The Evolutionary History of Peptidases Involved in the Processing of Organelle-Targeting Peptides

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Abstract

Most of the proteins present in mitochondria and chloroplasts, the organelles acquired via endosymbiotic events, are encoded in the nucleus and translated into the cytosol. Most of such nuclear-encoded proteins are specifically recognized via an N-terminal-encoded targeting peptide (TP) and imported into the organelles via a translocon machinery. Once imported, the TP is degraded by a succession of cleavage steps ensured by dedicated peptidases. Here, we retrace the evolution of the families of the mitochondrial processing peptidase (MPP), stromal processing peptidase (SPP), presequence protease (PreP), and organellar oligo-peptidase (OOP) that play a central role in TP processing and degradation across the tree of life. Their bacterial distributions are widespread but patchy, revealing unsurprisingly complex history of lateral transfers among bacteria. We provide evidence for the eukaryotic acquisition of MPP, OOP, and PreP by lateral gene transfers from bacteria at the time of the mitochondrial endosymbiosis. We show that the acquisition of SPP and of a second copy of OOP and PreP at the time of the chloroplast endosymbiosis was followed by a differential loss of one PreP paralog in photosynthetic eukaryotes. We identified some contrasting sequence conservations between bacterial and eukaryotic homologs that could reflect differences in the functional context of their peptidase activity. The close vicinity of the eukaryotic peptidases MPP and OOP to those of several bacterial pathogens, showing antimicrobial resistance, supports a scenario where such bacteria were instrumental in the establishment of the proteolytic pathway for TP degradation in organelles. The evidence for their role in the acquisition of PreP is weaker, and none is observed for SPP, although it cannot be excluded by the present study.

Key words: endosymbiosis, organellogenesis, chloroplast, mitochondria, subcellular targeting, proteolysis.

Significance

The peptidases involved in proteolysis of those peptides that allow the proper targeting of protein to mitochondria and chloroplast are of bacterial origin, but the evolutionary scenario of these acquisitions remained undetermined. Here, we compiled an exhaustive catalog of these peptidases across the tree of life and show how they were acquired once or twice, at the time of primary endosymbiosis, with the possible contributions of other bacteria than the chloroplast and the mitochondrial progenitors. These results shed new light on the origin of the degradation process of targeting peptides in organelles that may have involved bacteria resistant to antimicrobial peptides.

Introduction

Eukaryotic organelles, mitochondria and chloroplasts, evolved from ancestral bacteria through primary endosymbiosis. Mitochondria most probably derived from \( \alpha \)-proteobacteria (Wang and Wu 2015; Roger et al. 2017) and the chloroplast from an ancestral cyanobacterium, probably close to Gloeomargarita litophora (Jensen and Leister 2014; Ponce-Toledo et al. 2017). These endosymbiotic events were accompanied by massive horizontal gene transfer (HGT) of genetic material from the bacterial
progenitors to the host genome that is named endosymbiotic gene transfer (EGT) (Timmis et al. 2004). Today, most of the proteins found in mitochondria and chloroplast are encoded in the nucleus, translated in the cytosol, and subsequently imported into the organelles through dedicated translocon machineries: the Translocone of the Outer/Inner mitochondrial Membrane (TOM/TIM) for mitochondria (Wiedemann and Pfanner 2017) and the Translocon at the Outer/Inner envelope Membrane of Chloroplasts (TOC/TIC) for chloroplast (Chotewutmontri et al. 2017). In most cases, this import is driven by an N-terminal targeting peptide (TP).

The degradation of TPs in organelles is ensured by zinc metallopeptidases acting on different peptides according to their lengths (Johnson et al. 2006; Kmiec and Glaser 2012). The degradation follows common steps in the two organelles (Teixeira et al. 2017; Kmiec et al. 2018). First, two endopeptidases, the mitochondrial processing peptidase (MPP) (Taylor et al. 2001) and the stromal processing peptidase (SPP), cleave the TP, respectively, in the mitochondria and the chloroplast, thus releasing the mature protein. These enzymes have critical roles in cell viability. MPP is essential for cell viability in yeast (Yaffe et al. 1985) and nematodes (Nomura et al. 2006), and mutant of the MPP catalytic subunit cause neurodegeneration in early childhood (Vögtle et al. 2018). Similarly, SPP is essential in Arabidopsis thaliana (Trösch and Jarvis 2011) and also has been shown to impact protein import in pea (Zhong et al. 2003). The resulting TP is further degraded by the presequence protease (PreP) that cleaves peptides ranging from 10 to 65 amino acids (Kmiec and Glaser 2012) and by the organelar oligo-peptidase (OOP) that cleaves peptides ranging from 8 to 23 residues (Kmiec et al. 2013). In photosynthetic eukaryotes, both PreP and OOP are dually targeted to mitochondria and chloroplast. To the best of our knowledge, all described PrePs are targeted to organelles, while many OOP paralogs remain in the cytosol. In mitochondria, the cleavage by MPP may be followed by an additional cleavage by Oct1, an OOP homolog (Kmiec et al. 2013), or Icp55 in order to obtain the final mature protein, more stable than the intermediate form released after the MPP cleavage (Vögtle et al. 2009, 2011; Venne et al. 2013). Finally, single amino acids are released by the action of aminopeptidases from the M1 and M17 metalloprotease families acting on peptides <8 amino acids (Kmiec et al. 2018). MPP, SPP, and PreP are members of the M16 family, whereas OOP is part of the M3 family in the MEROPS hierarchical classification (Rawlings et al. 2018). In eukaryotes, MPP is a heterodimer, composed of a catalytic subunit, MPP-β in A. thaliana (Mas1 in yeast and PMPCB in human) and MPP-α (Mas2 and PMPCA) (Poveda-Huertes et al. 2017). Default of TP degradation can perturb organelle integrity and biogenesis, as free peptides may have a toxic effect (Kmiec et al. 2014). Also, human PreP is responsible for the degradation of amyloid-β peptides that accumulate into mitochondria in Alzheimer’s disease patients, which saturates the capacity for peptide degradation in the mitochondrion and leads to the accumulation of immature precursor proteins (Pinho et al. 2014). In yeast, in addition to Cym1 (the yeast PreP homolog), two other M16 proteases are involved in mTP degradation: Prd1 in the intermembrane space (Kambacheld et al. 2005) and Ste23, which is also able to cleave amyloid-β peptides (Taskin et al. 2017). For a complete overview of TP processing in mitochondria or chloroplast, including additional peptidases with a narrower set of substrates, see recent reviews (van Wijk 2015; Poveda-Huertes et al. 2017; Ghifari et al. 2019).

