunoblot analysis in a total protein lysate. The immunoblot analysis of WT TAZ1 and aconitase (Aco1) serve as controls for blotting and loading, respectively. (B) The lysate (T) of strains from panel A was fractionated into the M and PMS fractions, followed by immunoblotting. Cytosolic Hsp70 served as an additional control for fractionation. (C) Taz1p Δ MA^{215–232} was imported into isolated WT, *tim10-1*, and *tim10-1tim9^S* mitochondria and subjected to trypsin treatment to remove nonimported precursor. Samples in WT mitochondria were also subject to carbonate extraction (Na₂CO₃) and separated in supernatant (S) and pellet (P) fractions by centrifugation. Fragments that were initially resistant to protease were labeled “protected.” (D) Taz1p Δ MA^{215–232} was imported as for panel C into WT, Δ tom20, and Δ tom70 mitochondria.

Consequently, we investigated whether Taz1p Δ MA^{215–232} was imported into isolated WT mitochondria (Fig. 5C). When Taz1p Δ MA^{215–232} was imported and treated with protease to assess translocation across the outer membrane, a reproducible pattern of protected Taz1p Δ MA^{215–232} initially appeared within the first 5 min of import, but it disappeared by the later time point of 30 min (Fig. 5C). The protection pattern resulted from trypsin addition that was added to remove nonimported precursor, suggesting that the Taz1p Δ MA^{215–232} associated with mitochondria but was not imported completely into the intermembrane space. However, Taz1p Δ MA^{215–232} was not inserted into the membrane, because Taz1p Δ MA^{215–232} was not recovered from the pellet after carbonate extraction. Furthermore, this protection pattern was not present in *tim10-1* mitochondria, which lack functional Tim9p-Tim10p complex. However, Taz1p Δ MA^{215–232} showed the same protease protection in mitochondria from the suppressor *tim10-1tim9^S* strain (Fig. 5C).

As we previously showed that Tom70p and Tom20p were important for Taz1p translocation, we investigated the import of Taz1p Δ MA^{215–232} in Δ tom70 and Δ tom20 mitochondria (Fig. 5D). There was a decreased association of Taz1p Δ MA^{215–232} in these mutant mitochondria compared with the WT control. Taken together, the results show that Taz1p Δ MA^{215–232} was targeted to the TOM complex and engaged the Tim9-Tim10p complex but was not translocated across and into the outer membrane. The precursor may become initially “trapped” in the TOM complex, which renders a portion of the substrate protected from pro-

tease; however, because of incomplete translocation, the substrate may dissociate from mitochondria at the later time points.

The membrane anchor and flanking sequences are sufficient for import of Taz1p. To determine if the membrane anchor of Taz1p contained sufficient information for targeting to mitochondria, a series of DHFR fusion proteins were constructed and imported into wild type mitochondria (Fig. 6A). To ensure that there was adequate flexibility of the protein sequence for engaging the mitochondrial translocons, a linker sequence of NAAIRS flanked the N terminus and a glycine linker (GGGGG) flanked the C terminus of the Taz1p sequence, followed by DHFR fusions on both sides of the Taz1p region. Constructs that contained the membrane anchor (residues 215 to 232) as well as flanking sequences were included. For the constructs that contained only the membrane anchor (DHFR-MA^{215–232}-DHFR) or the membrane anchor with a C-terminal flanking region (DHFR-MA^{215–265}-DHFR), import was not robust. In contrast, constructs that contained the membrane anchor plus the flanking N-terminal region (DHFR-MA^{204–265}-DHFR and DHFR-MA^{204–232}-DHFR) showed strong import. Thus, the membrane anchor and adjacent N-terminal sequence, with several charged amino acid residues, seem to direct Taz1p import.

We also tested the ability of the V224R mutation and flanking sequences to direct import by generating a similar DHFR fusion construct that contained the membrane anchor with the V224R mutation and both N-terminal and C-terminal flanking sequences (DHFR-MA^{204–265/V224R}-DHFR) (Fig. 6B). Again, this region conferred import into mitochondria. Furthermore, the mi-

