



HHS Public Access

Author manuscript

Biochem J. Author manuscript; available in PMC 2023 September 08.

Published in final edited form as:

Biochem J. 2021 September 17; 478(17): 3205–3220. doi:10.1042/BCJ20210342.

Evolutionary rate covariation identifies SLC30A9 (ZnT9) as a mitochondrial zinc transporter

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Abstract

Recent advances in genome sequencing have led to the identification of new ion and metabolite transporters, many of which have not been characterized. Due to the variety of subcellular localizations, cargo and transport mechanisms, such characterization is a daunting task, and predictive approaches focused on the functional context of transporters are very much needed. Here we present a case for identifying a transporter localization using evolutionary rate covariation (ERC), a computational approach based on pairwise correlations of amino acid sequence evolutionary rates across the mammalian phylogeny. As a case study, we find that poorly characterized transporter SLC30A9 (ZnT9) coevolves with several components of the mitochondrial oxidative phosphorylation chain, suggesting mitochondrial localization. We confirmed this computational finding experimentally using recombinant human SLC30A9. SLC30A9 loss caused zinc mishandling in the mitochondria, suggesting that under normal

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Competing Interests

The authors declare that there are no competing interests associated with the manuscript.

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Kirill Kiselyov: Conceptualization, Resources, Data curation, Formal analysis, Supervision, Funding acquisition, Validation, Investigation, Visualization, Methodology, Writing — original draft, Project administration, Writing — review and editing.

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conditions it acts as a zinc exporter. We therefore propose that ERC can be used to predict the functional context of novel transporters and other poorly characterized proteins.

Introduction

Localization in a specific organelle or subset of organelles defines the functional context for many molecules, especially ion and metabolite transporters. For example, there are ion channels and transporters that localize to the plasma membrane or endo/sarcoplasmic reticulum, and even though they might conduct the same ions, they clearly have a different impact on cellular function [1–3]. Similarly, a large cohort of zinc transporters, responsible for pumping ionic zinc into distinct organelles or in or out of cells, subserve distinct functions, including synaptic modulation [4] and co-secretion with milk, in the latter case providing an essential dietary supplement and shaping mammary gland development [5]. The recent advances in genomic sequencing have resulted in identification of many putative transporters. While some aspects of their function can be inferred based on sequence homology or structural similarity to previously characterized molecules, important details, such as subcellular localization, are usually beyond the reach of these approaches. The main objective of the present studies is to evaluate whether a computational approach based on molecular evolution can accurately predict cellular localization of poorly characterized cellular proteins. Here, this idea was tested using zinc transporters as a model.

Zinc is an essential component of cellular function, as a key cofactor in enzymatic reactions, transcription and synaptic transmission [6]. Moreover, it has been estimated that ~10% of human proteome binds zinc [7]. Zinc deficit is a factor in growth delays, hair and skin pathologies and behavioral defects [8]. On the other hand, excess zinc promotes the production of reactive oxygen species and many forms of cell damage [9–12]. Because of zinc's duality, cellular zinc concentrations are tightly regulated by a system of transporters and metal-binding proteins. In particular, zinc entry into the cytoplasm is managed by the SLC39A (Zip) family of transporters, while its expulsion or sequestration into organelles is driven by the SLC30A family. Metallothioneins, in addition to other zinc-binding proteins, provide an important intracellular buffer for the metal, leading to extremely low (sub nanomolar), or perhaps non-existent, levels of free zinc within the cytoplasm [13]. The zinc transporter families contain two dozen of structurally related members, which were discovered as a result of advances in sequencing technology and gene annotation [6]. However, many of these molecules remain under-characterized at the level of subcellular localization or transport mechanisms. The full extent of annotations supported by several lines of experimental evidence indicates that, for example, SLC30A1 and SLC39A1 function in the plasma membrane [14,15], SLC30A3 functions in synaptic vesicles [16] and SLC30A2 and SLC30A4 function in lysosomes and related organelles [17,18]. For many of the remaining SLC30A and SLC39A transporters, however, functional annotations, including cellular localizations, are fragmentary. Importantly, some organelles appear to contain several zinc transporters with overlapping functions, and the reason for this is unclear [6,19].