Peptidases from the M16, M3, M1, and M17 families involved in TP cleavage and degradation are well conserved from bacteria to eukaryotes, and M16 and M3 peptides may have been present in the Last Universal Common Ancestor (LUCA) (Rawlings et al. 2018; Rawlings and Bateman 2019). However, the MEROPS families are very large, with members carrying varied functions, because the classification is solely based on the part of the protein directly responsible for the peptidase activity (substrate binding site and the catalytic residues). Therefore, they are not adapted to accurately retrace the evolutionary history of a specific peptidase. The possible presence of an M16 and an M3 peptidase in LUCA does not imply that the functions now ensured by MPP, SPP, PreP, and OOP were present as well. Rather, previous studies all pointed for a bacterial origin of the organelle peptidases, as peptidases acting similarly are also found in bacteria, which is not the case for the protein degradation processes observed in the nucleus and the cytosol where these are mainly performed by the proteasome (Van Dyck and Langer 1999; Adam and Clarke 2002). It is of note that in photosynthetic eukaryotes, both mitochondria and chloroplasts, which arose from two distinct primary endosymbiotic events, use the same players (PreP and OOP) and players from the same peptidase family (MPP and SPP) to ensure disposal of their TPs.

We recently provided experimental evidence for an antimicrobial origin of TPs. A subclass of antimicrobial peptides (AMPs) are synthesized by the ribosome and adopt an amphipathic helical structure (HA-RAMPs) (Garrido et al. 2020), supporting the idea that the import machineries into organelles may derive from an antimicrobial resistance machinery in the bacteria involved in early endosymbiotic events (Wollman 2016; Caspari and Lafontaine 2021). In this view, the translocone machineries and the peptidases involved in the proteolytic degradation of TP may derive, respectively, from bacterial transporters and bacterial peptidases that participated in antimicrobial resistance. In agreement with this hypothesis, it has been demonstrated that MPP most probably originates from a progenitor of a rickettsiales putative peptidase (RPP) in a parasitic
bacterium, and that extant bacterial RPP still have the capability to cleave mitochondrial TPs (Kitada et al. 2007). To challenge this hypothesis further, here, we focused on the peptidases involved in the first steps of the main proteolytic pathway (MPP, SPP, PreP, and OOP), leaving aside more specific peptidases, as well as the house-keeping amionopeptidases that degrade small peptides <10 amino acids, acting at the very end of the proteolytic pathway. We therefore built the exhaustive catalog of MPP, SPP, PreP, and OOP families across eukaryotes, bacteria and archaea in order to decipher accurately their evolutionary history and to estimate the possible contribution of HA-RAMP-resistant bacteria in the acquisition the proteolytic pathway degrading organelle TPs.

Results

An Exhaustive Catalog of Organelle Peptidase Families

We searched for homologs of MPP, SPP, PreP, and OOP in 8,340 reference proteomes from Uniprot comprising 291 archaea, 6,820 bacteria, and 1,230 eukaryotes (including 135 photosynthetic eukaryotes). The MPP, SPP, PreP, and OOP sequences of A. thaliana were thus used as a starting point to construct Hidden Markov Model (HMM) profiles with the retrieved homologs (supplementary table S1, Supplementary Material online). HMM profiles allowed the detection of 36,807 candidates (10,930 PreP; 2,716 SPP; 6,822 OOP; and 16,339 MPP), from which, only the 26,211 candidates that contain all the Protein Families (PFAM) motifs present in the A. thaliana peptidase of reference were considered its homologs (table 1).

Given the paucity of those peptidases in the 291 archaeal proteomes present in the Uniprot Reference proteomes, we searched the Uniprot database to retrieve all archaeal sequences in the ca. 440 archaeal species absent from the Reference proteomes that possess either a M16 domain (429 sequences from 143 species) or a M3 domain (1,185 sequences from 623 species) following the procedure described above. We also searched homologs in the nonredundant (nr) database at NCBI (see Methods). The outcome of this search is provided in supplementary data S1, Supplementary Material online. In total, we found peptidase homologs in <7% of the 1,937 different archaea species with sequences referenced in the three databases. Among the 44 referenced Asgard species, which are considered the most closely related to eukaryotes, only one has an MPP homolog: Candidatus Thorarchaeota archaeaeon (strain OWC), and another one has an OOP homolog: Heimdallarchaeota archaeaeon (strain LC_2). PreP and SPP are absent from Asgard species.

On the proximity graph of the four peptidase families based on the BLAST pairwise E-value (supplementary fig. S1, Supplementary Material online), MPP, PreP, and SPP group together within the M16 peptidases families, well apart from OOP, belonging to the M3 family. Each family of M16 peptidase localizes differently from the two others, with an overlap between MPP and SPP.

The M16 Family

To better analyze the relationships among the peptidases of the M16 family, we built a phylogenetic tree of the 264 M16 homologs from the sample set (see Methods), based on the multiple alignment of the two first conserved domains (M16 and M16C) between MPP, PreP, and SPP (fig. 1). Each peptidase groups in three different subtrees in which both eukaryotic and bacterial homologs are found. The eukaryotic MPP subtree is divided into two subtrees, in line with an early duplication at the basis of the eukaryotic lineage, leading to the two subunits (a) and (b) of the MPP heterodimer. Only 28 homologs are misclassified on the tree: 2 PrePs and 16 SPPs localize within the MPP subtree; 1 MPP and 3 SPPs are found within the PreP subtree and 5 MPPs are found within the SPP subtree. Given that the tree is based on the first two conserved domains, and not the entire protein sequence, this could be due to sequences experiencing contrasting evolutionary rates. The rickettsiales putative peptidase (RPP), putative MPP ancestral progenitor, groups within the MPP subtree, confirming previous reports (Kitada et al. 2007). The bacterial metalloproteases FucS and HrpP that play a role in antimicrobial resistance mechanisms in Gram-negative bacteria group within the SPP subtree.

MPP, PreP, SPP, and OOP have a Bacterial Origin

Figure 2 describes the repartition of the homologs across the tree of life. MPP, PreP, and OOP peptidases are found, respectively, in 98%, 87%, and 75% of the eukaryotic proteomes, with 69% of the studied eukaryotic species possessing the three peptidases. The Rotosphaerida species Fongicula alba bears only the MPP and the two Chaoanoflagellata species lack PreP. Their frequencies are lower in bacterial lineages 72%, 53%, and 12% for MPP, OOP, and PreP, respectively, where they are heterogeneously distributed. There are very few homologs in Archaea: MPP and OOP are present only in 16 out of 291 archaeal proteomes and no PreP was found in any of these proteomes. They are also scarce in viruses, with only 13 MPPs, 2 OOPs, and no PreP among the 9,424 viral proteomes.

