Molecular insights into the \textit{m-AAA} protease–mediated dislocation of transmembrane helices in the mitochondrial inner membrane

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Protein complexes involved in respiration, ATP synthesis, and protein import reside in the mitochondrial inner membrane; thus, proper regulation of these proteins is essential for cell viability. The \textit{m-AAA} protease, a conserved hetero-hexameric AAA (ATPase associated with diverse cellular activities) protease, composed of the Yta10 and Yta12 proteins, regulates mitochondrial proteostasis by mediating protein maturation and degradation. It also recognizes and mediates the dislocation of membrane-embedded substrates, including foreign transmembrane (TM) segments, but the molecular mechanism involved in these processes remains elusive. This study investigated the role of the TM domains in the \textit{m-AAA} protease by systematic replacement of one TM domain at a time in yeast. Our data indicated that replacement of the Yta10 TM2 domain abolishes membrane dislocation for only a subset of substrates, whereas replacement of the Yta12 TM2 domain impairs membrane dislocation for all tested substrates, suggesting different roles of the TM domains in each \textit{m-AAA} protease subunit. Furthermore, \textit{m-AAA} protease–mediated membrane dislocation was impaired in the presence of a large downstream hydrophilic moiety in a membrane substrate. This finding suggested that the \textit{m-AAA} protease cannot dislocate large hydrophilic domains across the membrane, indicating that the membrane dislocation probably occurs in a lipid environment. In summary, this study highlights previously underappreciated biological roles of TM domains of the \textit{m-AAA} proteases in mediating the recognition and dislocation of membrane-embedded substrates.

Maintaining protein homeostasis at the mitochondrial IM is thus essential and critical for cellular fitness. The \textit{m-AAA} protease and the \textit{i-AAA} protease form a major quality control system in the mitochondrial IM and mediate the degradation of misfolded or unassembled mitochondrial IM proteins (1–4). The \textit{i-AAA} protease is a homo-oligomeric complex formed by Yme1, and the catalytic domain is exposed to the intermembrane space (IMS) (5, 6). The \textit{m-AAA} protease is a heterohexameric complex composed of Yta10 and Yta12; each contains two N-terminal TM2, soluble AAA+ domains, and a proteolytic domain exposed to the matrix (7, 8).

The function of the \textit{m-AAA} proteases is not limited to quality control, but it also assists with protein folding of a mitochondrial ribosome subunit, MrpL32 (9), and maturation of cytochrome \textit{c} peroxidase (Ccp1) (10). In the case of Ccp1 maturation, the \textit{m-AAA} protease acts as an ATPase by pulling the Ccp1 TM to the matrix prior to processing by a rhomboid protease, Pcp1, in the IM (10, 11). Furthermore, an earlier study has shown that moderately hydrophobic TM segments can be dislocated by the \textit{m-AAA} protease (12).

It has been reported that deletion of TMs of Yta10 or Yta12 causes severe defects in the degradation of integral membrane proteins but only mild defects in the degradation of peripherally associated membrane proteins, implicating the importance of TMs of the \textit{m-AAA} protease in extracting the substrate TM segments from the membrane (13–15). However, it is poorly understood whether the \textit{m-AAA} protease TMs are actively involved in the dislocation process or simply act to anchor the catalytic domain near the membrane.

To better understand the molecular mechanism of the \textit{m-AAA} protease–mediated TM helix dislocation, this study focused on the role of TMs of the \textit{m-AAA} protease. Individual TMs of Yta10 and Yta12 were replaced one at a time, and the capacity of these \textit{m-AAA} protease variants to dislocate various membrane substrates was assessed. Our data show that replacement of TM2 of Yta10 impairs recognition of a subset of substrates, whereas replacement of Yta12 TM2 impairs membrane dislocation of all tested substrates, illustrating different roles of TMs in each subunit. Additionally, we have investigated the length of a soluble moiety in the IMS of the dislocation substrate the \textit{m-AAA} protease is capable of extracting from the membrane.
Results

Yta10 or Yta12 variants with a replaced TM complement the respiratory growth defect of the corresponding deletion strain

To elucidate the role of individual TMs of the m-AAA protease subunits, each TM of Yta10 and Yta12 was replaced with that of another mitochondrial IM protein, Mdl2, one at a time (Fig. 1A). These constructs were transformed into the yta10Δ or yta12Δ strain. As reported previously, the yta10Δ and yta12Δ strains were viable on fermentable carbon medium but could not grow on non-fermentable carbon medium, as the m-AAA protease is required for the maturation of the mitochondrial ribosome subunit MrpL32 (16).

