Discovery of EMRE in fungi resolves the true evolutionary history of the mitochondrial calcium uniporter

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Calcium (Ca²⁺) influx into mitochondria occurs through a Ca²⁺-selective uniporter channel, which regulates essential cellular processes in eukaryotic organisms. Previous evolutionary analyses of its pore-forming subunits MCU and EMRE, and gatekeeper MICU1, pinpointed an evolutionary paradox: the presence of MCU homologs in fungal species devoid of any other uniporter components and of mt-Ca²⁺ uptake. Here, we trace the mt-Ca²⁺ uniporter evolution across 1,156 fully-sequenced eukaryotes and show that animal and fungal MCUs represent two distinct paralogous subfamilies originating from an ancestral duplication. Accordingly, we find EMRE orthologs outside Holoza and uncover the existence of an animal-like uniporter within chytrid fungi, which enables mt-Ca²⁺ uptake when reconstituted in vivo in the yeast *Saccharomyces cerevisiae*. Our study represents the most comprehensive phylogenomic analysis of the mt-Ca²⁺ uptake system and demonstrates that MCU, EMRE, and MICU formed the core of the ancestral opisthokont uniporter, with major implications for comparative structural and functional studies.

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Mitochondria from several organisms play a key role in regulating intracellular Ca\(^{2+}\) signaling due to their ability to rapidly uptake and buffer cytosolic Ca\(^{2+}\). Mitochondrial Ca\(^{2+}\) (mt-Ca\(^{2+}\)) uptake is mediated by a highly selective channel, the mt-Ca\(^{2+}\)-uniporter, that resides in the inner mitochondrial membrane and is powered by the negative transmembrane potential\(^{2-4}\). However, the functional role of mt-Ca\(^{2+}\) uptake and its pathophysiology has remained largely unknown, as the molecular identity of the mt-Ca\(^{2+}\) uniporter channel was only recently discovered\(^{5-7}\). To this end, comparative genomics analyses based on a few eukaryotic species, in combination with RNAi assays, were instrumental in identifying the first components of the mammalian uniporter, MCU and MICU1 (refs. \(^{8-10}\)). While MCU constitutes the pore-forming and Ca\(^{2+}\)-conducting subunit of the uniporter\(^9\), the EF-hand-containing and Ca\(^{2+}\)-sensitive protein MICU1 has been shown to function as a channel gatekeeper and cooperative activator\(^9\). Importantly, their discovery paved the way to the identification of additional structural and regulatory components of the mammalian uniporter, including a MCU-dominant-negative beta subunit (MCUB)\(^{10}\), three tissue-specific MICU1 paralogs or splice variants (MICU2 (refs. \(^{11,12}\), MICU3 (ref. \(^{13}\)), and MICU1.1 (ref. \(^{14}\))), and an essential MCU regulator (EMRE)\(^{15}\). When co-expressed with MCU, EMRE was shown to be necessary and sufficient to form a functional Ca\(^{2+}\) channel and to reconstitute mammalian-like uniporter activity even in yeast mitochondria\(^{15,16}\), which are otherwise incapable of Ca\(^{2+}\) uptake\(^{16-18}\).

Altogether, these findings have highlighted a complex multi-meric nature for the mammalian uniporter, whose composition, stoichiometry, and regulation need to be fine-tuned to the physiological demands of each cell and tissue. However, the functional and mechanistic role of each uniporter component and the molecular basis of their interdependence are still unclear. Furthermore, although several observations have pointed to an ancient eukaryotic origin of mt-Ca\(^{2+}\) uptake, the identification of several organisms with a different uniporter’s composition in different clades have raised interest in understanding the structural basis of uniporter activity\(^{19-24}\). For example, while MCU and MICU1 showed highly correlated evolutionary histories across 138 sequenced eukaryotic organisms, EMRE apparently lacked any homolog outside the metazoan lineage and was therefore suggested to be an animal-specific innovation\(^{25,26}\). Consistently, Dictyostelium discoideum, an amoebozoan that diverged earlier than the origin of opisthokonts, expresses both MCU and MICU1 orthologs, which form a functional uniporter in the absence of EMRE\(^{16}\). Despite the ancient origin of their interaction and the overall observed high co-evolution between MCU and MICU1, most fungi represented an exception to this rule, with most fungal species encoding homologs of MCU but not of MICU1 (refs. \(^{25,26}\)). The identification of MCU as the only uniporter component in Basidiomycota and filamentous Ascomycota (e.g., Neurospora crassa and Aspergillus fumigatus) has been described as an evolutionary paradox (see ref. \(^{27}\)), but has also been interpreted as indicating that fungal MCUs could be sufficient for mt-Ca\(^{2+}\) uptake and regulated independently of MICU1 (refs. \(^{25,26}\)). Based on the assumption of an orthologous relationship between human and fungal MCUs\(^{25,26,28}\), several independent structural studies of Ascomycota MCUs have been performed to understand the basic principles of uniporter channel assembly and function\(^{21-24}\). However, those organisms had been shown to lack uniporter activity\(^{29,30}\) and their MCU homologs were unable to mediate mt-Ca\(^{2+}\) uptake when heterologously expressed in HeLa or yeast cells\(^{37,24}\). Not surprisingly, significant structural and sequence differences were found between fungal MCUs and their animal counterparts\(^{20-24}\), raising the question of whether fungal MCUs function as classical Ca\(^{2+}\) uniporters at all.