SPP is the least abundant peptidase, absent from archaea and viruses, present in only 21% of the bacterial proteomes and found only in photosynthetic eukaryotes (11% of the studied eukaryotic proteomes, including two of the three photosynthetic Stramenopiles present in our proteome database) and in nonphotosynthetic eukaryotes.
Table 1  
Composition of the Peptidase Families

| PFAM motif          | Archaea (291) |  | Bacteria (6,820) |  | Eukaryotes (1,230) |  |
|---------------------|---------------|---|------------------|---|--------------------|---|
|                     | #proteins     | #species | #proteins        | #species | #proteins         | #species |
| MPP                 | 11            | 10    | 10,812           | 4,972    | 3,966             | 1,204    |
| PreP                | 0             | 0     | 910              | 854      | 1,037             | 925      |
| SPP                 | 0             | 0     | 2,376            | 1,477    | 279               | 141      |
| OOP                 | 6             | 6     | 4,694            | 3,624    | 2,120             | 1,071    |

For each peptidase, the composition in PFAM motifs is given in the first column M16: PF00675; M16C: PF05193; M16C_assoc: PF08367; M3: PF01432. The number of homologs and number of corresponding proteomes across archaea, bacteria, and eukaryotes is given in the following columns. The total number of species considered is indicated in parenthesis for each domain.

**Fig. 1.**—Tree of a subset of M16 peptidases with 160 MPP, 60 SPP, and 44 PreP sequences used (see Methods). Branches are colored according to their bootstrap value (upper left box legend). The eukaryotic MPP subtree is divided into two subtrees, α and β, indicated with red dotted arrows. The inner circle gives the classification (upper left box legend) of the sequences among the three domains of life, with a distinction for bacteria with an experimentally documented resistance to AMP (supplementary data S1, Supplementary Material online). The outer circle gives a further classification (lower left box legend) distinguishing the taxonomic groups of interest in this study. Bacterial peptidases whose functions are discussed in the text are indicated, as well as homologs from model organisms present in the subset, with their names in parenthesis for bacteria and if paralogs exist for a given peptidase in eukaryotes. Cre: Chlamydomonas reinhardtii, Ath: Arabidopsis thaliana, Eco: Escherichia coli. Note that in this tree, the MPP-α2 of C. reinhardtii groups in the β subtree, while it groups correctly in the MPP-α subtree of figure 3.
Fig. 2.—Taxonomic distribution of the four peptidases among the tree of life. The cladogram represents the phylogenetic relationships between the different taxonomic groups (see Methods for the reference trees used). Circles next to leaves are colored according to the main groups of interest, with the proportion of AMP-resistant bacteria indicated in red. Minor bacterial phyla are colored in grey. Eukaryotes are divided into photosynthetic and nonphotosynthetic groups, according to the presence or the absence of a chloroplast. In the table, the first column indicates the number of proteomes included in each group. The number of proteomes where at least one homolog is found for MPP, PreP, SPP, and OOP are given in columns 2–5, respectively. The color gradient is proportional to the number of proteomes in the group with at least one homolog of the peptidase. On the right, the upper circle indicates the distribution of proteomes in the different groups; the circles below indicate the distribution of each peptidase among the different groups. PVC: superphylum composed of Planctomycetales, Verrucomicrobia, and Chlamydiales. FCB: superphylum composed of Fibrobacteres, Chlorobi, and Bacteroidetes. The color legend for the taxonomy is given at the upper right of the figure. For SPP, the homologs retrieved in Metazoa (1) and in Fungi (3) differ from those found in photosynthetic eukaryotes by an additional M16M PFAM domain that may impact their function (see text and supplementary data S2, Supplementary Material online).
containing a plastid like some Apicomplexa, Amoebozoa, and Alveolata species.

The taxonomic distribution of the four eukaryotic peptidases suggests that they most likely have a bacterial origin and that they were acquired early in eukaryote evolution. Only 23/1,230 (2%) eukaryotes are devoid of those four peptidases, including the 19 Microsporidia having lost their mitochondria, the parasitic nematode *Brugia timori*, the neuropathogenic flatworm *Trichobilharzia regenti*, as well as two Basidiomycota Fungi that are plant pathogens: the brown-rot fungus *Postia placenta* and *Moniliophthora perniciosa*. There are 9% (607 of 6819) of bacterial proteomes free of MPP, PreP, SPP, and OPP, among which there are four groups in which all species lack the peptidases: the 77 Mollicutes pathogens (55 Mycoplasmatales and 22 Entomoplasmatales), the 22 Enterococcus pathogenic species, and 48 Bifidobacteriales. The other 460 species are scattered among their respective groups.

The Evolutionary History of Peptidase Families

To get insights into the evolutionary history of each peptidase family from bacteria to eukaryotes, we built its phylogenetic tree and looked for the distribution of the
bacterial sequences that are close to eukaryotic homologs, which are the potential donors through gene transfers.

**MPP and SPP, the Cleavage Peptidases, were Inherited from a Single HGT**

On the MPP tree, all the eukaryotic MPP sequences form a monophyletic group among other bacteria (fig. 3), which suggests that the ancestral eukaryotic MPP has been acquired by a single HGT. Note that because there are more than 14,000 MPP homologs, the tree is collapsed. The eukaryote MPP subtree forms a monophyletic group with some homologs from α-proteobacteria (see supplementary fig. S2, Supplementary Material online for a tree with a sampling of bacterial homologs), in agreement with an EGT from the mitochondrial progenitor during or following the early stages of mitochondrial endosymbiosis. In some bacterial homologs, the M16–M16C region is repeated. Given their heterogeneous distribution on the tree presented in supplementary fig. S3A, Supplementary Material online, it is unlikely that it represents an ancestral form of MPP. 132 additional PFAM domains were identified in our retrieved candidates. These additional domains are heterogeneously distributed and shared by <8 species each, with two exceptions found in the leotiomyceta clade in the Ascomycota phylum (supplementary data S2, Supplementary Material online). The addition of these domains could suggest two independent subfunctionalizations, one in Eurotiomycetes and one in Leotiomycetes.

On the SPP tree, homologs from photosynthetic eukaryotes are divided into two subtrees (fig. 4). A first one as a sister group of sequences from the superphylum composed of Fibrobacteres, Chlorobi, and Bacteroidetes (FCB) in the lower part of the tree (euka1), and a second one at the basis of the larger group containing the euka1 subtree (euka2). The majority (93/141) of eukaryotic species have a paralog in both subtrees. This topology can be explained by an acquisition from a FCB bacteria by HGT at the basis of the photosynthetic eukaryotic lineage, followed by a duplication event and the divergent evolution of one of the paralogous copies. The conservation of the domain architecture of the SPP homologs from the lower subtree containing the euka1 and euka2 subtrees (supplementary fig. S3B, Supplementary Material online) supports this hypothesis. Note that homologs from euka2 appear to bear an additional N-terminal part, compared to all other homologs from the lower subtree that could explain their basal position. Noticeably, there are very few SPP homologs in cyanobacteria (64 from the 142 cyanobacterial proteomes studied), most closely related to chloroplast, and they form a monophyletic group distant from the euka1 and euka2 subtrees. 33 species (28 bacteria and 5 eukaryotes) and 8 others (7 bacteria and 1 eukaryote) have an additional M16M and M16C_assoc PFAM domain in their identified SPP homologs, respectively. The other 26 additional PFAM domains in SPP candidates are found in <4 species each (supplementary data S2, Supplementary Material online).