At the same time, some key aspects of organellar zinc transport remain poorly understood and/or lack an assigned transporter. This is especially the case for mitochondrial zinc, which, as a cofactor of several mitochondrial enzymes, is a critical component of mitochondrial physiology [20,21]. In parallel to full-cell consequences of zinc dysregulation, mitochondrial zinc overload is toxic and has been linked to traumatic brain injury and stroke [12]. The mechanisms of mitochondrial zinc regulation are poorly understood and neither SLC30A nor SLC39A members have been directly shown to localize in the mitochondria. Moreover, while the mitochondrial calcium uniporter MCU was proposed to be responsible for zinc uptake into the mitochondria from the cytoplasm [21,22], the pathway that dissipates the mitochondrial zinc has not been described.

These considerations illustrate a need for a predictive approach to match zinc transporters to specific organelles and focus experimental hypotheses to a reasonable set of targeted studies. We propose Evolutionary Rate Covariation (ERC) as the link between functional genomic information about zinc transporters and their associated organelles. ERC is based on the central hypothesis that proteins with shared functional context will share evolutionary pressure and thus evolve at similar rates, even as their rates change over time and between species [23,24]. This evolutionary rate covariation occurs in physically interacting proteins that coevolve as well as functionally related proteins such as gene regulatory elements, membrane traffic adaptors, and transporters. It is also important to note that many, but not all, recognized functional relationships between genes are reflected by high ERC correlations. Because of its predictive capability in some pathways, ERC has been used to identify the functions of uncharacterized genes, including novel DNA damage-induced apoptosis suppressor, sex peptide network components, regulators of cell adhesion, mitochondrial components and ion and aminoacid transporters [23,25–30]. ERC is calculated as correlation between evolutionary rates throughout a phylogeny. In fact, the work presented here will focus on values calculated across 33 mammal species. Genes are inferred to be functionally related if they have a high degree of correlation in evolutionary rates, or, in other words, if the genes in question have similar evolutionary rates across species.

We present ERC as a powerful tool for *de novo* prediction of molecular function with characterization of the zinc transporter SLC30A family, specifically SLC30A9, its most poorly characterized member. Although there are previously reported links to zinc transport [31,32], this transporter had previously not been linked to an organellar function in mammalian cells. We show here that ERC analysis suggests a strong and specific signal for SLC30A9 coevolution with several components of the mitochondrial oxidative phosphorylation chain including complex I, and the mitochondrial H⁺ driven ATP synthase (complex V [33,34]). The ERC signal between SLC30A9 and mitochondrial complexes was significantly higher than the signal between SLC30A9 and the vacuolar/vesicular/lysosomal H⁺ pump. We also find that recombinant SLC30A9 co-localizes with the mitochondrial protein marker TOM20, strongly suggesting mitochondrial localization. This is consistent with prior evidence contained in MitoCarta, which integrates multiple lines of evidence for mitochondrial localization and function of proteins, including APEX_matrix, targetP signal, yeast mitochondrial homolog, mitochondrial protein domain, induction, and MS/MS [35,36]. Furthermore, SLC30A9 knockdown in HeLa cells using siRNA suppresses the

dissipation of mitochondrial zinc after zinc overload. We therefore propose that SLC30A9 is a mitochondrial zinc exporter and thus possesses a unique functional profile unique for members of the SLC30A family. Of interest, further inquiry of SLC30A sequence revealed that SLC30A9 has followed a unique evolutionary trajectory — it is deeply and highly conserved from mammals through archaea and proteobacteria, while other SLC30As are likely a result of more recent gene duplication events. These findings both illuminate the function of SLC30A9 and distinguish it as a unique molecule among SLC30A family members.

Materials and methods

Discovery and categorization of SLC30 sequences

Mammalian and other multi-cellular eukaryotic amino acid sequences of SLC30 family members were obtained using Pubmed search and aligned using CLUSTALW [37]. The ExxHxxxD...(IVL)(IVL)xED and DxxHxxxD... HxxxD templates were used as wildcard searches of single-celled eukaryotic and prokaryotic sequences by means of BLASTp. The resulting sequences were verified for the presence of six full or partial transmembrane domains using TMHMM server.

Phylogenetic analysis

SLC30 family proteins and related homologs collected in the previous step were analyzed in terms of their divergence and phylogeny. Amino acid sequences were aligned using CLUSTALW [38]. The resulting alignment was used in PhyML 3.0 to infer their phylogenetic relationships and perform branch support analysis using 100 bootstrap replicates [39]. The 'LG' amino acid substitution model was used and rate heterogeneity was modeled using a class of invariable sites with freely estimated size and a Gamma rate shape parameter discretized as four rate categories. The tree figure in Figure 2 was made using Interactive Tree of Life (iTOL) [40,41].

