A New Function in Translocation for the Mitochondrial *i*-AAA Protease Yme1: Import of Polynucleotide Phosphorylase into the Intermembrane Space †

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Polynucleotide phosphorylase (PNPase) is an exoribonuclease and poly(A) polymerase postulated to function in the cytosol and mitochondrial matrix. Prior overexpression studies resulted in PNPase localization to both the cytosol and mitochondria, concurrent with cytosolic RNA degradation and pleiotropic cellular effects, including growth inhibition and apoptosis, that may not reflect a physiologic role for endogenous PNPase. We therefore conducted a mechanistic study of PNPase biogenesis in the mitochondrion. Interestingly, PNPase is localized to the intermembrane space by a novel import pathway. PNPase has a typical N-terminal targeting sequence that is cleaved by the matrix processing peptidase when PNPase engaged the TIM23 translocon at the inner membrane. The *i***-AAA protease Yme1 mediated translocation of PNPase into the intermembrane space but did not degrade PNPase. In a yeast strain deleted for Yme1 and expressing PNPase, nonimported PNPase accumulated in the cytosol, confirming an in vivo role for Yme1 in PNPase maturation. PNPase localization to the mitochondrial intermembrane space suggests a unique role distinct from its highly conserved function in RNA processing in chloroplasts and bacteria. Furthermore, Yme1 has a new function in protein translocation, indicating that the intermembrane space harbors diverse pathways for protein translocation.**

Polynucleotide phosphorylases (PNPases) encompass an evolutionarily conserved enzyme family whose members regulate RNA levels in bacteria and plants. In *Escherichia coli*, PNPase associates with an RNA helicase and enolase in a degradosome complex, where it functions as both an exonuclease and a poly (A) polymerase to control RNA stability (32) . In plants, PNPase localizes to the chloroplast stroma, where it performs polyadenylation and exonucleolytic degradation to mediate RNA turnover (3, 28, 29, 53). In both bacteria and plants, poly(A) addition to RNA constitutes a signal for PNPase-mediated RNA degradation.

Interestingly, PNPase is absent from archaea and single-cell eukaryotes including *Saccharomyces cerevisiae*, *Schizosaccharomyces pombe*, and *Plasmodium falciparum* but is present in animals including the worm, fly, mouse, and human (24). This odd phylogenetic distribution suggests that PNPase might possess a unique function in animals, rather than its evolutionarily conserved role in the degradosome, as shown for prokaryotes and chloroplasts.

Recently, human PNPase was identified as an up-regulated gene in terminally differentiated melanoma cells and in senescent progeroid fibroblasts, implying a potential role in growth inhibition (25). Human PNPase is a 85-kDa nucleus-encoded protein that localizes to the mitochondrion (4, 38) and functions in vitro as a phosphate-dependent exoribonuclease (8, 26) and poly(A) polymerase (33). Analogous to bacterial and chloroplastic PNPase, mammalian PNPase could be expected to regulate RNA levels in the mitochondrial matrix. However, robust PNPase overexpression inhibits cell growth and induces apoptosis with down-regulation of *MYC* and *BCL-xL* expression (42), suggesting that PNPase may degrade mRNAs in the cytosol. These observations deviate from a predicted role for PNPase in regulating mitochondrial RNA levels (33). A possible explanation for this disparity includes aberrant targeting of exogenous PNPase to the cytosol. Alternatively, PNPase might function within and outside mitochondria, akin to cytochrome *c*, apoptosis-inducing factor, Smac/DIABLO, HtrA2/ Omi, Bit1, and endonuclease G, with the intermembrane space providing a location for mitochondrial activity and conditions that cause outer membrane permeability enabling cytosolic access (41). Indeed, we have identified PNPase by its interaction with the cytosolic TCL1 oncoprotein in recent work (8). Furthermore, our accompanying paper shows that PNPase does not regulate mitochondrial RNA levels but, after treatment with apoptotic stimuli, is mobilized late after cytochrome *c* release (4).

Given that the specific localization of PNPase is not understood, we have characterized the PNPase biogenesis pathway in detail. Our accompanying study shows that endogenous PNPase resides in the intermembrane space in mouse liver mitochondria and, when heterologously expressed in *S. cerevisiae*, localizes in mitochondria and not the cytosol (4). Moreover, PNPase has a typical N-terminal targeting sequence that confers mitochondrial localization.

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Mitochondria possess an elaborate set of translocons on the outer membrane (TOM) and translocons on the inner membrane (TIM) to mediate the import and assembly of nucleuscoded proteins such as PNPase (20, 21, 36, 46). Whereas the TIM22 complex mediates the import of inner membrane proteins, including the carrier family and the import component Tim22, the TIM23 translocon imports proteins with a typical N-terminal targeting sequence (21). After passing through the TOM, the precursor engages the TIM23 complex; translocation initiation into the TIM23 complex requires the electrochemical membrane potential $(\Delta \psi)$ of the inner membrane. For proteins targeted to the matrix, the Hsp70 motor mediates translocation. The matrix processing protease (MPP), consisting of the heterodimer Mas1/Mas2, typically mediates cleavage of the presequence (9). Precursors destined for the intermembrane space, such as cytochromes b_2 and c_1 , contain a bipartite targeting sequence. The N terminus directs the precursor to TIM23. After cleavage by the MPP, the precursor arrests at the "stop-transfer" domain in the translocon (12). The inner membrane protease (IMP), consisting of the complex Imp1/Imp2 and facing the intermembrane space, mediates a second cleavage, thereby trapping cytochrome b_2 in the intermembrane space (9) .