**OOP and PreP Degrade the TP after their Cleavage has been Acquired Twice in Close Relationship with the Two Primary Endosymbioses**

On the complete OOP tree (fig. 5), a large group of 1,393 eukaryotic sequences, 110 of which are from photosynthetic eukaryotes, form a sister group in the lower left subtree with mainly Proteobacteria and Terrabacteria, suggesting an acquisition from these bacteria at an early step of eukaryogenesis. Another 171 sequences from photosynthetic eukaryotes, 120 cyanobacteria, 15 PVC, 5 Proteobacteria, and 1 Chlamydiales, a potential partner of the chloroplast endosymbiosis (Ball et al. 2016) form a monophyletic group in the upper left part of the tree (fig. 5), suggesting a second acquisition of a bacterial OOP during chloroplast endosymbiosis. The majority of photosynthetic eukaryotes (85 over 123) has a paralog in both eukaryotic subtrees, thus they conserved the two acquired OOP copies. As the OOP family contains many paralogs (172 in eukaryotes on average) some of which being cytosolic (three in A. thaliana), we built a tree with eukaryotic OOP that are predicted to be addressed to either the chloroplast or the mitochondria and their closest bacterial homologs (one per species). The taxonomic distribution of the peptidases remains unchanged and so are the evolutionary conclusions (supplementary fig. S5, Supplementary Material online). 135 additional PFAM domains were found in OOP candidates, found in <10 species each (supplementary data S2, Supplementary Material online).

The eukaryotic PreP homologs are distributed among three subtrees (fig. 6). The vast majority of homologs from nonphotosynthetic eukaryotes (770) form a monophyletic group with a sister bacterial clade mainly composed of Proteobacteria (NPSeuka subtree, fig. 6) and all but 5 (148) PreP from photosynthetic eukaryotes form a monophyletic group with a sister bacterial clade composed of homologs form Proteobacteria and Terrabacteria (PSeuka subtree, fig. 6). Another 97 Opisthokonta sequences (from 91 distinct species) form a long-branched group at the basis of the tree (Baseuka subtree, fig. 6), suggesting they are highly diverged homologs that could be related to either the NPSeuka or the Pseuka subtree. Ignoring the Baseuka subtree, the tree topology suggests two independent acquisitions of PreP in eukaryotes: one associated with the mitochondrial endosymbiosis (within the NPSeuka subtree) and a posterior one that could be associated to the chloroplast endosymbiosis (within the Pseuka subtree). Among bacteria, PreP is present most exclusively in Terrabacteria (493 homologs) and in Proteobacteria (263) (fig. 2 and supplementary data S1, Supplementary
Material online), containing the cyanobacteria and α-proteobacteria, respectively, the extant bacteria most closely related to chloroplast and mitochondria. The nearly complete absence of homologs from photosynthetic eukaryotes in the NPSeuka subtree suggests that the PreP copy acquired at the time of the mitochondrial endosymbiosis has been lost after the acquisition of the chloroplast, except in five microalgae. The timing of the second PreP acquisition relatively to the chloroplast endosymbiosis is difficult to determine with the present data when considering the Baseuka subtree: half of these Opisthokonta homologs correspond to paralogs from the NPSeuka subtree, supporting an independent duplication before the Opisthokonta divergence followed by rapid evolution, and a second PreP acquisition during or after the chloroplast endosymbiosis. But the remaining half sequences in the Baseuka subtree have no paralog in the NPSeuka subtree and could also correspond to highly diverged homologs from the Pseuka subtree. This would imply an alternative, but less parsimonious scenario, in which the two acquisitions of PreP occurred before the chloroplast endosymbiosis so that ancestral eukaryotes had two PrePs and one copy has been differentially lost depending on the presence of a chloroplast in the majority of the lineages. In support of this alternative scenario, the domain composition of PreP homologs in the Baseuka subtree is identical to the one in the Pseuka subtree (supplementary fig. S3C, Supplementary Material online). 27 additional PFAM domains were found in PreP candidates, found in <7 species each (supplementary data S2, Supplementary Material online).

**FIG. 4.**—Phylogenetic tree of the 2,652 SPP homologs. Five additional sequences are used as outgroup (see supplementary data S1, Supplementary Material online). Branches are colored according to the classification given in the left lower part of the figure. The inner circle recapitulates the taxonomy. Branches with support higher than 0.8 are depicted in bold. Homologs from model organisms are indicated next to the inner circle, with the name of the *Escherichia coli* homolog in parenthesis. Cre: *Chlamydomonas reinhardtii*, Ath: *Arabidopsis thaliana*, Eco: *E. coli*. See supplementary figure 3A, Supplementary Material online for a collapsed version of this tree.
Molecular Evolution of the Peptidase Motifs from Bacteria to Eukaryotes

To get further insights into the evolution of the function of these peptidases, we compared the conserved motifs containing their catalytic sites. The logo representations of the multiple alignments of the M16 motif in the MPP family are presented in figure 7. Eukaryotic homologs in the MPP tree were annotated as MPP-$\alpha$ or MPP-$\beta$ according to their localization in either the subtree $\alpha$ or the subtree $\beta$ (fig. 3). The 27 MPP homologs grouping outside these two subtrees were not considered.

The logos for MPP-$\beta$ (fig. 7B) and for MPP in the 200 bacteria that are the most closely related to eukaryotic homologs, hereafter referred to as top200-bacteria (fig. 7D) show a much higher overall conservation (indicated by the height of the stacks in the logo) than those of MPP-$\alpha$ (fig. 7A) and MPPs in the bulk of bacteria (fig. 7C). Further comparison of the logos on figure 7B and D shows however that the most frequent amino acid at a given position is not conserved in most cases. A noticeable exception is the zinc-binding motif HXXEHX$_7$E (red boxes in fig. 7) required for the activity of the catalytic subunit MPP-$\beta$ (fig. 7B) which is well conserved in the top200-bacteria and in the bulk of bacterial homologs (fig. 7C and D), suggesting that it was already present in the bacterial donors of MPP.

Note that HXXEH has been lost in MPP-$\alpha$ which is consistent with it being an inactive form in the MPP heterodimer found in eukaryotes (fig. 8).