Here, we perform a comprehensive evolutionary analysis of the mt-Ca\(^{2+}\) uniporter and show that a gene duplication at the opisthokont common ancestor of animals and fungi resulted in two distinct MCU paralogous clades, differentially retained in the two groups. When only the animal-like clade is considered, we observe fully consistent co-evolutionary patterns between MCU and MICU (MICU1’s family), across euukaryotes, and of these with EMRE, across opisthokonts. We find that only three early diverging fungi contain both the fungal paralog and the complete animal-like MCU complex (MCU, MICU, and EMRE). Consistently, we find that the heterologous expression of MCU and EMRE from fungal species with an animal-like MCU complex results in the functional reconstitution of Ca\(^{2+}\) uptake in mitochondria of HeLa and yeast cells. Conversely, representatives of the fungal paralogs do not show uniporter activity, suggestive of an alternative function or of the requirement for a yet unidentified regulator. Altogether, our phylogenomic and functional analyses of the mt-Ca\(^{2+}\) uptake system demonstrate that MCU, EMRE, and MICU represent the minimal core components of the ancestral opisthokont uniporter and pinpoint key targets for comparative structural and functional studies. Finally, this study also confirms the importance and power of thorough evolutionary analyses to understand the molecular basis of functional interactions within a protein complex.

### Results

#### The mitochondrial calcium uniporter evolution

We assessed the evolution of each uniporter component across 1156 fully-sequenced eukaryotic genomes (see Supplementary Data 1), using a combination of profile-based sequence searches, protein domain composition assessment, and phylogenetics. As shown in Fig. 1 (Supplementary Fig. 1), the overall taxonomic distributions of MCU and MICU1 homologs were largely congruent with that of previous genomics surveys\(^{25,26}\). We confirmed the presence of MCU in at least some species of the major eukaryotic groups (Unikonts, the SAR clade, Plants, and Euglenozoa), and its absence in all sequenced Apicomplexans, Microsporidia, Trichomonas, and Giardia, and all yeasts in Saccharomycotina and most in Spirochaetes. Hence, mt-Ca\(^{2+}\) uptake appeared to have been lost many times independently during the evolution of eukaryotes. A significant number of these losses correlated with extreme streamlining of mitochondrial metabolism, as most MCU/MICU-lacking lineages encompassed relict forms of anaerobic mitochondria, such as mitosomes (microsporidians, Entamoeba, Giardia, and Cryptosporidium) or hydrogenosomes (Trichomonas)\(^{31}\). Our homology-based results confirmed the above-mentioned anomaly that most fungal genomes, for which our dataset is particularly rich —776 species as compared to 50 in previous studies\(^{25,26}\)—encode for MCU but not MICU or EMRE. Unexpectedly, our analysis uncovered the presence of EMRE outside Holozoa, identifying reliable orthologs in three chytrid fungi—an early diverging zoosporic fungal lineage. Allomyces macrognus, Catenaria anguillulae, and Spicellomyces punctatus. Additional searches in public databases confirmed that EMRE was not present in other sequenced fungi.