Evolutionary rate covariation

The calculation of ERC values was performed as in previous publications [23,24]. In this case, orthologous gene sequences from 33 mammalian species were obtained from the 100-way alignment at the University of California Santa Cruz Genome Browser [42]. The species chosen were: *Homo sapiens* (human), *Pongo pygmaeus abelii* (orang-utan), *Macaca mulatta* (rhesus macaque), *Callithrix jacchus* (marmoset), *Tarsius syrichta* (tarsier), *Microcebus murinus* (mouse lemur), *Otolemur garnettii* (bushbaby), *Tupaia belangeri* (tree shrew), *Cavia porcellus* (guinea pig), *Dipodomys ordii* (kangaroo rat), *Mus musculus* (mouse), *Rattus norvegicus* (rat), *Spermophilus tridecemlineatus* (squirrel), *Oryctolagus cuniculus* (rabbit), *Ochotona princeps* (pika), *Vicugna pacos* (alpaca), *Sorex araneus* (shrew), *Bos taurus* (cow), *Tursiops truncatus* (dolphin), *Pteropus vampyrus* (megabat), *Myotis lucifugus* (micro-bat), *Erinaceus europaeus* (hedgehog), *Equus caballus* (horse), *Canis lupus familiaris* (dog), *Felis catus* (cat), *Choloepus hoffmanni* (sloth), *Echinops telfairi* (tenrec), *Loxodonta africana* (elephant), *Procavia capensis* (rock hyrax), *Dasyurus novemcinctus* (armadillo), *Monodelphis domestica* (opossum), *Macropus eugenii* (wallaby), and *Ornithorhynchus anatinus* (platypus). Those 17 486 coding sequence alignments were

used to calculate gene-specific branch lengths over the mammalian species tree topology using codeml of the PAML package [43]. For each orthologous gene group/tree, the branch lengths were normalized into relative evolutionary rates through a projection operator [44]. Those relative rates were then used to calculate the Pearson correlation coefficient, the ERC value, between each pair of genes.

Enrichment analysis of organellar proteomes

Pairwise ERC values between SLC30 genes and all other genes were extracted from the ERC website: https://csb.pitt.edu/erc_analysis/. Values for gene subsets of interest were used to generate visualizations and calculate enrichment statistics. Shown in Figure 3A are ERC values using proteomes included in Supplementary File S1 represent key elements of mitochondrial and lysosomal proteomes that would interact with zinc transporters. Enrichment statistics were calculated using two sets of annotations to map genes to functions. First, search terms were used (shown in Figure 3B) to search gene descriptors for each gene taken from the UCSC Genome Browser gene track. If a descriptor contained the term, the gene was mapped to that search term annotation. Also included were MitoCarta annotations [35,36] describing mitochondrial genes. After extracting terms, enrichment was calculated in two ways. First, a one-tailed Fisher's exact test was performed to detect an increased proportion of annotated genes with ERC values above 0.3. This test indicates that a gene being included in a particular annotation and the gene having a high ERC value are related. Second, a Wilcoxon rank-sum test was performed to look for shifts in distributions of ERC values for genes in a particular annotation compared with genes not in that annotation. AUC values, which range from 0 to 1, were calculated directly from the Wilcoxon rank-sum W statistic as $W/(\text{number of annotated genes} \times \text{number of unannotated genes})$. Small AUC values indicate that annotated genes have *lower* ERC values than unannotated genes and large AUC values indicate that annotated genes have *higher* ERC values than unannotated genes.

Clustering

Clusters depicted in Figure 4A were constructed using igraph [45]. Genes included were selected from genes listed in Supplementary File S2 that represent mitochondrial complexes and ATP transporters. Lines were drawn to connect genes with ERC values greater than or equal to 0.3 and singleton genes that did not connect to other genes were removed. The optimal clustering algorithm implemented in igraph was used to define clusters.