Contrary to the eukaryotic MPPs, bacterial ones do not function in a heterodimeric context. A possible signature of this functional shift from bacteria to eukaryotes can be found in the HXXEH motif that reads HFLEH in MPP-$\beta$ but HLLEH in the MPP from the top200-bacteria (fig. 7B and C). Also, the
positions 98–103 (green boxes, fig. 7) read HLNAYT in MPP-β but EXNAXT in the top200-bacteria and XXNAXT in the bulk of bacteria. In MPP-β, this region corresponds to the beta strand located at the active site (strand 1, fig. 8) within the binding region of the mitochondrial TP, delimited by the edges of four β sheets (Taylor et al. 2001). In the inactive MPP-α, there is no conservation at these positions, which correspond to strand 4 of the substrate binding region (fig. 8). Several other changes in the logos between MPP-β and bacterial MPPs are also likely to be related to the functional shift from bacteria to eukaryotes, like positions 158–167 (grey box, fig. 7) that are not conserved in the bulk of bacterial MPPs except for the proline at position 158, read PDDVVFDRFX in the top200-bacteria but becomes EEVVFDHLH in MPP-β. Logos of the multiple alignments for the other peptidases are given in supplementary figures S6, S7, S8, Supplementary Material online. Similar observations suggesting a possible functional shift from bacteria to eukaryotes can be made for OOP and SPP, but not for PreP.

### Some Peptidases from AMP-Resistant Bacteria are Close to their Eukaryotic Homologs

Next, we estimated the extent at which AMP-resistant bacteria could have been involved in the acquisition of the peptidases from the proteolytic degradation pathway of organellar TP. It is worth mentioning here that our definition of AMP-resistant bacteria is restricted to those bacteria for which there are publications that document an AMP-resistance experimentally. A number of bacteria has not been studied yet in that respect.

For each peptidase family, we determined the distance of each bacterium to its closest eukaryotic homolog according to three different metrics: 1) the evolutionary distance between each pair of species, that is, the sum of branch length separating two leaves of the complete gene family tree, 2) the topological distance between each pair of species, corresponding to the number of internal nodes separating a pair of species on the complete gene family tree,
and 3) the log(ε-value) of the best Blast hit between two sequences (see Methods). We took these three metrics into account to avoid erroneous conclusions due to phylogenetic reconstruction uncertainties, considering they will be somehow compensated. Figure 9 presents the evolutionary distance of all bacterial sequences to their closest eukaryote counterparts. We defined as the top200 set, the 200 bacteria that are most proximal to their eukaryotic homologs in each group, for a given metric, and pan-top200 corresponds to the union of the top200 sets of three metrics (supplementary table S2, Supplementary Material online).

For the MPP family, 22% (114/521) of homologs of the pan-top200 belongs to resistant-bacteria among which 46 Pseudomonas and 35 Rhizobium (α-proteobacteria). For the OOP family, 10% (40/402) of homologs of the pan-top200 belongs to resistant-bacteria, mainly γ-proteobacteria, with 30 Enterobacteria and 8 Pseudomonadales. Also, a significant part (85%) of the homologs from resistant proteobacteria groups with the eukaryote subtree at the upper left (fig. 6).

At the genus level, Pseudomonas species have homologs in the pan-top200 of MPP, OOP, and SPP, and Acinetobacter...
have close homologs in the pan-top200 of SPP, PreP, and OOP. *Borreliella* species have homologs in the pan-top200 of SPP and PreP; *Escherichia* and *Acinetobacter* species have homologs in the pan-top200 of OOP and SPP. However, only few bacterial species have peptidases homologs in the top-200 of more than one family. The most notable case is four *Pseudomonas* species (*P. agarici, P. cremoricolorata, P. fluorescens F113* and *P. sp. 5*) with a homolog in the OOP pan-top200 and a homolog in the MPP pan-top200.

Regardless of their distance to eukaryotic homologs, co-occurrences of pairs of the four peptidases in AMP-resistant bacteria are significantly less frequent than observed in the whole bacterial clade. However, the most frequent co-occurrence of peptidases among the 658 AMP-resistant bacteria is observed for OOP and MPP, in 167 species (25%), notably in all studied *Pseudomonas* and FCB species, and in the majority of the α and γ-proteobacteria. This co-occurrence of OOP and MPP is significantly higher in resistant α–proteobacteria than in non-resistant α–proteobacteria. Similarly, the co-occurrence of MPP and PreP is higher in resistant Terrabacteria than in non-resistant Terrabacteria. The co-occurrence of OOP and SPP is observed in 58 AMP-resistant bacterial species (8%), including 28 *Vibrio*. The co-occurrence of PreP and OOP is observed only in the 15 Acinetobacter species (supplementary fig. S9, Supplementary Material online). Note that only 98 out of the 658 AMP-resistant bacteria studied (15%) are devoid of homolog for the four studied peptidases.

### Discussion

We established a complete version of the homologous families of the peptidases involved in the proteolytic pathway of TPs in mitochondria and chloroplast, recapitulated in figure 10. Indeed, deep phylogenies are based on very ancient signal and incorrectly account for the very different modes of genetic transmission between bacteria and eukaryotes, and are therefore hardly devoid of errors. In an attempt to avoid, as much as we could, erroneous conclusions due to phylogenetic artifacts, we used a simple strategy-mixing metrics based on evolutionary models and a simple similarity distance. If our phylogenies are error-free, our results confirm that MPP, PreP, SPP, and OOP were inherited from bacterial ancestors, but not necessarily the organelle progenitors. Indeed, we found no SPP and PreP homologs in Archaea and less than a dozen of archaeal MPP and OOP homologs. The scarce distribution on the archaeal homolog on the MPP and OOP gene trees provide no evidence for their presence in LUCA, but rather for an acquisition from bacteria. Such HGT from Bacteria to Archaea are estimated to be 5-fold more frequent than HGT from Archaea to Bacteria, and have been largely documented, see, for example, Nelson-Sathi et al. (2015).

In the presentation of our results, we assumed that HGT occurred shortly after or during the endosymbiosis, directly from ancestral bacteria close to the extant ones, to the ancestral eukaryotes. This implies that bacterial partners other than the endosymbiont ancestors would have been involved in such events, as proposed earlier (Gray 2015;
Another and non-mutually exclusive possibility is that the ancestors of mitochondria and chloroplast have transferred to the host cell part of their ancestral pan-genome, that is, the entire gene set of their species, which was different from the present pan-genome. Thus, the present phylogenetic proximities observed with bacteria other than cyanobacteria and proteobacteria could also be caused by posterior HGT events among prokaryotes and may reflect that they were part of the ancestral pan-genome of mitochondria and chloroplast precursors (Ku et al. 2015). Whatever the actual source of these transfers, genetic properties that
are absent from the present repertoire of extant bacteria close to the endosymbiont ancestors are likely to have played an important role for the establishment and the maintenance of the endosymbionts, notably from pathogenic partners that helped to escape host defense mechanism. Also, we found several molecular signatures specific of either bacterial or eukaryotic homologs, which could play a role in the functional shift in their activity in eukaryotes, except for PreP. Further biochemical investigations should be undertaken to assess the significance of these observations.