To clarify the underlying evolutionary history of the uniporter, we reconstructed and inspected the molecular phylogenies of MCU (Fig. 2a, Supplementary Fig. 2a) and MICU1 (Fig. 2b, Supplementary Fig. 2b) homologs across euukaryotes. We found that the evolution of both MCU and MICU gene families was driven by numerous gene duplications and losses, some of them having occurred in parallel in different lineages, implying an ancient and tight functional relationship. Furthermore, we
showed that evolutionary-independent duplications at the base of several main eukaryotic lineages—vertebrates, streptophytes, oomycetes, kinetoplastids, and ciliates—resulted in the existence of multiple MCU paralogs in each of these clades. As a result, orthology relationships within the MCU gene family are complex and of the type many-to-many. This means, for instance, that human MCU and MCUb are equally distant evolutionarily (co-orthologous) to each non-vertebrate animal clade, with the same pattern, these have all been collapsed to the species level, resulting in 969 terminal nodes shown. The phylogenetic distribution of MCU (red), MICU (green), and EMRE (gray) homologs across 1156 eukaryotic genomes is shown on the NCBI taxonomy tree. Viridiplantae and Rhodophyta (red algae) have been grouped together as Archaeplastida, and alveolates, stramenopiles (Str/es), and rhizarians as the SAR clade. In all cases where data from various strains or species are present with the same pattern, these have all been collapsed to the species level, resulting in 969 terminal nodes shown. The mt-Ca\(^{2+}\) uniporter complex has been completely lost in Apicomplexa within Alveolates, Rhizaria (5 genomes), and red algae (3 genomes), Cryptophytes (5 genomes), Haptophytes (1 genome), and the Entamoeba clade within Amoebozoa. Within fungi (in purple), all major clades that have completely lost MCU homologs are indicated with a darker purple color, namely Onygenales, Saccharomycetales, Pucciniomycotina, Mucoromycotina (Mucor/a), and Microsporidia (Micro/a). The only three early diverging fungal species (A. macrognous-Am, C. anguilulae-C, S. punctatus-Sp) that encode also MICU and EMRE are highlighted with a red arrowhead. The NCBI taxonomy and the presence/absence profile were visualized using the ETE toolkit\(^{41}\). For a version of the profile, which includes the species names, see Supplementary Fig. 1.

Reconstitution of mt-Ca\(^{2+}\) uptake by chytrid MCU and EMRE. The above-mentioned finding of bona-fide EMRE orthologs in these three chytrids (Fig. 1) placed back the origin of an animal-like mt-Ca\(^{2+}\) uptake in the opisthokont ancestor, preceding the diversification of animals and fungi. Consistently, the heterologous expression of MCU from Dictyostelium discoideum, representing an amoebozoan lineage that diverged earlier than the origin of opisthokonts, is alone sufficient to reconstitute mt-Ca\(^{2+}\) uptake in yeast mitochondria, while human (Hs-) MCU only does so in the presence of EMRE\(^{16}\). Similarly, we hypothesized that co-expression of animal MCU and EMRE proteins from chytrids would be necessary and sufficient to reconstitute uniporter activity. The phylogenetic distribution profile (presence/absence) across the MCU complex components revealed a strong co-evolution pattern, when only the true orthologous sequences were considered (Fig. 3a). Strikingly, we detected mt-Ca\(^{2+}\) uptake in yeast strains expressing animal MCUs from either A. macrognous (Am-MCUa) or S. punctatus (Sp-MCU) with their respective EMREs (Am-EMRE1, Sp-EMRE) (Fig. 3b, c, Supplementary Fig. 3). Particularly, in A. macrognous that encodes two MCU proteins, Am-MCUa and Am-MCUb, mt-Ca\(^{2+}\) uptake activity was detected only for the former, and showed different efficiency with its two encoded EMRE variants (Am-EMRE1 higher than Am-EMRE2). In contrast, we did not detect any mt-Ca\(^{2+}\) uptake in yeast strains expressing MCP proteins from A. macrognous (Am-MCU1P) and S. punctatus (Sp-MCU), despite proper expression and localization (Supplementary Fig. 3, Supplementary Fig. 4). Similar results would be expected in other Holozoa despite the inability to detect EMRE by similarity searches. Indeed, the co-expression of MCU from the sea anemone Nematostella vectensis (Nv-MCU) with Hs-EMRE in yeast was able to reconstitute mt-Ca\(^{2+}\) uptake to a similar extent of a strain expressing Hs-MCU and Hs-EMRE (Supplementary Fig. 5). These results, together with the absence of MICU proteins in most fungal lineages, indicate that mt-Ca\(^{2+}\) uptake in fungal mitochondria, if it exists, is not mediated by MCU\(_{s}\), or that a different—yet unknown—regulator is necessary. Instead, animal-like MCUs from chytrid fungi and Holozoa function similarly to the mammalian uniporter, in an EMRE-dependent fashion. Altogether, our evolutionary analyses and experimental results confirmed that MCU–EMRE interaction is conserved, and was already present in the last common ancestor of fungi and animals.