To examine whether Yta10 and Yta12 variants are functional, a growth complementation assay was performed (Fig. 1B). All of the constructs rescued the growth defect of the corresponding deletion on non-fermentable medium. Furthermore, MrpL32 maturation in the Yta10 and Yta12 variants were unaffected (Fig. 1C). These results suggest that Yta10 or Yta12 variants with a replaced TM are correctly targeted, assembled into the m-AAA protease complex, and processed by MrpL32 in the mitochondria.

Dislocation substrates for the m-AAA protease

Mgm1 is an IM protein that naturally exists in two forms: l-Mgm1 and s-Mgm1. It carries an N-terminal presequence that is processed by mitochondrial processing peptidase (MPP) and two putative TMs. A recent report has shown that Mgm1 is co-translationally imported into mitochondria (17). Uniquely, the insertion efficiency of the TM1 into the IM is roughly 50% (18–20). When the TM1 is inserted into the IM, it becomes l-Mgm1. On the other hand, when the TM1 of Mgm1 is translocated to the matrix, the TM2 subsequently enters the IM, where it is further processed by Pcp1, and s-Mgm1 is formed (Fig. 2A, left). Here, the sorting of the TM1 of Mgm1 occurs at the level of the TIM23 complex.

However, when the TM1 of Mgm1 is replaced with the model TM segment composed of varying number of alanines and leucines (19-n alanines and n leucines), this segment is inserted...
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Figure A

Figure B

Figure C
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into the IM by the TIM23 complex, resulting in l-Mgm1 (21). Subsequently, l-Mgm1 can be dislocated by the m-AAA protease (12) (Fig. 2A, right). The dislocation efficiency has been shown to be dependent on the hydrophobicity; moderately hydrophobic TM segments are more readily dislocated than sufficiently hydrophobic ones (12). Thus, for the Mgm1 constructs carrying the engineered TM segments (Mgm1(A/L) constructs), the generation of s-Mgm1 is dependent on the dislocation activity of the m-AAA protease, as the cleavage of the TM2 by Pcp1 only occurs after dislocation of the TM1 by the m-AAA protease.

Replacement of TM2 of Yta10 and Yta12 impairs membrane dislocation of model TM segments in Mgm1

To assess the membrane dislocation efficiency of Yta10 and Yta12 TM mutants, the yta10Δ or yta12Δ strain carrying each Yta10 and Yta12 variant was transformed with a set of Mgm1(A/L) constructs. We reasoned that, if the role of Yta10 or Yta12 TMs is limited to anchoring the catalytic domain (ATPase domain and the metalloprotease domain) near the membrane, then substitution of TMs would have a minimal effect. In contrast, if the TMs of Yta10 or Yta12 are required for the recognition or active membrane dislocation, then substitution of Yta10 TMs would compromise the membrane dislocation activity of m-AAA protease.

Replacement of TM1 (RTM1) of Yta10 or Yta12 did not have significant effects on the dislocation of the tested TM segments, as the ratio of l-Mgm1 to s-Mgm1 was comparable with the Yta10 or Yta12 WT strain (Fig. 2, B and C). Although subtle, a decrease in membrane dislocation of a sufficiently hydrophobic TM (Mgm1(14A/5L)) has been observed in both the Yta10 and Yta12 RTM1 variants (Fig. 2, B and C). In comparison, replacement of TM2 (RTM2) of Yta10 or Yta12 significantly reduced generation of the s-Mgm1 form, indicating that the membrane dislocation activity is severely impaired upon swapping of TM2 of the m-AAA protease (Fig. 2, B and C).