Our results suggest that some AMP-resistant bacteria could have contributed to the setup of the TP proteolytic pathway. MPP was most probably inherited by EGT from an α-proteobacteria-like mitochondrion ancestor, most probably pathogenic. This is in line with previous findings that the rickettsiales putative peptidase (RPP) in the obligate intracellular *Rickettsiales* is able to cleave mTP from a couple of yeast, human, and mouse preproteins. Most notably, RPP can induce the peptidase activity of the yeast MPP-β subunit for the processing of proteins, that MPP-β normally does not perform (Kitada et al. 2007). It is thus a reasonable hypothesis to consider that the ancestral eukaryotic MPP already had this property, which would have been crucial for adaptation of the ancestral endosymbiont. A first copy of OOP would also have been acquired from proteobacteria, at the time of the mitochondrial endosymbiosis, with a possible role of AMP-resistant α and γ-proteobacteria. A second acquisition of OOP probably occurred at the time of chloroplast endosymbiosis, with a possible role of a pathogenic Chlamydia, in agreement with the “Ménage à trois” hypothesis (MATH), in which pathogenic Chlamydiae provided critical help for maintenance of the chloroplast ancestor into the protist host (Greiner and Bock 2013; Ball et al. 2016; Cenci et al. 2016, 2017). As for OOP, PreP would have been acquired twice most probably from a resistant Terrabacteria or a Proteobacteria. The first copy would have been acquired at the time of mitochondrial endosymbiosis and the second copy at the time of the chloroplast endosymbiosis, with the first acquired copy being lost after the chloroplast endosymbiosis in photosynthetic eukaryotes. If this hypothesis is true, then the second acquired copy works not only in the chloroplast, but also in the mitochondria, as in *A. thaliana* where it has been experimentally demonstrated (Kmiec et al. 2014). The chloroplast counterpart of MPP, SPP would have been acquired during chloroplast endosymbiosis from an FCB bacteria with no evidence of contribution from resistant-bacteria.

Our results are based on a minimal group of bacteria for which experimental evidence of resistance to AMPs has been described, with a considerable and expected bias towards Enterobacteria. In particular, to our knowledge there are no reports of studies regarding possible resistance to

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**Fig. 10.**—Scenario of gains and losses of the four peptidases across the tree of life. Up arrows represent horizontal transfer. Peptidase names and up arrows are colored according to the bacterial donor group. Down arrows represent loss. Superscript “D” indicates a basal duplication in the group. In the table, the number of proteomes containing at least one homolog of each peptidase is indicated. The heatmap indicates the percentage of the proteomes with at least one homolog within each group. This figure has been created with Biorender.com.
AMPs among cyanobacteria, from which originated the chloroplast. On the other hand, we considered a whole bacterial genus as potentially antimicrobial resistant but functional evidence is not available for every single species, leading to possible over-interpretation that should however be minimal if orthologous conservation is a proxy for functional conservation.

In conclusion, AMP-resistant bacteria have OOP, MPP, and PreP homologs in the closest vicinity to their eukaryotic counterparts, which is consistent with their possible implication in the early stages of mitochondrial endosymbiosis and for OOP and PreP, also in the early stage of the chloroplast endosymbiosis. It is of note that very few SPP homologs from resistant-bacteria were found close to SPP in photosynthetic eukaryotes.

As Microsporidia lost their mitochondria after the divergence of Fungi, the absence of MPP and PreP, found exclusively in organelles is expected. As many OOP paralogs remain cytosolic, the fact that Microsporidia also lack OOP is consistent with a possible action of OOP on proteins that failed to be imported or that were released from mitochondria. The fact that the four other eukaryotes devoid of the four peptidases are all pathogens could suggest an unusual mitochondrial metabolism in those species.

With our previous finding that a certain class of HA-RAMPs and organelle TPs are likely evolutionary related, based on their shared physico-chemical properties, and their ability to functionally complement each other (Garrido et al. 2020; Caspari et al. 2021) the present phylogeny of organellar peptidases is fairly consistent with the hypothesis that the organelle-targeting machineries derive from an antimicrobial resistance (Wollman 2016; Caspari and Lafontaine 2021), through an import and destroy mechanism that has been documented in several studies. In that respect, the fact that the co-occurrence of MPP and PreP/OOP is higher in AMP-resistant α-proteobacteria/Terrabacteria compared to sensitive α-proteobacteria/Terrabacteria suggests that MPP and PreP/OOP were likely to be involved in the destroy steps. Moreover, we note that the Gram-negative γ-proteobacteria Pseudomonas and Acinetobacter are opportunistic pathogen that possess MPP, OOP, and SPP homologs closely related to their eukaryotic counterparts, making them model organisms of choice that possess the major part of the proteolytic pathways for the degradation of AMP and AMP-derived TPs. In Pseudomonas aeruginosa HA-RAMP activity requires its binding to outer membrane proteins. This could probably promote HA-RAMP uptake and cytoplasmic protrusion as it has been proposed for the HA-RAMP hRNase 7 with the outermembrane protein OprI (Lin et al. 2010), and for the HA-RAMP LL-37 with the porin OmpA (Lin et al. 2015).

Other HA-RAMPs are actively internalized via import pump, like the bacteriocin pyocin S2 that can hijack the iron transporter FpvAI, a β-barrel TonB-dependent transporter (TBDTs) in P. aeruginosa (White et al. 2017). Such hijacking of TBDTs by bacteriocins is also observed in the Gram-negative phytopathogens Pectobacterium species in which pectocin M1 and M2 containing a ferredoxin domain can enter the bacterial cell through iron-uptake (Grinter et al. 2012, 2014). In those phytopathogens, FucS is associated with the TBDTs to promote the import of ferredoxin across the outer membrane (Grinter et al. 2018). This protease-associated import system capable of HA-RAMP uptake is widespread in Gram-negative bacteria (Grinter et al. 2019) and suggests that such association could have played a role in the establishment of an early protein import system upon endosymbiosis.