Consistent with our hypothesis that MCU\(_{s}\) do not represent true functional orthologs of Hs-MCU, when comparing MCU sequences across eukaryotes we found that MCU\(_{s}\) lacked key residues conserved in the animal-like MCUs, despite retaining a DXXE motif (Fig. 4a, Supplementary Fig. 6a). Those residues have been previously shown to be important for MCU function and its interaction with EMRE\(^{36}\). Notably, animal and chytrid
EMREs appeared highly divergent (Fig. 4b, Supplementary Fig. 6b), although the MCU interacting domain GXXxA/S/G and the polyaspartate tail necessary for the binding to MICU1 (ref. 36) were fully conserved. Interestingly, the fungal EMRE sequences contained an extra C-terminal domain that was not found in Holozoa, suggesting some degree of specialization. Thus, we hypothesized that Am-MCUa and Sp-MCU would have evolved to interact with EMRE proteins from the same or related species. To this goal, we measured mt-Ca^{2+} uptake in HeLa cells infected with lentiviral particles expressing either Hs-MCU or Am-MCUa and Sp-MCU (Supplementary Fig. 7a). Those proteins were expressed in addition to the endogenous Hs-MCU and Hs-EMRE. We found that mt-Ca^{2+} uptake in HeLa cells expressing either Am-MCUa or Sp-MCU was similar to uninfected control cells in response to histamine, differently from the overexpression of Hs-MCU, which resulted in a gain-of-function phenotype. We then measured mt-Ca^{2+} uptake in HeLa cells where MCU was stably knocked-down (sh-MCU) (Supplementary Fig. 8), with a peak concentration of mt-Ca^{2+} similar to that of cells with MCU knock-down, due to the absence of a fungal EMRE. Furthermore, the expression of Am-MCUa and Sp-MCU in yeast mitochondria was unable to reconstitute mt-Ca^{2+} uptake in the presence of Hs-EMRE (Fig. 4c, Supplementary Fig. 9). Instead, Hs-MCU was functional when co-expressed with either Am-EMRE or Sp-EMRE (Fig. 4d, Supplementary Fig. 9). Altogether, these results suggested that the C-terminal domain of chytrid EMREs was dispensable for a functional interaction with Hs-MCU but necessary to activate animal-like fungal MCUs. Accordingly, we observed that the co-expression of Am-MCUa and Sp-MCU with Am-EMRE and Sp-EMRE lacking the extra C-terminal domain (EMRE-t) was unable to efficiently reconstitute mt-Ca^{2+} uptake (Fig. 4e, Supplementary Fig. 10a). However, while the fusion of Hs-EMRE with this extra chytrid C-terminal domain did not affect the function of Hs-MCU and thus of mt-Ca^{2+} uptake, it was not sufficient to reconstitute mt-Ca^{2+} uptake when co-expressed with Am-MCUa and Sp-MCU (Fig. 4f, Supplementary Fig. 10b). On the one hand, these findings hint at a possible activating role of the extra C-terminal domain of the fungal EMRE for the efficient regulation of animal-like MCUs function. On the other hand, they suggest that functional domains that are conserved between fungal and human EMREs are sufficient to regulate Hs-MCU activity.

**Discussion**

Altogether, our identification of a strong co-evolution pattern between MCU, MICU, and EMRE provides an explanation to an elusive evolutionary paradox: the presence of uniporter homologs in species with MCU but no MICU1 homologs and no detectable mt-Ca^{2+} uptake. We demonstrate that an animal-related MCU complex has been lost early within the evolution of most fungal clades, which retained only fungal-specific paralogous
Fig. 3 Functional reconstitution of mt-Ca^{2+} uptake by fungal MCU and EMRE homologs. a Phylogenetic distribution profile (presence/absence) across MCU complex components. The distribution pattern of MICU (across eukaryotes) and EMRE (in opisthokonts) largely overlaps with that of the animal-like MCU, but not the MCUP. b, c Representative traces and quantification of mt-Ca^{2+} transients in yeast cells expressing human MCU and EMRE (n = 4) or animal-like or fungal-specific MCU orthologs from S. punctatus (n = 3) (b) and A. macrogynus (n = 3) (c) with either their respective EMRE proteins or empty vector (p425) upon glucose-induced calcium (GIC) stimulation in presence of 1 mM CaCl_{2}. All data represent mean ± SEM. P values are indicated in the different panels (b, c *p < 0.001, one-way ANOVA with Dunnnett’s multiple comparisons test). The silhouette images in a representing the different lineages were downloaded from Phylopic (http://phylopic.org/) or wikipedia (https://www.wikipedia.org/), or designed by A.A.P. in Inkscape (https://inkscape.org/). All downloaded images were available for reuse under a Public Domain license and do not require attribution. Source data of b and c are provided as a Source Data file.