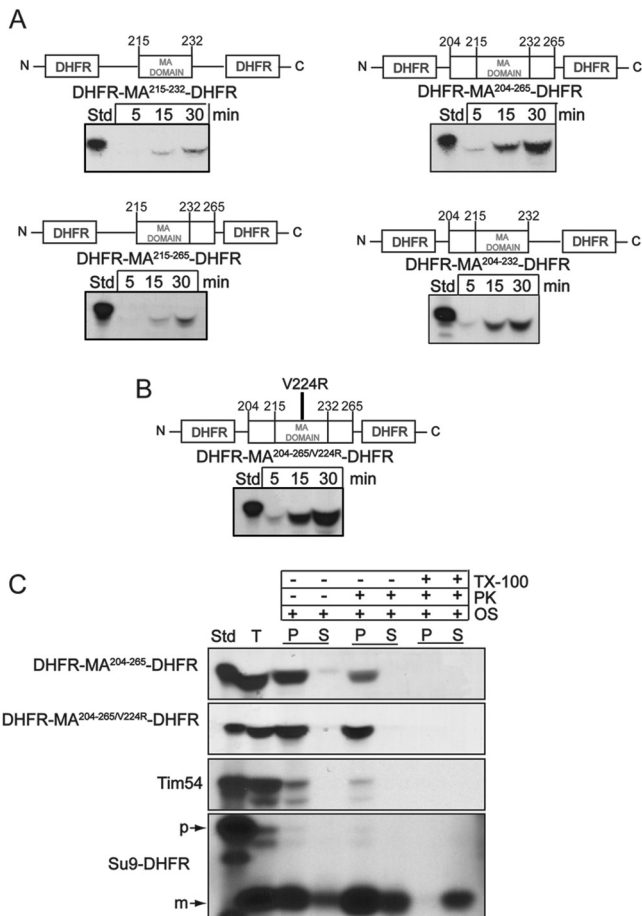


FIG 6 The membrane anchor region and its flanking sequences are sufficient for mitochondrial import. (A) Constructs were made in which the membrane anchor (MA) of Taz1p (amino acids 215 to 232) and flanking sequences (amino acids 215 to 265, 204 to 265, and 204 to 232) were inserted between two DHFR moieties and imported into isolated WT mitochondria. Nonimported precursor was removed by protease treatment. (B) The V224R mutation was introduced, creating DHFR-MA^{204-265/V224R}-DHFR, and imported as for panel A. (C) Imported DHFR-MA²⁰⁴⁻²⁶⁵-DHFR and DHFR-MA^{204-265/V224R}-DHFR were localized within mitochondria by osmotic shock in the presence and absence of proteinase K and Triton X-100. Controls include Tim54p for the intermembrane space and Su9-DHFR for the matrix.

tochondrial localization was tested with osmotic shock in the presence of protease (Fig. 6C) to assess whether the construct was imported into the matrix or intermembrane space. As predicted, DHFR-MA²⁰⁴⁻²⁶⁵-DHFR was imported into the intermembrane space, but DHFR-MA^{204-265/V224R}-DHFR was imported into the mitochondrial matrix. As a control, imported Tim54p was detected in the intermembrane space, but Su9-DHFR localized to the matrix. These fusion constructs confirm that the membrane anchor sequence alone is not sufficient for targeting to the mitochondria, but critical targeting information is contained N terminal to the transmembrane domain.

DISCUSSION

The range and variety of Taz1p mutations that result in Barth syndrome hint at both the importance and the complexity of this protein and its mitochondrial function (40). Taz1p has a complex localization pattern in mitochondria such that it localizes to the

inner and outer membranes and lines the intermembrane space as an interfacial membrane protein (6). In contrast, cardiolipin synthase has been localized to the inner leaflet of the inner membrane (13). A hot spot for mutations seems to be the membrane anchor domain, and mutations in this region result in mislocalization of Taz1p to the mitochondrial matrix (6). Together, these findings warranted further investigation of the biogenesis of Taz1p, especially to develop strategies to understand BTHS.

This study has shown that Taz1p first binds the TOM complex and that Tom20p and Tom70p are both required. It was important to first implicate the TOM receptors, because some outer membrane proteins, such as the proapoptotic Bcl-2 proteins and various alpha-helical signal- and tail-anchored proteins, may be inserted into the outer membrane without passing through the TOM40 channel (41, 42). Furthermore, this study supports the notion that Taz1p is imported via the TOM translocator because Taz1p was cross-linked to Tom40p during translocation. However, the inner membrane translocons seem to be surprisingly bypassed, because the wild type Taz1p showed no import defects in the *tim22-4* or *tim23-6* strains, and Taz1p import was independent of a $\Delta\Psi$.

The endogenous Taz1p protein resides in both the outer and inner membranes of the mitochondrion, as shown with sucrose gradients, immunoelectron microscopy (6), and, as shown here, a digitonin solubilization assay. The evidence presented in this study indicates that tafazzin first integrates into the outer membrane. Therefore, how Taz1p migrates from the outer membrane to the inner membrane, which is the site of CL synthesis and enrichment and the submitochondrial residence of two binding partners of tafazzin, the ATP synthase and AAC (39), is an open question. Perhaps the enzyme moves through contact sites between the two membranes or is transferred with the lateral migration of phospholipids. Recent studies support the notion that a multisubunit protein complex (termed MitOS/MINOS/MICOS) has a scaffold-like structure to connect outer and inner membranes (43–45), providing a potential conduit for lipids and Taz1p trafficking from the outer to the inner membrane. From the experimental approach in which imported Taz1p was fractionated with digitonin, we attempted to extend the length of the import reaction from 30 min up to 2 h to chase the Taz1p precursor into the inner membrane, but we did not observe a difference in the precursor's distribution profile. It may be reasonable to assume that normal lipid trafficking and diffusion do not occur or cannot be recapitulated in purified mitochondria, even with a $\Delta\Psi$.