Replacement of TM2 of Yta10 selectively impairs membrane dislocation for only a subset of substrates

To distinguish whether the replacement of Yta10 or Yta12 TM2 causes a general defect in membrane dislocation activity or in recognizing substrates, other previously characterized m-AAA protease substrates were tested. Ccp1 is a natural substrate of the m-AAA protease. For maturation of Ccp1, precursor Ccp1 undergoes two-step cleavage by the m-AAA protease in the matrix and Pcp1 in the IM (Fig. 3A). Replacement of its TM1 with the model TM segment composed of 15 alanines and four leucines (Ccp1(15A/4L)) does not affect its maturation (12). For the Ccp1 constructs, membrane dislocation efficiency is measured by the production of mature Ccp1(m-Ccp1) (Fig. 3A, left).

Cox5aT-MFP (MFP, Mgm1 fusion protein) is a chimera protein where Cox5a protein is truncated at residue 128 and fused to the C terminus of Mgm1 (22). Previous studies have shown that this C-terminal truncation at the IMS side converts Cox5a into the m-AAA protease substrate. Another m-AAA protease substrate, Cox5aT(15A/4L)-MFP, was prepared by replacing the Cox5a TM with the TM segment composed of 15A/4L in the background of Cox5aT-MFP. The membrane dislocation efficiency of Cox5aT-MFP variants was judged by the formation of s-Mgm1 (Fig. 3A, right).

First, the membrane dislocation efficiency of Ccp1 and Cox5aT-MFP variants in cells carrying the Yta10 variants was assessed. In contrast to the results with Mgm1(A/L) variants, the generation of m-Ccp1 or s-Mgm1 in Yta10 RTM2 strain was comparable with that of the Yta10 WT strain, indicating that dislocation of the Ccp1 and Cox5aT-MFP variants was unaffected by the TM2 replacement of Yta10 (Fig. 3B). These results thus implicate that replacement of Yta10 TM2 does not impair the general membrane dislocation activity of the m-AAA protease.

Replacement of TM2 of Yta12 causes a general defect in membrane dislocation activity of the m-AAA protease

Next, the dislocation efficiency of the Ccp1 and Cox5aT-MFP variants was tested in the Yta12 strains. Here, the generation of m-Ccp1 or s-Mgm1 was significantly reduced compared with the WT strain, indicating that the membrane dislocation of tested substrates was severely impaired (Fig. 3C). This suggests that replacement of TM2 of Yta12 causes a general defect in the membrane dislocation activity of the m-AAA protease, different from the effects of the TM2 replacement of Yta10.

Dislocation of a TM segment with a hydrophilic IMS moiety is inhibited in a length-dependent manner

When the m-AAA protease mediates degradation of a membrane protein, not only the TMs but also the loop regions in the IMS should be pulled across the IM. How large can the soluble domain be for the m-AAA protease to handle? To determine the dislocation capacity of the m-AAA protease, a hydrophilic segment containing many polar and charged residues from the Escherichia coli leader peptidase periplasmic (P2) domain was introduced in between the two TMs of Mgm1(15A/4L) at various lengths (Fig. 4A).

Dislocation of TM1 of Mgm1(15A/4L) was gradually decreased by 24- and 50-amino acid-long extensions and completely inhibited by 105-amino acid-long extensions (Fig. 4B). An earlier study has shown that a tightly folded domain in downstream of the TM segment prevents membrane protein degradation by the m-AAA protease (15, 23). To check whether the hydrophilic segments used in our experiment form a tightly