MCU proteins (MCUPS). These MCUPS, which are paralogous to Hs-MCU, have been so far wrongly considered functionally equivalent to Hs-MCU and deeply studied for this reason. This is the case, for instance, of the Ascomycota MCUP proteins (e.g., N. crassa) that have been recently used as models for understanding structure and regulation of the human uniporter channel^{20-24}. Consistently, we and others find that fungal MCUPS are unable to reconstitute or rescue mt-Ca^{2+} uptake in yeast or HeLa cells lacking MCU, respectively^{17,24,29}. Similarly, the same fungal MCU sequences used for investigating structure and function of the mammalian uniporter^{21,22,24} failed to mediate mt-Ca^{2+} uptake when expressed in yeast mitochondria or in MCU knockout HeLa cells (Supplementary Fig. 11). Instead, we show that only fungal species having both MCU and EMRE sequences (animal-like MCUs), such as A. macrogynus and S. punctatus, are able to mediate mt-Ca^{2+} uptake. Those observations together with the significant structural and sequence differences found between MCUs and MCUPS^{20} question whether fungal MCUPS function as classical Ca^{2+} uniporters.

The remarkable presence/absence pattern across eukaryotes of MCU and MICU—when only the orthologs of the two protein families are considered—indicates strong co-evolution and functional interdependence. Indeed, MICU1 plays a key role in regulating the gating and activation properties of MCU^{9}. Thus, the evolutionary coupling of uniporter’s gating mechanism with its regulator suggests a dependency on MICU for Ca^{2+} homeostasis and cell physiology. Consistently, across the 1156 species in our analysis, MCU and MICU co-occur in 1144 (~99%), whereas only 12 species have one of the two components. Most such incongruencies are likely to result from assembly, annotation, or identification errors. In fact, we expect MICU to be encoded in all species with a functional MCU complex, with the only likely exception being Onchoercaidae, a family of parasitic nematodes. In our dataset all three species in the clade (Brugia malayi, Onchoerca volvulus, and Loa loa) appear to lack MICU1 homologs in the presence of MCU (Supplementary Fig. 1), raising the possibility that MICU is indeed lost in the clade rather than being undetected. If this preliminary observation is confirmed, this would make these parasites interesting targets to explore further the physiological significance of MICU1.

Our phylogenomic analysis identifies non-metazoan EMRE sequences and demonstrates the ancestrally essential role of EMRE in mt-Ca^{2+} homeostasis. Our results imply that EMRE, previously thought to be an animal-specific innovation^{15,16,27}, formed part of an animal-like machinery in the common ancestor of opisthokonts, and was lost secondarily in the evolution of fungi, together with the other components of the animal-like mt-Ca^{2+} uptake machinery (Fig. 5). These results have major implications for structural and functional studies of the uniporter. Indeed, members of the orthologous MCU complex in basal fungi constitute relevant targets for future research and comparative structural analyses, particularly for identifying key MCU–EMRE
interactions. Our results show that human MCU can function in the presence of both human and chytrid EMREs, whereas chytrid MCUs such as Am-MCUa and Sp-MCU can only reconstitute mt-Ca\(^{2+}\) uptake when co-expressed with their corresponding chytrid EMREs. These findings indicate a tight co-evolution between MCU and EMRE proteins, which we know to functionally and physically interact, and provide the framework to understand the sequence determinants of this interaction. Finally,
our work underscores that accurate phylogenomic analyses can resolve apparent evolutionary rhythms while making explicit functional predictions that can drive future experiments.