Additional proteases in the mitochondrial inner membrane have been implicated in import pathways in recent studies (2, 47). Beyond their role in protein degradation, the *m-*AAA (ATPases associated with a number of diverse cellular activities) proteases Yta10 and Yta12 are required for maturation of the ribosomal protein MrpL32 and cytochrome *c* peroxidase (Ccp1) (7, 34). The evolutionarily conserved rhomboid protease processes the substrates Mgm1, involved in mitochondrial dynamics, and Ccp1 (7, 16). These recent studies confirm that the mitochondrial proteolytic system is crucial for the maturation and assembly of precursors in the mitochondrion.

Import mechanisms into the intermembrane space are diverse (17). Translocation of proteins such as cytochromes b_2 and c_1 requires the hydrolysis of matrix ATP and $\Delta \psi$. For proteins that lack a typical N-terminal targeting sequence and do not require a $\Delta\psi$ for translocation (thereby bypassing the TIM), import mechanisms rely on (i) the energy gain of protein folding, often around a cofactor, or (ii) the association of a protein with a high-affinity binding site, such as another protein.

Here we use the conserved import machinery of *S. cerevisiae* to investigate the pathway by which PNPase is localized to the intermembrane space. Import depends on the presence of $\Delta \psi$, and the presequence is cleaved by MPP. Translocation of PNPase into the intermembrane space requires the *i*-AAA protease Yme1, which possesses chaperone-like activity (23); this constitutes the first evidence of a translocation function for this protease, presenting a new twist on biogenesis mechanisms in the intermembrane space. Conceivably, other intermembrane space proteins, particularly those involved in the execution events of apoptosis, may also utilize Yme1 for import.

MATERIALS AND METHODS

Plasmids and strains. Standard genetics techniques were used for growth, manipulation, and transformation of yeast strains (14). Human PNPase cDNA was cloned by PCR from a Nalm-6 B-cell cDNA library with 5'-ccggtattatccAT GGCGGCCTGCAGGTAC-3' and 5'-ccgcacgtcgacTCACTGAGAATTAGAT G-3' primers (lowercase indicates inserted restriction site). PNPase was subcloned into pSP65 (Promega) and pcDNA3.1myc-His (Invitrogen) for in vitro transcription/translation. For expression in yeast, PNPase was subcloned into pYEX-BX (2μm, *URA3*; Clontech), and induction was regulated by the copperinducible *CUP1* promoter. The pYEX-BX-PNPase construct was transformed into GA74-1A or the corresponding mutant yeast strains (22). The Δ *yme1* strains were purchased from ResGen. The yeast strains MY111-2 (as the *mas1* strain is designated) and JN174 (as the $\Delta imp1 \Delta imp2$ strain is designated) have been previously described (35, 51). The *yme1*^{E541Q} strain was generously provided by Thomas Langer (23). The yeast strain deleted for \lim_{54} (the $\Delta \lim_{54}$ strain) has been previously described (D. Hwang et al., submitted for publication). The *E. coli* strain for expressing His-tagged MPP was a gift from Vincent Geli (30). His-tagged Omi/HtrA2-(134-458) was generously provided by Antonis Zervos (5).

The PNPase-dihydrofolate reductase (DHFR) fusion construct was generated by subcloning the first 157 amino acids of PNPase in frame with DHFR-His $_6$ derived from pQE16 (QIAGEN) into pET28a (Novagen). Recombinant PNPase-DHFR was induced according to the manufacturer's protocols, and the recombinant protein was purified under denaturing conditions according to the manufacturer's protocols. For in vitro transcription/translation, PNPase-DHFR and PNPase(1-392) were amplified by PCR with 5'-ggggatttaggtgacactatagaaggagcta agcttATGGCGGCCTGCAGGTA-3' (encoding the SP6 promoter at the 5' end) and 5'-AAGCTTAGTGATGGTGATGG-3' (PNPase-DHFR) or 5'-CTGTGT TTGTCCTCTTTGA-3' [PNPase(1-392)] and isolated using a Nucleospin extraction kit (Clontech).

Purification of MPP and Omi/HtrA2 from *E. coli* **and in vitro processing assay.** MPP and Omi/HtrA2 were purified as described previously (5, 30). Radiolabeled precursors (5 μ l) or purified PNPase(1-157)-DHFR (2.5 μ g) was incubated in a final volume of 20 μ l or 165 μ l cleavage buffer (50 mM NaCl, 20 mM HEPES-KOH [pH 7.4], 1 mM ZnCl₂), respectively, in the presence or absence of MPP for 30 min at 30°C. The Omi protease cleavage reactions were performed as described above, except in 200 mM NaCl, 20 mM NaPO₄ (pH 7.4), 5% glycerol at 37°C. Twenty micrograms of casein was used as a control substrate for Omi cleavage, and the inhibitor UCF-101 (Calbiochem) was used at a final concentration of 20 μ M. Samples were analyzed by sodium dodecyl sulfatepolyacrylamide gel electrophoresis (SDS-PAGE) and autoradiography (radiolabeled precursors), immunoblotting with anti-DHFR (recombinant PNPase-DHFR), or Coomassie staining (casein). For Edman degradation, the cleaved PNPase DHFR was transferred to a polyvinylidene fluoride (PVDF) membrane for analysis. N-terminal sequencing was performed on an Applied Biosystems model 494 precise protein sequencer by the DNA core facility at the USC Norris Comprehensive Cancer Center.