Figure 2. Dislocation of Mgm1(A/L) from the IM requires TM2 of Yta10 and Yta12. A, alternative topogenesis of Mgm1 WT and processing of Mgm1 (A/L). The sequences of the TM1 segment of Mgm1, Mgm1(16A/3L, 15A/4L, 14A/5L, 13A/6L), are shown. B, dislocation of Mgm1(A/L). Yeast transformants expressing the indicated Yta10 variant and Mgm1(A/L) were lysed in the presence of 25% TCA (final concentration). The lysates were analyzed by SDS-PAGE and Western blotting. The blots were immunodecorated with anti-HA antibody, which detects the HA tag at the C terminus of Mgm1(A/L) (left panel). Relative amounts of s-Mgm1 were quantified from three independent experiments and plotted with standard deviations. A t test was performed to examine the statistical significance of the observed results (right panel). C, Yta12 variants were tested in the same manner as described in B. The differences in dislocation efficiency were statistically significant between WT and EV/RTM2 (*, p < 0.05).
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folded structure, the structure of the P2 domain was examined (PDB code 1T7D) (24). 24- and 50-amino acid-long segments are mostly unstructured, and a 105-amino acid segment consists of some β strands and unstructured loops. Thus, it is unlikely that membrane dislocation was prevented because of the folding structure of the hydrophilic extension. These results suggest that the m-AAA protease has limited capacity to dislocate a large IMS moiety.

Drastic dislocation impairment in the presence of a hydrophilic linker led us to think about the molecular environment from which the proteins are dislocated. We reasoned that, if dislocation occurs through the hydrophobic core of the lipid bilayer, then a large sum of energy is required; therefore, a long hydrophilic extension would prevent membrane dislocation, whereas, if it occurs through a proteinaceous pore, then the energy barrier would be lower, and the extension length would influence the dislocation efficiency less. To test the translocation efficiency of the P2 domain through the proteinaceous channel, increasing lengths of hydrophilic extensions were added in between the TM1 and the downstream Pcp1 cleavage site in Mgm1 WT (Fig. 4A). For the generation of s-Mgm1 in Mgm1 WT, the TM1 is translocated through a pore formed by the TIM23 complex (Fig. 2A); thus, Mgm1 WT with the hydrophilic domain extension could serve as a control for membrane dislocation through a proteinaceous environment. Here, the relative ratio of s-Mgm1 to J-Mgm1 was unaffected by the presence of a 24-, 50-, or 105-amino acid-long hydrophilic segment, demonstrating that addition of a hydrophilic moiety between the two TMs did not interfere with the translocation/insertion efficiency of Mgm1 (Fig. 4B). Further, this result showed that
the presence of a long hydrophilic moiety upstream of the rhomboid cleavage site did not impair the cleavage efficiency of Pcp1. Taken together, these results suggest that the \( m \)-AAA protease is incapable of dislocating a large hydrophilic domain across the IM, subsequently implicating that membrane dislocation probably occurs through the lipid environment.

**Discussion**

The \( m \)-AAA TMs have been shown to be indispensable for integral membrane protein degradation (13), but their roles in extracting proteins out of the membrane remain poorly understood. By swapping each TM one at a time, we present data showing that replacement of the TM2s of \( m \)-AAA protease causes defects in the membrane dislocation of substrate proteins. Interestingly, replacement of TM2 of Yta10 and Yta12 exhibited different defects in substrate dislocation. Replacement of TM2 of Yta10 impaired the dislocation of Mgm1(A/L) variants only, whereas replacement of TM2 of Yta12 impaired the dislocation of all tested substrates.

The general dislocation impairment observed with TM2 replacement of Yta12 was unexpected because Ccp1 dislocation/maturation was shown to be unaffected in a yeast strain expressing a TM deletion mutant of Yta12 (13). Therefore, it is less likely that TM2 of Yta12 is dispensable for membrane protein dislocation or substrate recognition. Why would TM2 replacement cause a more severe defect than TM deletion in membrane dislocation? Augustin et al. (25) have shown that coordinated intersubunit signaling between AAA+ domains of Yta12 and Yta10 is especially critical for membrane dislocation. TM2 of Yta12 is located upstream of the AAA+ domain (Fig. 1A). If the TM2 replacement causes subtle misalignment between the AAA+ domains of Yta12 and Yta10, then it could impair efficient ATP hydrolysis relay and subsequent membrane dislocation. However, in the complete absence of a TM, the AAA+ domain of Yta12 could be better positioned via interaction with the AAA+ domain of Yta10.

Compared with Yta12, TM2 replacement of Yta10 impaired the membrane dislocation of Mgm1(A/L) variants only. Because dislocation of the Ccp1 and Cox5aT-MFP variants is normal, the dislocation function of the \( m \)-AAA protease is not abolished. Thus, these results imply that replacement of TM2 of Yta10 causes impairment in the process of selective substrate recognition.