Further investigations are needed to estimate the evolutionary relationships between bacterial peptide uptake mechanisms in resistant bacteria and the TIM/TOM and TIC/TOC complexes. Indeed, several subunits of the extant translocons are of bacterial origin. For example, the protein core of TOC/TIC, composed of Toc75, Toc136, and TIC110, is thought to be derived from the bacterial β-barrel protein export system BAM/TAM, composed of BamA/TamA (OmpB5 and Toc75 homologs) and TamB (Tic136 homolog) (Day and Theg 2018; Richardson and Schnell 2020). Similarly, mitochondrial outer membrane β-barrels (VDAC, TOM40, but also lineage-specific outer membrane β-barrels like ATOM in trypanosoma) are homologous and thought to derive from bacterial primitive channels (Pereira and Lupas 2018), and Tim23, Tim44 and Tim 14 of the TIM23 complex also have bacterial homologs (Rassow et al. 1999; Clements et al. 2009).

**Methods**

Peptidase Sequences and Proteomes

Reference sequences were selected for each of the four peptidases in the photosynthetic model plant A. thaliana: the two members of the PreP family PreP1, PreP2; the three MPP subunits: MPPα1, MPPα2, MPPβ; OOP and SPP, see supplementary table S1, Supplementary Material online for accession numbers. The search for homologs was performed against a database composed of the 8,340 nonvirus proteomes from the Uniprot reference database (released version 26 February 2020), including 291 archaea, 6,819 bacteria, and 1,230 eukaryote proteomes.

Family Reconstruction

First, the proteome database was searched for significant hits (minimum query coverage of 70% and E-value <10^-5) with BLASTP (standalone version 2.6.0) (Altschul et al. 1997) for each of the reference proteases (table 1). In order to retrieve remote homologs from the A. thaliana sequences, we built some HMM profiles for each peptidase with hmmbuild. Hits for the highly similar paralogs PreP1
ogeny is taken from Burki et al. (2020) and Torruella et al. was determined with hmmsearch (hmmer version 3.2.1) as homologs. The presence of additional conserved domains against all (El-Gebali et al. 2019). The phylogenetic relationships between the organisms included in our analysis were taken from the literature. The phylogeny of the Archaea as well as the placement of eukaryotes is based on Spang et al. (2017). The Eukaryota phylogeny is taken from Burki et al. (2020) and Torruella et al. (2015) for Opistokontha. The bacterial phylogeny is extracted from Castelle and Banfield (2018) and Zhu et al. (2019).

Peptidase Tree

The concatenation of the aligned PFAM motifs identified in each peptidase family was performed with MAFFT v7.450 (Katoh and Standley 2013). Trees were reconstructed with the Approximately Maximum-Likelihood method FastTree (Price et al. 2010) (v2.1.11) with the JTT+CAT model. The local support values given by FastTree are based on the Shimodaira–Hasegawa (SH) test (Price et al. 2010).

We used the NCBI taxonomy retrieved with ETE3 toolkit to perform trees annotation. Tree display was performed with iTOL website (Letunic and Bork 2019). The phylogenetic analysis of the M16 family was performed with the sample set, using only the two first motifs M16 and M16C shared by the three M16 peptidases. For each family, MPP, SPP, PreP, and OOP, the alignments were performed with only the conserved motifs given in table 1. The sequences used as outgroups for the M16 peptidases are given in supplementary data S1, Supplementary Material online. The OOP tree was rooted with the midpoint method.

Minimal Distance to Eukaryote

The evolutionary distance is estimated by the sum of the branch length between a given bacterium and its closest eukaryotic homolog. The topological distance corresponds to the number of internal nodes separating a bacterium to its closest eukaryotic homolog. Both distances were computed with python ETE3 toolkit (3.1.1). The pairwise distance corresponds to the log(E-value) Blastp between a bacterial sequence and its eukaryote best hit, with the BLAST version 2.6.0 (Altschul et al. 1997).

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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Data Availability

Supplementary data S2, phylogenetic trees in newick format, all Python and R in-house scripts are available at https://github.com/UMR7141/Peptidase_Analysis.

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Literature Cited

Adam Z, Clarke AK. 2002. Cutting edge of chloroplast proteolysis. Trends Plant Sci. 7:451–456. doi:10.1016/S1360-1385(02)02326-9

Altschul SF, et al. 1997. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25:3389–3402. doi:10.1093/nar/25.17.3389

Ball SG, Bhattacharya D, Weber APM. 2016. Pathogen to powerhouse. Genome Biol. Evol. 8:3702–3704. doi:10.1093/gbe/evw256

Burki F, Roger AJ, Brown MW, Simpson AGB. 2020. The new tree of eukaryotes. Trends Ecol Evol. 35:43–55. doi:10.1016/j.tree.2019.08.008

Caspari OD, Garrido C, Choquet Y, Wollman F-A, Lafontaine I. 2020. The role of antimicrobial peptides in organelle proteolysis. PLoS Pathogens 17:e1009466. doi:10.1371/journal.ppat.1009466

Castelle CJ, Banfield JF. 2018. Major new microbial groups expand diversity and alter our understanding of the tree of life. Cell 172:1181–1197. doi:10.1016/j.cell.2018.02.016

Cenci U, et al. 2017. Biotic host–pathogen interactions as major drivers of plastid endosymbiosis. Trends Plant Sci. 22:316–328. doi:10.1016/j.tplants.2016.12.007

Cenci U, Ducatez M, Kadouche D, Colleoni C, Ball SG. 2016. Was the evidence supporting an antimicrobial origin of targeting peptides acquired by bacteria for iron piracy against plants. PLOS Biol. 14:e1002345. doi:10.1016/j.jmb.2019.03.008

Chotewutmontri P, Holbrook K, Bruce BD. 2017. Chapter six - plastid endosymbiosis. BioEssays 35:354–365. doi:10.1002/bies.20160262

Kmiec B, Glaser E. 2012. A novel mitochondrial and chloroplast peptidase, PreP. Physiol Plant. 145:180–186. doi:10.1111/j.1399-5137.2011.01531.x

Kmiec B, Teixeira PF, Glaser E. 2014. Shredding the signal: targeting peptide degradation in mitochondria and chloroplasts. Mol Plant 11:342–345. doi:10.1090/molplant/1190. doi:10.1093/molbev/msv183

Greiner S, Bock R. 2013. Tuning a ménage à trois: co-evolution and co-adaptation of nuclear and organellar genomes in plants. BioEssays 35:354–365. doi:10.1002/bies.201200137

Kmiec B, et al. 2018. FusC, a member of the M16 protease family acquired by bacteria for iron piracy against plants. PLOS Biol. 16:e2006026. doi:10.1371/journal.pbio.2006026