Import of radiolabeled proteins into isolated mitochondria, manipulation of mitochondria, and cross-linking studies. Mitochondria were purified from yeast cells grown in rich ethanol/-glycerol medium (13) and assayed for in vitro protein import as described previously (39). Proteins were synthesized in a rabbit reticulocyte lysate in the presence of $[35S]$ methionine after in vitro transcription of their corresponding genes by SP6 or T7 polymerase. The reticulocyte lysate containing the radiolabeled precursor was incubated with isolated mitochondria at 25°C in import buffer (1 mg/ml bovine serum albumin, 0.6 M sorbitol, 150 mM KCl, 10 mM MgCl₂, 2.5 mM EDTA, 2 mM ATP, 2 mM NADH, 20 mM HEPESKOH [pH 7.4], 2 mM KH₂PO₄, 5 mM L-methionine). Where indicated, the potential across the mitochondrial inner membrane was dissipated with $1 \mu M$ valinomycin and 25 μ M carbonyl cyanide *p*-(trifluoromethoxy)phenylhydrazone (FCCP). Nonimported radiolabeled proteins were removed by treatment with 100 μ g/ml trypsin for 30 min on ice, and trypsin was inhibited with 200 μ g/ml soybean trypsin inhibitor. Osmotic shock experiments and cross-linking studies were performed as previously described (22, 27). After import, mitochondria were washed, suspended at 1 mg/ml in import buffer without bovine serum albumin and methionine, and incubated with 1 mM dithiobis(succinimidyl propionate) on ice for 30 min followed by a quench with 100 mM Tris-HCl, pH 8.0. For immunoprecipitation, solubilized mitochondria were incubated with the corresponding monospecific rabbit immunoglobulin Gs coupled to protein A-Sepharose (39).

Miscellaneous. Mitochondrial proteins were analyzed by SDS-PAGE and a Tricine-based running buffer (43). Proteins were detected by immunoblotting using nitrocellulose and PVDF membranes and by visualization of immune complexes by use of horseradish peroxidase-conjugated secondary antibodies and enhanced chemiluminescence (Pierce). PNPase (1:1,000), AAC (1:5,000), cytochrome b_2 (1:2,000), DHFR (1:2,000), and Mge1 (1:1,000) antisera were used in immunoblot analyses. For autoradiography, 6 or 8% gels were used for PNPase and 10 or 12% gels were used for PNPase-DHFR. Degradation assays

FIG. 1. PNPase processing is inhibited in yeast lacking the MPP and *i*-AAA protease Yme1. (A) Schematic showing the N-terminal targeting sequence within the N terminus of PNPase. The basic residues are indicated with a " $+$," the sequence of the mature PNPase is underlined, and the arrows mark the MPP cleavage site. (B) The first 26 amino acids were placed on a helical wheel (40). The boxes mark basic residues, the diamonds mark hydrophobic residues, and the circles mark the remaining residues. (C) PNPase was expressed hetero-
logously in yeast under control of the Cu²⁺-inducible *CUP1* promoter. Expression was analyzed in a total lysate by immunoblotting with polyclonal anti-PNPase antibody. PNPase was expressed (+PNPase) in strains defective in MPP (mas1), IMP (Δ imp1 Δ imp2), and Yme1 $(\Delta y$ me1) and in the WT strain. The precursor form (p) is marked for the in vitro-translated standard (Std), and the mature form (m) is marked for PNPase expressed in WT yeast (WT +PNPase). The asterisk marks products that accumulate when PNPase import and assembly are impaired.

for PNPase-DHFR and the model substrate Yta10-DHFR were performed as described previously (23).

RESULTS

PNPase requires the mitochondrial proteases for maturation. The N-terminal 37 amino acids are predicted to target PNPase to the mitochondrion (Fig. 1A), and this targeting sequence is absent from prokaryotic PNPase proteins (24). PNPase features a typical N-terminal targeting sequence in which the basic residues cluster on one face of the predicted α -helix and the hydrophobic residues cluster to other regions (Fig. 1B) (48). As we show in the accompanying paper, PNPase resides in the intermembrane space (4); however, to specifically localize PNPase to this compartment, we investigated the PNPase biogenesis pathway in detail. Because PNPase is processed at the N terminus (4), we first determined which mitochondrial proteases are required. We took advantage of yeast for in vivo studies and yeast mitochondria for in organello studies, as protein import is highly conserved between animals and fungi (18, 21), and yeast are amenable to genetic and biochemical manipulations. Specifically, heterologous PNPase expressed in yeast and endogenous PNPase from mouse liver fractionated identically in isolated mitochondria, displayed similar sensitivities to protease treatment during osmotic shock, and assembled in similarly sized complexes (4).