Because Yta10 and Yta12 share high sequence homology (supplemental Fig. S1), at present we cannot exclude the possibility that TM2 of Yta10 and Yta12 has a redundant function in the recognition of membrane substrates. Although most substrates need one or the other, for some substrates, such as Mgm1(A/L), require both. In the case of the \( i \)-AAA protease Yme1, it has been shown that both N- and C-terminal helical domains serve to recognize different degradation substrates (26, 27). Given that the \( m \)-AAA protease plays a diverse role in mitochondrial proteostasis, it would not be surprising if it has
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multiple modes of substrate recognition. Further study is needed to clarify this issue.

Next, we probed the length of the hydrophilic polypeptides the m-AAA protease can dislocate through the membrane by introducing different lengths of hydrophilic linker in the m-AAA protease substrate. Our results show that the dislocation efficiency is decreased significantly as the hydrophilic extension lengthens. We reason that, if the dislocation occurs in a proteinaceous pore, then it would be less sensitive to the length of the hydrophilic moiety; however, if the dislocation occurs in the lipid bilayer, then the presence of a long hydrophilic segment would be energetically too costly. Hence, these results implicate that the m-AAA protease dislocates a TM segment from the lipid environment and that it would be incapable of dislocating a large IMS domain across the membrane. Subsequently, it can be predicted that, for the m-AAA protease to extract IM proteins for degradation in the matrix, either the IMS domain should be small or the large IMS domain should be cleaved off by corroborating with other proteases prior to extraction from the membrane.

Experimental procedures

Yeast strains

W303-1a (MATa, ade2, can1, his3, leu2, trp1, and ura3) was used as a parental strain in this study. yta10Δ (MATa, ade2, can1, his3, leu2, trp1, ura3, yta10::HIS3MX6) was made by standard homologous recombination, substituting YTA10 with an amplified HIS3MX6 (28). The primers used for the amplification of the HIS3MX6 cassette were 5′-CAGCGTTT-GGACACGTATTATCCTCGATCCCCGGTATTAAAC-3′ and 5′-TTGGGTAGAACGTTATCGAATGAGCAGCTCGTTTAAAC-3′. yta10Δ was used to make yta10Δyta12Δ (MATa, ade2, can1, his3, leu2, trp1, ura3, yta10::HIS3MX6, yta12::KanMX6) by substituting YTA12 against KanMX6. The primers used for the amplification of the KanMX6 cassette were 5′-TATCGGTTCCATATAAGAAAGTC-3′ and 5′-GCCCTTAAGATGACCTACGTTTATT-3′. These yeast strains were cultured at 30 °C in this study.

Plasmid construction

YTA10 was amplified from the genomic DNA using the set of primers 5′-ATAGGGCGAATTGGAAGGTCCACCCGGTGGT-GCGTTGTGACATATATGCTC-3′ and 5′-TATCGGATAA-GTTGATATCGATTTCCGAGGTTATAATGAGAAGGTGTT-3′. YTA12 was amplified from the genomic DNA using the set of primers 5′-GGTGGGAGCGCCGCTTAGAAGAATGATTGCGATCATACATATATATGCTC-3′ and 5′-ATCGATAGCTGATTATCGAATGAGGAGGACAAAAATGATCGTTT-3′. The set of primers amplified YTA10 and YTA12 with 1 kb upstream sequences and 500 bp downstream sequences to include endogenous promoters and potential transcriptional regulators and contained nucleotides for cloning the amplified product into pRS314 or pRS316, respectively, by homologous recombination ([pRS314 YTA10 WT], [pRS316 YTA12 WT]).