Methods

Sequence data and homology searches. The protein sequences encoded in 1156 completely sequenced eukaryotic genomes were retrieved from Ensembl DB v91, and v37 of Ensembl Metazoa, Plants, Fungi, and Protists (see Supplementary Data 1). Only the genome of Catenaria anguillulae PL171 was added from Ensembl Fungi v41. For each of the protein families studied, homologs were selected on the basis of sequence similarity and phylogenetic analysis. HMMER searches were performed using HMMER 3.1b2 (ref.37) and using the Gathering Cut-Off threshold (−cut_ga) when the raw HMM profile from Pfam was used, an e-value threshold of 10\(^{-3}\) otherwise. For all BLAST searches low-complexity regions in the query sequence (default parameter) were filtered out to minimize the number of false positives and an E-value threshold of 10\(^{-5}\) was used. Conserved domains in all retrieved sequences were annotated using the HMM profiles of Pfam release 30.0. For MCU, proteins in our database containing at least one MCU (Pfam: PF04678 [https://pfam.xfam.org/family/PF04678]) domain were detected using an HMMsearch. In total, 1076 protein sequences were selected for subsequent analysis. For MICU, 2105 protein sequences were retrieved from a BLAST search using Hs-MICU1 (UniProt: Q9BPX6 [https://www.uniprot.org/uniprot/Q9BPX6]) as a query. HMBase was used to search for additional domains in the retrieved sequences using all the Pfam domain profiles, and all the sequences with at least one Mito_carr domain (Pfam: PF00153 [https://pfam.xfam.org/family/PF00153]) were classified as the members of the mitochondrial carrier family (scl25a12-Aralar homologs). The Aralar-related sequences (Mito_carr domain containing) were clustered together clearly as a monophyletic clade in a phylogenetic tree, in the exclusion of known MICU sequences, and were excluded. The remaining 651 MICU sequences were re-aligned and a new phylogenetic tree with the same methods was reconstructed. EMRE sequences are characterized by a DDDD (Pfam: PF10161 [https://pfam.xfam.org/family/PF10161]) domain. Their short length and low sequence conservation makes detection strikingly difficult, which explains why in some cases EMRE appears to be missing even from some animal genomes. HMMER searches with the “gathering” (−cut_ga) threshold were performed, and all detected homologs were retrieved and re-aligned, for a new HMM profile to be built and used to search back the genome database in a second iteration. One EMRE sequence was detected in A. macrogynus in the first search, while one more sequence in the same species, as well as in S. punctatus and C. anguillulae, were detected in the second iteration. The detected EMRE sequences in this second iteration are those that were further considered for all analyses. For NCLX (NCKX6), the Na\(_\text{Ca}\text{ex}\) Pfam domain was initially used to retrieve protein sequences. However, this domain characterizes a ubiquitous superfamily of sodium/calcium exchangers that regulate intracellular Ca\(^{2+}\) concentrations in many cell types. Therefore, selection of NCKX6 (NCLX) homologs based on the Na\(_\text{Ca}\text{ex}\) Pfam domain using HMMER returned 4024 hits. To narrow down the number of hits for more accurate alignment and phylogenetic reconstruction, we used the human NCLX sequence as a query (uniprot: Q6H4K2 [https://www.uniprot.org/uniprot/Q6H4K2]) for a blast search, retrieving 2105 sequences for phylogenetic analysis. Using the human members as reference, 1391 sequences across eukaryotes were selected as related to the NCLX clade (NCLX orthologs).

Phylogenetic analysis. Sequences of all the protein families described were aligned, and the alignment was trimmed and used to compute a phylogenetic tree. The selected homologous proteins were aligned with MAFFT v7.394 (ref.38) (E-INS-I for MICU, EMRE, and NCLX families and L-INS-I for the MCU family, based on their multi/single-domain architecture) and a soft trimming was applied, filtering out positions in the alignment with gaps in more than 99% of the sequences using trimAl\(^{(9)}\) (−gt 0.01). IQ-TREE v1.6.8 (ref.40) was used to derive Maximum Likelihood (ML) trees. LG was selected as the base model to test all rate-heterogeneous models using the “-mset LG” parameter (26 protein models, combinations of invariant sites +I, discrete Gamma model with 4 rate categories +G, the FreeRate model +R with 2–10 categories, and empirical AA frequencies estimated by the data). The best-fit models, chosen according to the Bayesian information criterion (BIC), were LG + R10 for the MCU and MICU alignments and LG + F + R10 for NCLX alignment. Branch supports were obtained using the ultrafast bootstrap implemented in the IQ-TREE program with 1000 replicates. The ETE Toolkit\(^{(41)}\) was used for all taxonomic and phylogenetic tree operations and visualization. Sequence alignments were visualized using Jalview (ref.42), and
The Multi-Harmony26 method was used identify patterns of variation across the different protein clades, positions conserved across the animal MCUs and the animal-like fungal MCUs but not in the MUCPs. Raw phylogenetic trees in newick format can be found in Supplementary Data 2.