We expressed PNPase under control of the Cu^{2+} -inducible *CUP1* promoter (31) and analyzed expression in strains that contained mutations in the known mitochondrial proteases (Fig. 1C) (7, 51). The MPP cleaves the targeting sequence of TIM23 substrates as the N-terminal targeting sequence enters

the matrix (9). As MPP is essential for viability, we utilized the temperature-sensitive *mas1* mutant to assess PNPase maturation (51). We also anticipated that PNPase would be cleaved by a protease in the intermembrane space due to its intermembrane space localization; likely candidates included the IMP (Δ *imp1* Δ *imp2* strain) and the *i*-AAA protease Yme1 (Δ *yme1* strain) (9). Total protein lysates from cells expressing PNPase were separated by SDS-PAGE and immunoblotted with PNPase antiserum. The mature form of PNPase was observed in wildtype (WT) yeast cells, as previously shown (4), and in cells lacking the IMP, which migrates faster in the gel than the control in vitro translation of the PNPase precursor (Fig. 1C). However, mature PNPase was not observed in cells defective in the MPP and the Yme1 protease. Aberrantly sized bands were detected when PNPase import was defective; these most likely correspond to degradation products that accumulate because PNPase import and processing are impaired.

MPP mediates cleavage of PNPase. Using the in organello import assay (12, 51), we imported radiolabeled PNPase into WT mitochondria (Fig. 2A) (4). As a control, we included the matrix-targeted fusion protein Su9-DHFR, which consists of subunit 9 of the *Neurospora crassa* ATPase fused to the reporter dihydrofolate reductase (37). The import assay was performed for 10 min at 25 $^{\circ}$ C in the presence and absence of $\Delta \psi$; import was confirmed by removing nonimported precursor with trypsin treatment. As expected, PNPase translocated into isolated WT mitochondria, and the N-terminal sequence was processed (Fig. 2A). We used the same assay to investigate import of radiolabeled PNPase and Su9-DHFR into *mas1* mitochondria, which are defective in MPP (Fig. 2B) (12, 51). In a control reaction, we included the TIM22 substrate, the ADP/ ATP carrier (AAC), which is targeted to the inner membrane via the TIM22 translocon (22). Mitochondria from the temperature-sensitive *mas1* strain were prepared after being shifted to the restrictive temperature for 6 h as reported previously (51). The mature form of Su9-DHFR did not accumulate in the presence of a $\Delta\psi$ in the *mas1* mitochondria (12); further, AAC was imported into the inner membrane, indicating that the TIM22 import pathway functions in the *mas1* mitochondria. PNPase import and processing failed in *mas1* mitochondria irrespective of $\Delta\psi$, confirming the in vivo results from Fig. 1C.

To validate that MPP is required for the putative first cleavage step and that the N terminus of PNPase indeed functions as a targeting sequence, we fused PNPase amino acids 1 to 157 to DHFR, generating the chimeric construct PNPase-DHFR. This radiolabeled fusion protein was incubated with WT mitochondria, and aliquots were removed during a time course and treated with protease to remove nonimported precursor (Fig. 2C). As shown for native PNPase, the fusion protein was imported into isolated mitochondria. To confirm that the first 157 amino acids were sufficient to localize PNPase to the intermembrane space, we coupled the import assay with hypotonic lysis treatment (Fig. 2D, left panel); the mitochondria were diluted into buffer with a final sorbitol concentration of 0.06 M, which selectively ruptures the outer membrane but leaves the inner membrane intact, to generate mitoplasts (12). After osmotic shock, the PNPase-DHFR was not soluble but remained associated with the membranes, as has been shown for endogenous PNPase (4). However, trypsin treatment de-

FIG. 2. MPP mediates cleavage of PNPase. (A) Radiolabeled PNPase was synthesized in vitro and incubated with isolated WT mitochondria in the presence or absence of a $\Delta\psi$ at 25°C for 10 min. After import, samples were divided in equal aliquots for protease treatment with trypsin (Tryp) to remove nonimported precursor; protease activity was halted with soybean trypsin inhibitor. As a control, matrix-localized Su9-DHFR was also imported. Samples were analyzed by SDS-PAGE and fluorography. Standard (Std) refers to 10% of the radioactive precursor added to each assay. p, precursor; m, mature. (B) Import of PNPase and Su9-DHFR was performed as described for panel A into *mas1* mitochondria. The inner membrane marker ADP/ATP carrier (AAC) was imported as a control; import reactions with AAC, which lacks a cleavable targeting sequence, were treated with protease followed by carbonate extraction to confirm insertion into the inner membrane. (C) Radiolabeled PNPase-DHFR (the N-terminal 157 amino acids of PNPase fused to DHFR) was imported into isolated WT mitochondria, and aliquots were removed at the designated time points. Nonimported precursor was removed with protease treatment. Standard (Std) represents 10% of the radioactive precursor in each time point. (D) PNPase-DHFR was imported into WT mitochondria (M) at 25° C for 10 min in the presence and absence of $\Delta \psi$. Samples were incubated in hypotonic buffer to swell the outer membrane, generating mitoplasts (MP), in the presence and absence of trypsin (Tryp), followed by inactivation with trypsin inhibitor. Mitoplasts were recovered by centrifugation (P) and separated from the supernatant (S) containing the soluble intermembrane space contents. As a control, Su9-DHFR was imported and treated identically; relevant reactions representing import in the presence of $\Delta\psi$ and the recovered mitoplast fraction (P) treated with trypsin (+Tryp) are shown.

graded the imported PNPase-DHFR, indicating that the PNPase-DHFR was localized to the intermembrane space. As a matrix-targeted control, imported Su9-DHFR remained protected from protease (Fig. 2D, right panel; note that only relative lanes for the control have been shown). Therefore, the first 157 amino acids contain the necessary targeting information to direct PNPase to the intermembrane space.