Gray MW. 2015. Mosaic nature of the mitochondrial proteome: Implications for the origin and evolution of mitochondria. Proc Natl Acad Sci U S A. 112:10133–10138. doi:10.1073/pnas.1421379112
cytochrome bc1 complex of *Caenorhabditis elegans*, a model parasitic nematode. J Biochem. 139:967–979. doi:10.1093/jb/mvj114
Pereira J, Lupas AN. 2018. The origin of mitochondria-specific outer membrane β-barrels from an ancestral bacterial fragment. Genome Biol Evol. 10:2759–2765. doi:10.1093/gbe/evy216
Pinho CM, Teixeira PF, Glaser E. 2014. Mitochondrial import and degradation of amyloid-β peptide. Biochim Biophys Acta (BBA) - Bioenergetics 1837:1069–1074. doi:10.1016/j.bbapap.2014.02.007
Ponce-Toledo RI, et al. 2017. An early-branching freshwater cyanobacterium at the origin of plastids. Curr Biol 27:386–391. doi:10.1016/j.cub.2016.11.056
Poveda-Huertes D, Mulica P, Vögtle FN. 2017. The versatility of the enzyme, their substrates and inhibitors in 2017 and a comparison approximately
Price MN, Dehal PS, Arkin AP. 2010. FastTree 2 – approximately maximum-likelihood trees for large alignments. PLoS One 5: e9490. doi:10.1371/journal.pone.009490
Rassow J, Dekker PJ, van Wilpe S, Moll J, Soll J. 1999. The preprotein translocase of the mitochondrial inner membrane: function and evolution. J Mol Biol. 286:105–120. doi:10.1006/jmbi.1998.2455
Rawlings ND, et al. 2018. The MEROPS database of proteolytic enzymes, their substrates and inhibitors in 2017 and a comparison with peptidases in the PANTHER database. Nucleic Acids Res. 46: D624–D632. doi:10.1093/nar/gkx1134
Rawlings ND, Bateman A. 2019. Origins of peptidases. Biochimie 166: 4–18. doi:10.1016/j.biochi.2019.07.026
Richardson LG, Schnell DJ. 2020. Origins, function, and regulation of the TOC–TIC general protein import machinery of plastids. J Exp Bot. 71:1226–1238. doi:10.1093/jxb/erz517
Roger AJ, Muñoz-Gómez SA, Kamikawa R. 2017. The origin and diversification of mitochondria. Curr Biol 27:R1177–R1192. doi:10.1016/j.cub.2017.09.015
Spang A, Caceres EF, Ettema TJG. 2017. Genomic exploration of the diversity, ecology, and evolution of the archaeal domain of life. Science 357:eaaf3883. doi:10.1126/science.aaf3883
Taskin AA, et al. 2017. The novel mitochondrial matrix protease Ste23 is required for efficient presequence degradation and processing. Mol Biol Cell 28:997–1002. doi:10.1091/mbc.e16-10-0732
Taylor AB, et al. 2001. Crystal structures of mitochondrial processing peptidase reveal the mode for specific cleavage of import signal sequences. Structure 9:615–625. doi:10.1016/S0969-2126(01)00621-9
Teixeira PF, et al. 2017. A multi-step peptidolytic cascade for amino acid recovery in chloroplasts. Nat Chem Biol 13:15–17. doi:10.1038/nchembio.2227
Timmis JN, Ayliffe MA, Huang CY, Martin W. 2004. Endosymbiotic gene transfer: organelle genomes forge eukaryotic chromosomes. Nat Rev Genet. 5:123–135. doi:10.1038/ng1271
Torruella G, et al. 2015. Phylogenomics reveals convergent evolution of lifestyles in close relatives of animals and fungi. Curr Biol. 25: 2404–2410. doi:10.1016/j.cub.2015.07.053
Trösch R, Jarvis P. 2011. The stromal processing peptidase of chloroplasts is essential in arabidopsis, with knockout mutations causing embryo arrest after the 16-cell stage. PLoS One 6:e23039. doi:10.1371/journal.pone.0023039
Van Dyck L, Langer T. 1999. ATP-dependent proteases controlling mitochondrial function in the yeast Saccharomyces cerevisiae. CMLS, Cell Mol Life Sci. 56:825–842. doi:10.1007/s000180050029
van Wijk KJ. 2015. Protein maturation and proteolysis in plant plastids, mitochondria, and peroxisomes. Annu Rev Plant Biol. 66:75–111. doi:10.1146/annurev-arplant-043014-115547
Venne AS, Vögtle F-N, Meisinger A, Zahedi RP. 2013. Novel highly sensitive, specific, and straightforward strategy for comprehensive N-terminal proteomics reveals unknown substrates of the mitochondrial peptidase lcp55. J Proteome Res. 12: 3823–3830. doi:10.1021/pr400435d
Vögtle F-N, et al. 2009. Global analysis of the mitochondrial N-proteome identifies a processing peptidase critical for protein stability. Cell 139:428–439. doi:10.1016/j.cell.2009.07.045
Vögtle F-N, et al. 2011. Mitochondrial protein turnover: role of the precursor intermediate peptidase Oct1 in protein stabilization. Mol Biol Cell 22:2135–2143. doi:10.1091/mbc.e11-02-0169
Vögtle F-N, et al. 2018. Mutations in PMPCB encoding the catalytic subunit of the mitochondrial presequence peptidase cause neurodegeneration in early childhood. Am J Hum Genet. 102: 557–573. doi:10.1016/j.ajhg.2018.02.014
Wang Z, Wu M. 2015. An integrated phylogenomic approach toward pinpointing the origin of mitochondria. Sci Rep. 5:7949. doi:10.1038/srep07949
White P, et al. 2017. Exploitation of an iron transporter for bacterial protein antibiotic import. Proc Natl Acad Sci U S A. 114:12051–12056. doi:10.1073/pnas.1713471114
Wiedemann N, Pfanner N. 2017. Mitochondrial machineries for protein import and assembly. Annu Rev Biochem. 86:685–714. doi:10.1146/annurev-biochem-060816-060151
Wollman F-A. 2016. An antimicrobial origin of transit peptides accounts for early endosymbiotic events. Traffic 17:1322–1328. doi:10.1111/tra.12446
Yaffe MP, Ohta S, Schatz G. 1985. A yeast mutant temperature-sensitive for mitochondrial assembly is deficient in a mitochondrial presequence activity that cleaves imported precursor polypeptides. EMBO J. 4: 2069–2074. doi:10.1002/j.1460-2075.1985.tb03893.x
Zhong R, Wan J, Jin R, Lampaa G. 2003. A pea antisense gene for the chloroplast stromal processing peptidase yields seedling lethals in Arabidopsis: survivors show defective GFP import in vivo. Plant J. 34:802–812. doi:10.1046/j.1365-313X.2003.01772.x
Zhu Q, et al. 2019. Phylogenomics of 10,575 genomes reveals evolutionary proximity between domains Bacteria and Archaea. Nat Commun. 10:5477. doi:10.1038/s41467-019-13443-4