**Cell lines.** HeLa cells stably expressing a wild-type mitochondrial matrix-targeted GFP-sequitoir (mt-AEQ) were generated as in ref. 18. Mt-AEQ HeLa cells stably overexpressing Hs-MCU, Sp-MCUP, Sp-MCU, Am-MCUP1, Am-MCUa, Am-MCUb, Ma-MCUP, and Nf-MCUP from the plasmid containing the target sgRNA sequences were annealed and cloned into the pLX304 destination vector by site-specific recombination (Life Technologies). All cell lines were grown in high-glucose Dulbecco’s modified Eagle’s medium supplemented with 10% FBS at 37 °C and 5% CO2.

**Yeast strains.** A yeast strain expressing mt-AEQ with Hs-MCU and Hs-EMRE was generated as in refs. 17,18. To generate yeast strains co-expressing mt-AEQ together with different combinations of fungal and human MCU and EMRE orthologues, cDNAs were amplified from genomic DNA transfection reagent (Roche), according to the manufacturer’s instructions. The YPH499 strain was isolated from pX330 (Addgene #42230) by BbsI digestion and cloned upstream of the ubiquitous CAG promoter. Two sets of single-guide RNA (sgRNA) were designed using an online tool CRISPOR (MCU-KO gRNAs: 5′-CAGCGGTTTCATCCGACGGG-3′, MCU-KO gRNAs: 5′-CAGCGGTTTCATCCGACGGG-3′). For each sgRNA, complementary oligonucleotides containing the sgRNA sequences were annealed and cloned into the pX330-Puro-cdc8 vector35. After which, pX330-Puro-cdc8 vector containing sgRNAs were transfected into mt-AEQ HeLa cells using the Lipofectamine 3000 Reagent (Life Technologies), according to the manufacturer’s instructions and selected with 2 µg/ml puromycin for 2 days. Cells were then seeded at a density of 1 cell per well. Gene knockout was confirmed by PCR (MCU-KO Forward1: 5′-GGTGTTAGTGAGTTTACAGGC-3′, MCU-KO Reverse: 5′-GGTGTTCCGTCATCGGTACC-3′) and confirmed by sequencing and western blot using the following antibodies: anti-MCU (Sigma-Aldrich, HPA01648, 1:1000), anti-EMRE (Santa Cruz Biotechnology, sc-86337, 1:1000), anti-V5 (Life Technologies, R96025, 1:5000), anti-AEQ (Merck/Millipore, MAB4405, 1:1000); anti-YME1; anti-PGK1 (Life Technologies, 459250, 1:10,000).

**MCU knockout in HeLa cells.** Mt-AEQ HeLa cells with knockout of MCU were generated by CRISPR targeting35. In brief, a cDNA encoding NLS-Cas9 was isolated from plasmid pX330 (Addgene #42230) by EcoRI digestion and cloned upstream of the ubiquitous CAG promoter. Two sets of single-guide RNA (sgRNA) were designed using an online tool CRISPOR targeting MCU-KO gRNAs: 5′-CAGCGGTTTCATCCGACGGG-3′, MCU-KO gRNAs: 5′-CAGCGGTTTCATCCGACGGG-3′. For each sgRNA, complementary oligonucleotides containing the sgRNA sequences were annealed and cloned into the pX330-Puro-cdc8 vector35. After which, pX330-Puro-cdc8 vector containing sgRNAs were transfected into mt-AEQ HeLa cells using the Lipofectamine 3000 Reagent (Life Technologies), according to the manufacturer’s instructions and selected with 2 µg/ml puromycin for 2 days. Cells were then seeded at a density of 1 cell per well. Gene knockout was confirmed by PCR for MCU-KO1 Forward1: 5′-GGTGTTAGTGAGTTTACAGGC-3′, MCU-KO Reverse: 5′-GGTGTTCCGTCATCGGTACC-3′) and confirmed by sequencing and western blot using the following antibodies: anti-MCU (Sigma-Aldrich, HPA01648, 1:1000), anti-EMRE (Santa Cruz Biotechnology, sc-86337, 1:1000), anti-V5 (Life Technologies, R96025, 1:5000), anti-AEQ (Merck/Millipore, MAB4405, 1:1000); anti-YME1; anti-PGK1 (Life Technologies, 459250, 1:10,000).