To test the requirement for MPP, mitochondria were incubated with MPP inhibitors, EDTA and 1,10-*o*-phenanthroline, prior to import of PNPase, Su9-DHFR, and AAC (Fig. 3A) (52). Mitochondria treated with the chelators lacked processed PNPase and Su9-DHFR; AAC was imported into the inner membrane, confirming that the chelator treatment did not inhibit the TIM22 import pathway and that the mitochondria were still functional (6). We then purified recombinant MPP containing a hexahistidine tag for an in vitro cleavage assay (Fig. 3B) (11). Recombinant MPP was eluted from the $Ni²⁺$ agarose and incubated with in vitro-translated substrates. Purified MPP cleaved radiolabeled PNPase and Su9-DHFR (Fig. 3C). In addition, MPP mediated the first cleavage of cytochrome c_1 to the intermediate form. The purified MPP was used in a dilution assay to compare the cleavage efficiencies of PNPase, PNPase-DHFR, and Su9-DHFR (Fig. 3D). The Su9 targeting sequence is a unique substrate for the MPP because it contains two cleavage sites (44), both of which are cleaved very efficiently. MPP at a 1:10 dilution cleaved approximately 50% of the PNPase constructs and Su9-DHFR. Therefore, the MPP cleavage efficiencies for PNPase and PNPase-DHFR

were similar to that for the control substrate, and MPP cleaved PNPase at one location.

Endogenous and heterologously expressed PNPase sequences have identical N termini. To compare the biogenesis of PNPase in yeast with that in mammals, we identified the N terminus of PNPase in mammalian cells by use of mass spectrometry (Fig. 1A) (4). PNPase was purified from mammalian cells by coimmunoprecipitation with TCL1 (8) and analyzed by mass spectrometry. The tandem mass spectrometry spectra showed three peptide fragments with a nontryptic amino terminus for 38-LWSSAGSR-45 that marked the 5' boundary of recovered PNPase sequences; peptide spectra were not detected amino terminal to this position. Tandem mass spectrometry coverage was excellent, at $\sim 85\%$ of the full-length protein, including the missing N-terminal 37 residues.

We purified recombinant PNPase-DHFR for N-terminal sequencing using Edman degradation. PNPase-DHFR was cleaved after the addition of yeast MPP (Fig. 3E), and the cleaved DHFR fragment was recovered after transfer to a PVDF membrane. From the Edman sequencing assay, leucine-38 was also identified as the N-terminal amino acid, as indicated in Fig. 1A. These results indicate that the N terminus of PNPase is identical whether cleaved with recombinant yeast MPP or isolated from mammalian cells. Additionally, the cleavage site resides at a consensus $R-2$ site for MPP, in which an arginine residue is located at the -2 position from the cleavage point (Fig. 1A) (49). Thus, PNPase processing in yeast mitochondria is identical to that in mammalian mito-

FIG. 3. Recombinant MPP mediates cleavage of PNPase. (A) WT mitochondria were left untreated or were incubated in 10 mM EDTA, 2 mM *o*-phenanthroline (o-phe) for 15 min prior to the start of the import reaction to inhibit the endogenous MPP (52). Radiolabeled PNPase, Su9-DHFR, and AAC were imported as described for Fig. 2A. Nonimported AAC was removed by protease treatment, but protease addition was omitted for Su9-DHFR and PNPase. Standard (S) refers to 10% of the radioactive precursor added to each assay. p, precursor; m, mature. (B) Recombinant MPP (10) was purified over a Ni^{2+} column from a total lysate (T), followed by washing to remove unbound proteins. Purified MPP was eluted in fractions 2 and 3 in the presence of 300 mM imidazole. (C) In an in vitro cleavage assay (10) , MPP was incubated with radiolabeled PNPase, Su9-DHFR, and cytochrome c_1 . The standard (Std) is an equal volume of the radioactive precursor. i, intermediate. (D) Serial MPP dilutions were incubated with PNPase, PNPase-DHFR, and Su9-DHFR for 30 min at 30°C. The reactions were stopped with SDS sample buffer. Samples were separated by SDS-PAGE and detected by fluorography. (E) Recombinant PNPase-DHFR was purified using a C-terminal hexahistidine tag for an in vitro cleavage assay, and cleavage was confirmed by immunoblot analysis with a polyclonal antibody against DHFR (α DHFR). The cleaved DHFR was recovered for analysis by Edman degradation.

FIG. 4. Yme1 is required for PNPase assembly in the intermembrane space. Radiolabeled PNPase, Su9-DHFR, and cytochrome *c*¹ were imported into Δyme1 (A), Δimp1 Δimp2 (B), yme1^{E514Q} (Su9-DHFR control omitted) (C), or Δt *im54* (D) mitochondria in the presence and absence of $\Delta\psi$ as described for Fig. 2A. In half of the samples, nonimported precursor was removed by trypsin treatment, and analysis was done subsequently with SDS-PAGE and fluorography. p, precursor; i, intermediate; m, mature.

chondria, confirming yeast as a valid model for mitochondrial import studies of mammalian proteins. Further, MPP is the sole protease required for PNPase processing.