The TMs of Yta10 were replaced with a TM of Mdl2 using [pRS314 YTA10 WT] as template for site-directed mutagenesis. Two sets of primers were used to replace TMs of Yta10 with a TM of Mdl2 using [pRS314 YTA10 WT] as template for site-directed mutagenesis. The Mgm1, Ccp1, and Cox5aT model proteins used in previous studies were adopted (12, 29). To make Mgm1 with a P2 domain inserted between residues 130 and 131 of Mgm1 WT and Mgm1(A/L), overlap PCR was performed (30). First, the N-terminal part of Mgm1 WT and Mgm1(A/L) was amplified using 5′-TGTTACGAGAGTATGGTACATTCAAGATGTAATTCTACCTCTTCAAAGTG-3′ and 5′-TTAAATCCTTACTCAGTTATTTTAAAGG-3′. The C-terminal part of Mgm1 WT was amplified using 5′-GGTGAATCGATGAAGGAAAAG-3′ and 5′-TTAGAGCGTATACGGAAAC-3′.

The P2 domain was amplified from LepH2 protein (31) using the primer 5′-AACACTAGATCGACGTTATATGATA-TCTGTTCTCCAGGACGTTTAC-3′ in combination with 5′-CTTTTCTCT-CTATCATCGAAGGAAAGATCGCTCGTTTC-3′ or 5′-CTTCT CCTTCATCGAAGGAAAGATCGCTCGTTTC-3′ or 5′-CTTTTCTCCTATCGAAGGAAAGATCGCTCGTTTC-3′. The set of primers amplified the plasmid pH84HA plasmid (32) by homologous recombination. The N-terminal part of Mgm1 WT and Mgm1(15A/4L) was each fused to the P2 domain by overlap PCR, and the stitched products were further annealed to the C-terminal part of Mgm1 WT. The resulting PCR product was cloned into the pH84HA plasmid (32) by homologous recombination.

Growth assay

yta10Δ expressing [pRS314 YTA10 variants] were cultured in -Trp(glucose) medium overnight at 30 °C. The overnight culture was diluted to 0.1 A600 and cultured to 0.5 A600 at 30 °C. 10 μl of culture was spotted on YPD (yeast extract peptone dextrose) or YPEG (yeast extract peptone ethanol, glycerol) plate and further incubated for 2 days at 30 °C. Photos were taken on a Chemi-doc-XRS+ system using Epi Light (Bio-Rad).
**Protein preparation and Western blotting**

Preparation of lysates, SDS-PAGE, and Western blotting were conducted as described previously (22). Briefly, yeast transformants were grown overnight in 5 ml of synthetic complete medium at 30 °C. Proteins were precipitated from 1 Aopt unit of yeast cells by addition of trichloroacetic acid (TCA) (Sigma). Precipitated proteins were resuspended in 40 μl of sample buffer and incubated for 5 min at 95 °C prior to SDS-PAGE. The samples were separated on 6.5% or 12.5% Tris-HCl gels (Bio-Rad) and followed by Western blotting. Membranes were immunodecorated with an anti-HA antibody (Covance) and developed with the Amersham Bioscience advanced ECL kit on a Chemi-doc-XRS+ system (Bio-Rad). Quantification of detected bands was done using Image Lab 5.0 (Bio-Rad).

**MrpL32 processing assay**

From overnight cell cultures of yta1A expressing [pRS314 YTA10 variants] and yta10A/yta12 [pRS314 YTA10 WT] expressing [pRS316 YTA12 variants], whole-cell lysates were prepared by resuspending cells in sample buffer (50 mM Tris-HCl, 5% SDS, 5% glycerol, 50 mM DTT, 2 μg/ml leupeptin, 2 μg/ml pepstatin A, 1 μg/ml chymostatin, ECL 15 mg/ml benzamidine, 0.1 mg/ml Pefabloc, 8.8 μg/ml aprotinin, 3 μg/ml Aidapin and bromphenol blue) and heating for 15 min at 60 °C. Prior to gel loading, the samples were centrifuged at 14,000 rpm for 5 min, and the supernatant fractions were loaded onto 15% Tris-HCl gels (Bio-Rad). Western blotting was followed with anti-MrpL32 antiserum (a kind gift from Drs. Langer and Tatsuta from the University of Cologne).

**Author contributions**—H. L. and H. K. conceived the project. H. L., S. L., and S. Y. conducted the experiments. H. L., S. L., and S. Y. designed the study, analyzed data, and wrote the manuscript. H. L., S. L., and S. Y. conducted the experiments.

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