**Isolation of crude mitochondria from HeLa cells.** Crude mitochondria were isolated from HeLa cells as described17. Briefly, HeLa cells were grown to confluence, rinsed with PBS and resuspended in ice-cold isolation buffer (IB: 220 mM sucrose, 25 mM HEPES-KOH pH 7.4, 1 mM EDTA-KOH pH 7.4, protease inhibitors). Cells were permeabilized by nitrogen cavitation at 600 psi for a 30 s cooling interval in between to break cell wall and plasma membrane. After the first centrifugation step at 1000 x g for 10 min at 4 °C, the supernatant was further centrifuged at 20,000 x g for 10 min at 4 °C to obtain the mitochondrial fraction (pellet). The supernatant (cytosolic fraction) was precipitated with trichloroacetic acid at −20 °C for 1 h, washed once with cold acetone, and centrifuged at 20,000 x g for 10 min at 4 °C. Both cytosolic and mitochondrial fractions were analyzed with immunoblotting according to the standard procedures using the following antibodies: anti-MCU (Sigma-Aldrich, HPA01648, 1:1000); anti-EMRE (Santa Cruz Biotechnology, sc-86337, 1:1000); anti-V5 (Life Technologies, R96025, 1:5000); anti-AEQ (Merck/Millipore, MAB4405, 1:1000); anti-YME1; anti-PGK1 (Life Technologies, 459250, 1:10,000).

**Mitochondrial calcium uptake in yeast and HeLa cells.** In vivo analyses of mitochondrial Ca2+ uptake in intact yeast and HeLa cells were performed using an aequorin-based measurement46. Aequorin is widely used as a genetically encoded Ca2+ indicator (GECI) for measurements of mt-Ca2+ kinetics because it offers several advantages. Besides being targeted with precision to the organelle, it functions over a wide range of Ca2+ concentrations, and shows low buffering capacity, despite achieving a limited spatial resolution compared to fluorescent GECIs48. Briefly, yeast cells were collected at an OD ~0.8, washed three times with milliQ water, and starved for 1.5 h at room temperature in a nutrient-free buffer (NFB, 100 mM Tris, pH 6.5 (1 x 10⁸ cells per ml). Afterwards, cells were collected at 3500 t.p.m. for 5 min and resuspended in NFB to a higher density (25 x 10⁶ cells per ml) in the presence of 50 nM native coelenterazine (Abcam, ab145165) to reconstitute the photoprotein aequorin. After 30 min in the dark at room temperature, 0.5 x 10⁶ cells per well were plated into a white 96-well plate of white 96-well plate in growth medium overnight. Afterwards, mt-AEQ was reconstituted with 2 µM native coelenterazine for 2 h at 37 °C and Mt-Ca2+-dependent light kinetics were recorded upon illumination with 1 mM CaCl₂ and 100 mM glucose, at 469 nm every 0.5 s interval in a MicroBeta2 LumijET Microplate Counter. At the end of each experiment, cells were lysed with 1 mM digitonin for 5 min at 37 °C and any residual aequorin counts were collected upon the addition of CaCl₂ to a final concentration of 140 mM. For mt-Ca2+ uptake measurement in mt-AEQ HeLa cells, those were seeded at 25,000 cells/well of white 96-well plates in growth medium overnight. Afterwards, mt-AEQ was reconstituted with 2 µM native coelenterazine for 2 h at 37 °C and Mt-Ca2+-dependent light kinetics were recorded upon illumination with 100 µM histamine stimulation at 469 nm every 0.1 s interval. At the end of each experiment, cells were lysed with a solution containing 0.5% Triton X-100 and 10 mM CaCl₂ to record the released residual aequorin counts.

**Quantification of calcium transients.** Quantification of mt-Ca2+ concentration was performed using a MATLAB software as in ref. 18. A description of the Ca2+-dependent luminescence signal was modeled by the cubic spline function:
and data analysis. Normal distribution was tested by Shapiro–Wilk normality test. Statistical tests between multiple datasets and conditions were carried out using one-way analysis of variance followed by Dunnett’s multiple comparison tests. Statistical analyses were performed using GraphPad Prism (GraphPad Software, version 8).

**Reporting summary** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

The data that support the findings of this study are available from the corresponding authors upon reasonable request. All genome data and predicted peptide sets are publicly available and were downloaded from Ensembl-v91 (https://wwwensembl.org/), and Ensembfungi (fungi.ensembl.org/), metazoa (metazoas.ensembl.org/), plants (plants.ensembl.org/) and protists (protists.ensembl.org/index.html). The raw HMM profiles of the different protein families were downloaded from Pfam release 30.0 (https://pfam.xfam.org/). Hs-MICU1 (MICU1_HUMAN) and NCLX v37. The raw HMM protein

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**Author contributions**

A.A.P., T.G., and F.P. conceived the project, designed the experiments, and wrote the manuscript. V.G. performed most of the experiments with help from J.W. and A.C.-S. A.A.P. performed computational analysis. T.G. and F.P. supervised work and acquired funding.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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