Yme1 is required for PNPase import. Given the necessity of Yme1 for PNPase maturation in yeast mitochondria (Fig. 1C), we characterized the translocation pathway of PNPase into the intermembrane space. PNPase and controls Su9-DHFR and cytochrome c_1 were imported into mitochondria lacking Yme1 (Δyme1 mitochondria) or IMP (Δimp1 Δimp2 mitochondria) (Fig. 4A and B). PNPase and Su9-DHFR imported efficiently into Δ *imp1* Δ *imp2* mitochondria; however, mature cytochrome $c₁$ was not generated, as IMP is required for the second cleavage that occurs during cytochrome c_1 import (12). Note that the intermediate form of cytochrome c_1 is protected from added protease, indicating that the C terminus has completely crossed the outer membrane (50). In contrast to Su9-DHFR and cytochrome c_1 , which imported correctly, PNPase did not import into Δ *yme1* mitochondria. In addition, marker proteins were localized identically in WT and Δ *yme1* mitochondria (see Fig. S1 in the supplemental material), indicating that Δy *me1* mitochondria were not generally defective in assembly. We analyzed whether PNPase import was dependent upon functional Yme1 by testing import into mutant *yme1*E541Q mitochondria, expressing a proteolytically inactive variant of Yme1 with a point mutation in the proteolytic site (23). As in the -*yme1* mitochondria, PNPase import was selectively inhibited (Fig. 4C).

FIG. 5. Yme1 is required for efficient translocation of PNPase and PNPase-DHFR. (A) PNPase-DHFR was imported into WT, *mas1*, Δy me1, and Δ *imp1* Δ *imp2* mitochondria (mitos) as described for Fig. 2A. (B) As shown in Fig. 2D in the same set of experiments, imported PNPase-DHFR was localized to the intermembrane space after osmotic shock. (C) Time course assays were performed for the import of PNPase, PNPase-DHFR, and Su9-DHFR (control) into WT, Δy *me1*, and Δt *im54* mitochondria at the indicated times. Nonimported precursor was removed by trypsin treatment, and equal aliquots were separated by SDS-PAGE. The percent substrate imported was quantitated from fluorographs by use of a scanning densitometer. The amount of precursor that imported into WT mitochondria at the end time point was set at 100%.

Previously, we have shown that Tim54 is required for assembly of Yme1 into a proteolytically active complex (Hwang et al., submitted). To confirm that Yme1 is essential for PNPase biogenesis, we investigated import of PNPase and controls Su9-DHFR and cytochrome c_1 into mitochondria deleted for *TIM54* (Δ *tim54* mitochondria) (Fig. 4D). Whereas Su9-DHFR and cytochrome c_1 were imported, PNPase import was impaired in Δ *tim54* mitochondria. This result verifies that the fully assembled and functional Yme1 complex is required for PNPase accumulation in the intermembrane space.

We next probed the events at the outer membrane in more detail. PNPase-DHFR import into WT, *mas1*, Δ *yme1*, and Δ *imp1* Δ *imp2* mitochondria and subsequent protease treatment to remove nonimported precursor were performed (Fig. 5A). PNPase-DHFR import was inhibited in *mas1* mitochondria but occurred normally in WT and Δimp1 Δimp2 mitochondria. Interestingly, PNPase-DHFR was imported into Δy *me1* mitochondria, as PNPase-DHFR remained protected from added protease. As expected from the osmotic shock experiments illustrated in Fig. 2D, PNPase-DHFR localized to the intermembrane space in Δy *me1* mitochondria because PNPase-DHFR was sensitive to exogenous protease introduced during osmotic shock (Fig. 5B).

We performed time course assays and quantitated import for PNPase, PNPase-DHFR, and Su9-DHFR into WT, Δ yme1,

and Δ *tim54* mitochondria (Fig. 5C). Whereas the import of Su9-DHFR was efficient, PNPase import was essentially undetectable in mitochondria lacking functional Yme1. PNPase-DHFR import also required functional Yme1 for efficient import, because import was decreased by approximately 75 to 80% in Δ *tim54* and Δ *yme1* mitochondria. Given that PNPase-DHFR (\sim 40 kDa) is shorter than PNPase, we also tested the import of truncated PNPase $(1-392)$ $(\sim 43 \text{ kDa})$ into Δ *yme1* mitochondria in time course assays (see Fig. S2 in the supplemental material); as with PNPase-DHFR, the import of truncated PNPase(1-392) was similarly decreased, suggesting that Yme1 facilitates translocation across the outer membrane. Taken together, these data show that Yme1 is required for the maturation of PNPase in the intermembrane space.

The PNPase translocation intermediate engages the TOM and TIM23 and binds to Yme1. Based on our import analysis, we predicted that, during translocation, PNPase should engage the TOM and TIM23 and bind directly to Yme1. PNPase and PNPase-DHFR were imported into WT mitochondria followed by cross-linking with the thiol-cleavable cross-linker dithiobis(succinimidyl propionate) (Fig. 6A and B). After a quenching with 0.1 M Tris, an aliquot of the PNPase-DHFR was separated by SDS-PAGE in the absence of reductant, whereas the remainder was immunoprecipitated with antibodies against Tim23, Yme1, Tom40, and the negative control,