Taz1p interacts with the Tim9-Tim10p complex after it passes the TOM complex. First, a noticeable import defect was shown with mitochondria from the *tim10-1* strain, and Taz1p import was restored in the suppressor strain *tim10-1Tim9S*, in which the mutation in Tim9p restores function of the Tim9p-Tim10p complex (31). Second, the Taz1p precursor was cross-linked to Tim10p during translocation. Endogenous Taz1p had an altered digitonin solubilization profile in the *tim10-1* strain, which we interpreted as an inability to correctly insert into the outer mitochondrial membrane due to the lack of functional Tim10p. This may reveal a role for the Tim9-Tim10p complex post-outer membrane insertion in facilitating the migration of Taz1p to the inner membrane. In the absence of the Tim9-Tim10p complex, Taz1p may be inserted into the outer membrane but not traffic to the inner membrane. The Tim9-Tim10p complex is known to mediate insertion of components of the SAM and TOM40 complex into the outer

membrane (17), but tafazzin is one of the few non- β -barrel outer membrane proteins to use the Tim9p-Tim10p complex. Additionally, unlike the β -barrel proteins, tafazzin does not require the SAM complex for membrane integration (15), indicating that Taz1p uses a new mitochondrial import pathway. Whether or not other proteins mediate the insertion of Taz1p into the lipid bilayer is currently unknown and requires additional investigation.

When a single amino acid in the membrane anchor is mutated (V224R), Taz1p alters its normal import course and does not reach the outer mitochondrial membrane. Instead, it interacts with the TIM23 translocon and is pulled into the matrix, where it likely resides on the inner leaflet of the inner membrane. The mislocalization event was seen by subjecting mitochondria to osmotic shock and treating with protease, both *in vivo* (6) and, in this current study, with an *in vitro* import assay. We hypothesize that the point mutation, in which a hydrophobic valine is changed to a charged arginine residue, changes the hydrophobicity of this crucial region and interferes with tafazzin's ability to interact with the Tim9p-Tim10p complex, which is known to bind to the hydrophobic regions of membrane proteins and protect them from the aqueous environment of the intermembrane space. In a default pathway, Taz1p is then passed to the TIM23 translocon, which perhaps supports the model that the TIM23 translocon interacts directly with the TOM complex (46). Alternatively, membrane integration may be required for Tim9p-Tim10p complex association with Taz1p. Because the membrane anchor and flanking regions of Taz1p contain several mutations found in BTHS patients (40) and mitochondrial membrane proteins typically have targeting and sorting information in or immediately adjacent to the membrane domains (20), we subsequently focused on the membrane anchor and its role in mitochondrial targeting and sorting. The membrane anchor and the N-terminal region are required for mitochondrial targeting and sorting. When construct Taz1 Δ MA²¹⁵⁻²³² that lacks the membrane anchor was imported into mitochondria, it initially engaged the TOM translocon and was partially protease protected (Fig. 5), but then the construct seemed to disengage from mitochondria because it was subsequently degraded by protease. Moreover, the protection from protease was dependent on Tim10p. We suggest that the Tim9-Tim10p complex binds to the membrane anchor of Taz1p. When Taz1 Δ MA²¹⁵⁻²³² was expressed in a Δ taz1p yeast strain, the protein was quickly degraded by cellular proteases, because the mutant protein was poorly detected in whole-cell extracts and not detected in an enriched mitochondrial fraction. These experiments postulate that the membrane anchor region not only enables the endogenous protein to integrate tightly into the lipid bilayer of the mitochondrial membranes, making it resistant to alkali and high salt extractions, but also is necessary if the protein is to be accurately sorted by the Tim 9/10 import protein machinery. Similar to inner membrane carriers (47), Taz1p likely inserts as a loop across the TOM complex, because appended DHFR at the N and C termini did not inhibit import. Given the abundance of mutations in the membrane anchor and flanking sequences, this study suggests that the molecular basis for BTHS in a subset of patients is caused by defects in mitochondrial targeting and localization. Thus, detailed analysis of Taz1p biogenesis in *Saccharomyces cerevisiae* provides a framework for further studies to understand the molecular basis of BTHS.

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