Ayme1 mitos

FIG. 6. Yme1 binds directly to PNPase and mediates translocation into the intermembrane space. (A) PNPase-DHFR was imported into WT mitochondria for 10 min at $25^{\circ}C$ (-XL) and an aliquot was removed. Proteins were cross-linked $(+XL)$ by the addition of 1.0 mM dithiobis(succinimidyl propionate) for 30 min followed by quenching with 0.1 M Tris-HCl. After an aliquot was removed, mitochondria were subsequently solubilized followed by immunoprecipitation (IP) with antibodies (α) against Yme1, Tim23, Tom40, and the control cytochrome b_2 (cyt b_2) Because cross-linked PNPase-DHFR migrates at a high molecular mass (indicated by the diamond), the immunoprecipitated cross-linking reactions were released with β -mercaptoethanol addition in the sample buffer, whereas reductant was omitted from the cross-linking reactions. The asterisks indicate cross-linked PNPase-DHFR (released by treatment with reductant) that copurified with the target protein of the antibody. (B) Cross-linking and immunoprecipitation assays were performed with full-length PNPase as described for panel A. (C) Cross-linking and immunoprecipitation assays were performed with PNPase-DHFR import into Δ *yme1* mitochondria. Note that PNPase-DHFR was not immunoprecipitated with antibodies against Yme1, confirming the specificity of the Yme1-PNPase interaction.

cytochrome b_2 . Cross-linked PNPase migrated near the top of the gel; this large cross-linked product was expected because Yme1 assembles into a large homo-oligomeric complex (23) and our cross-linking conditions indicated that several Yme1 monomers were simultaneously cross-linked (data not shown). After the immunoprecipitation reaction, we cleaved the crosslinked products with the addition of B-mercaptoethanol to release cross-linked PNPase-DHFR to its native molecular mass and separated the products by SDS-PAGE. As predicted, PNPase and PNPase-DHFR bound directly to Yme1, Tom40, and Tim23 but not to cytochrome b_2 .

As further confirmation, we investigated the association of PNPase-DHFR into *yme1*^{E541Q} and Δ *yme1* mitochondria using the same cross-linking protocol (Fig. 6C) (see Fig. S3 in the supplemental material). PNPase-DHFR bound directly to Yme1, Tom40, and Tim23 in *yme1*^{E541Q} mitochondria. As expected, a PNPase-DHFR–Yme1 interaction was not detected in mitochondria lacking Yme1 (Fig. 6C), although interactions with TIM23 and TOM were detected, indicating that the precursor can engage the translocons. Consequently, a new function has been determined for Yme1: translocation of precursors into the intermembrane space.

PNPase-DHFR is not a proteolytic substrate for Yme1. To determine if Yme1 might bind to PNPase-DHFR because Yme1 functions as a proteolytic component rather than as an import mediator, we compared PNPase-DHFR degradation with that of the Yme1p model substrate for degradation assays, Yta10-DHFR^m (23), which consists of a fusion between the N-terminal 161 amino acids of Yta10p and a loosely folded mutant of DHFR. As described previously, PNPase-DHFR and Yta10-DHFR^m constructs were imported into WT mitochondria for 30 min in the presence of an ATP-regenerating system at 37°C (23), and stability was then measured over a 60-min period (Fig. 7A). Whereas approximately 50% of the Yta10-DHFR^m was degraded in WT mitochondria, PNPase-DHFR was stable in WT mitochondria, similar to the stability of Yta10-DHFRm in mitochondria deleted for *yme1*. If PNPase was indeed a proteolytic substrate for Yme1, mature PNPase would be expected to accumulate in mitochondria when expressed in the Δ *yme1* yeast strain. We examined the targeting and stability of PNPase in WT and Δ *yme1* mitochondria in vivo (Fig. 7B). With WT mitochondria, we found that both PNPase and the marker AAC localized to mitochondria. However, with Δ *yme1* mitochondria, we found that PNPase did not accumulate in the mitochondrion, supporting a role for Yme1 in the import of PNPase. We assign a new function to Yme1 in the translocation of PNPase (but interestingly not in proteolytic processing or degradation) and suggest that other proteins, particularly sequestered mammalian apoptotic proteins residing in the intermembrane space, may also rely on Yme1 for biogenesis.

DISCUSSION

PNPase is localized to the intermembrane space. In contrast to studies that suggest a localization and function for PNPase in the cytosol or mitochondrial matrix (26, 33, 38), our accompanying study finds that PNPase is a new resident in the intermembrane space (4). Previous studies have suggested that PNPase functions in the cytosol to degrade specific cytosolic

FIG. 7. PNPase is not a proteolytic substrate of Yme1. (A) Radiolabeled Yta10-DHFR^m and PNPase-DHFR were imported into WT mitochondria for 30 min, and nonimported precursor was removed by protease treatment. The mitochondria were then incubated in the presence of an ATP-regenerating system at 37°C, and aliquots were removed at the indicated times to measure stability. The rate of proteolysis during the time course from three independent assays was quantitated using a Bio-Rad FX molecular imager and the affiliated Quantity 1 software. The amounts of Yta10-DHFR^m and PNPase-DHFR remaining at each chase time point are expressed as the percentages of the amounts detected for the respective strains at $t = 0$ (set at 100%; mean \pm standard deviation; $n = 3$). As a control for Yme1dependent degradation of Yta10-DHFRm, a control reaction experiment was performed with Δ *yme1* mitochondria. (B) Mitochondria (Mito) and the postmitochondrial supernatant (PMS) were fractionated from Δ yme1 and WT mitochondria expressing PNPase. Samples were separated by SDS-PAGE and blotted for PNPase and the mitochondrial marker AAC.

RNAs. However, these results may be artifact, because PNPase was expressed with an N-terminal tag that blocked translocation into the mitochondria and/or PNPase was overexpressed (25, 26). Indeed, in our companion work, we specifically show that an N-terminal tag appended to the N terminus blocked import and processing of PNPase (4). In another study, PNPase was not implicated in the regulation of RNA dynamics in the mitochondrial matrix (33). From our detailed analysis of PNPase biogenesis, we place PNPase precisely in the intermembrane space and define its specific function with respect to mitochondrial localization.

A new function for Yme1 in protein translocation. We have used a systematic approach, taking advantage of yeast and isolated mitochondrion mutants to directly investigate the biogenesis pathway of PNPase. Our studies reveal an unexpected twist on import mechanisms into the intermembrane space (Fig. 8). PNPase is imported via the TOM and TIM23. MPP mediates cleavage (step 1) and Yme1 serves as a nonconventional translocation motor to pull PNPase into the intermembrane space (step 2). PNPase then assembles in mature complexes similarly sized in both yeast and mammalian mitochondria (step 3) (4); this complex has been postulated to

FIG. 8. Yme1 mediates PNPase import into the intermembrane space. The schematic shows the import pathway of PNPase. See Discussion for details. OM, outer membrane; IM, inner membrane; IMS, intermembrane space.

be a trimer or dimer of trimers, based on the structure of the bacterial PNPase (45).

PNPase contains a typical N-terminal targeting sequence, and MPP processes PNPase at a predicted cleavage site. In addition, the region following the N-terminal targeting sequence (amino acids 45-AVAVDLG-51) is hydrophobic and may serve as a stop-transfer domain to arrest translocation in TIM23, similar to the stop-transfer domains in cytochromes b_2 and c_1 (12). The substrate NADH-cytochrome b_5 reductase shows similar properties (15), requiring MPP cleavage for localization in the intermembrane space. However, NADH-cytochrome $b₅$ reductase biogenesis is more complicated, since it resides in both the outer membrane and the intermembrane space and is dependent upon folding and interactions with the TOM and TIM23.

Given that cross-links between PNPase and Tom40, Tim23, and Yme1 can be trapped, Yme1 is seemingly in close proximity to TIM23. Surprisingly, a processed PNPase intermediate is not detected in mitochondria lacking Yme1; therefore, Yme1 may play an indirect role in mediating MPP cleavage, potentially by stabilizing PNPase in a conformation that is amenable for MPP cleavage in TIM23. Yme1 pulls the C terminus of PNPase across the TOM complex, as evidenced by defective full-length PNPase import in *yme1* mutant mitochondria (Fig. 1C and 4A). The normal function assigned to Yme1 is that of a protease; however, PNPase-DHFR was not efficiently degraded by Yme1 in comparison to the model substrate Yta10-DHFR^m.

Although other AAA protease family members have been implicated in translocation, this study uniquely assigns a translocation function to Yme1. Previously, Yme1 was characterized only for its chaperone and protease activities (2). However, Yme1's chaperone function in translocation is not unexpected, because these functions have been observed for the AAA proteases. Leonhard and colleagues have reported that Yme1 specifically senses the folding state of solvent-exposed domains and degrades nonnative membrane proteins (23), extracting them from the membrane. The recombinant AAA domain of Yme1 binds to unfolded peptides and suppresses their aggregation. Additionally, the *m*-AAA proteases Yta10 and Yta12, which face the matrix, mediate the assembly of cytochrome oxidase (1). Finally, the AAA protease Cdc48 is proposed to function in retrograde translocation, pulling a misfolded protein from the endoplasmic reticulum (19). Our studies demonstrate that Yme1 binds directly to unfolded PNPase during translocation, a feature reminiscent of chaperones, and then diverts it into the intermembrane space rather than targeting it for degradation. Plausibly, partner proteins may participate in this pathway, as has been reported for other AAA proteases. Future studies to elucidate the mechanism by which Yme1 targets PNPase for import rather than degradation will provide interesting insights into the function of this protease.

Since mammalian mitochondria contain the Omi protease, which is absent in yeast (41), we ruled out recombinant Omi protease as an import mediator of PNPase (5) (see Fig. S4 in the supplemental material). We have attempted to investigate the role of Yme1 in PNPase import in cultured mammalian cells by down-regulating Yme1 levels using RNA interference (4). Unfortunately, the cells died quickly, and unprocessed full-length PNPase was undetectable in isolated cytosol and mitochondrial fractions. Because of difficulties manipulating cultured mammalian cells, yeast mitochondria have proven advantageous, given the conservation of protein import pathways among plants, animals, and fungi.

What is the function of PNPase in the intermembrane space? The detailed localization presented herein creates a framework for assigning a function to PNPase. Our accompanying paper shows that down-regulation of PNPase via RNA interference impacts mitochondrial morphology and respiration and that PNPase is mobilized late with apoptotic stimuli and may subsequently bind to TCL1 (4). That PNPase is found only in animals and not in fungi suggests it may have acquired a unique function distinct from its conserved role in RNA dynamics. Additional intermembrane space proteins, particularly those implicated in apoptosis, may require Yme1 for translocation. Given the recent unique discoveries in mitochondrial biology, PNPase may participate in a novel pathway in the intermembrane space, which may be reflected by the new import pathway reported here.

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