Protein expression and widefield microscopy

YFP-tagged C- and N-terminal in-frame fusions of human SLC30A9 sequence (NP_006336.3) were synthesized by Vectorbuilder (Chicago, IL, U.S.A.) and transfected in HeLa cells using Lipofectamine 3000 (ThermoFisher, Waltham, MA, U.S.A.). Before transfection, the cells were grown in plastic culture dishes in DMEM supplemented with 10% FBS, at 37°C, 5% CO₂, in a humidified incubator and plated on glass coverslips at ~70% confluency. The cells were fixed in 4% formaldehyde (in phosphate-based solution, 5 min incubation) 16–24 h post-transfection, permeabilized with 0.1% Triton X100 in phosphate-based solution, 5 min incubation, transferred to blocking buffer (1% bovine serum albumin and 1% goat serum in phosphate-based solution, 1–24 h) and then treated

with primary and secondary (fluorescent) antibodies. TOM20 antibodies (polyclonal, cat number PA5–52843) and Alexa 568-tagged secondary antibodies were from ThermoFisher. The imaging was performed using Nikon SP-1 microscope and analyzed using FIJI [46].

Zinc imaging and qPCR

HeLa cells cultured as above and grown in 12-well culture plates or 35 mm Mattek dishes (MatTek Corp) were transfected with SLC30A9 esiRNA (Sigma, St Louis, MO, U.S.A.) and used 16–48 h post-transfection. qPCR analysis was performed as before [47], and the data were analyzed using the DDCT method. *ACTB* was used as a housekeeping gene; primers were optimized as described before [47]. For zinc-loading experiments, the cells were exposed to 100–300 μM ZnCl_2 in DMEM/FBS for 16 h. Next, the cells were washed, loaded with 5 mM Rhod-2, am (ThermoFisher) in a zinc-containing HEPES-based salt solution (HBSS, in mM: 140 NaCl, 5 KCl, 1 MgCl_2 , 1 CaCl_2 , 10 HEPES (pH 7.4) and 1 g/l glucose, supplemented with 100–300 μM ZnCl_2) and washed again. The dishes were inserted into a closed, thermo-controlled (37°C) stage top incubator (Tokai Hit Co.) above the motorized stage of an inverted Nikon TiE fluorescent microscope equipped with a 60 \times optic (Nikon, CFI Plan Fluor, NA 1.4), a diode-pumped light engine (SPECTRA X, Lumencor). Emissions were detected using an ORCA-Flash 4.0 sCMOS camera (Hamamatsu) and excitation and emission filters were from Chroma. Zinc removal was performed by manual aspiration and gravity-fed application of nominally zinc-free HBSS. The recorded image stacks were analyzed using FIJI. Specifically, discrete regions of interest containing clearly identifiable mitochondria were outlined by hand and the dynamics of average fluorescence intensity in these images was analyzed and compared. Data are presented as the mean \pm standard error of the mean.

Results

SLC30A9 phylogeny

We sought to investigate SLC30A9 function by analyzing its evolution throughout the tree of life. Although SLC30A9 variants have been previously identified in some species, including rat, mouse, zebrafish, and fruit fly (Pubmed search), we performed a more rigorous, targeted search. Due to heterogeneity of the SLC30A/SLC39A superfamily, we began by establishing defining features of SLC30A9 and other SLC30As. Zinc permeation by SLC30A is coordinated by aspartate and histidine residues contributed by the HxxxD motif (flanked by D at position 39 of human SLC30A1, forming DxxHxxxD signature) and by the HxxxD motif located downstream of the first motif (at position 250 of human SLC30A1) (Figure 1A; full alignment in Supplementary Figure S1). Previously published crystal structures and homology modeling show that these motifs are located in the transmembrane domains 2 and 5 of human SLC30As [48–50]. Amino acid alignment of human and other organisms' SLC30A9 with other SLC30As, coupled with homology modeling, reveals a distinct and conserved arrangement (Figure 1A and Supplementary Figure S1). In human SLC30A9, the HxxxD sequence is flanked by a glutamate residue (ExxHxxxD at position 268 of human SLC30A9) and the downstream sequence is ILLED.

These substitutions provide a unique signature that identifies SLC30A9 and distinguishes it from other SLC30A family members. The signatures are surprisingly well conserved: homologous sequences containing these substitutions and transmembrane domain arrangements reminiscent of human SLC30A9, are found throughout the tree of life (Figure 1A and Supplementary Figure S1). Critically, a detailed phylogenetic analysis of SLC30A9 candidates and other members of the SLC30A family identifies SLC30A9 as a distinct phylogenetic entity, components of which are more related to each other across the tree of life, than to other SLC30A members.

The substitutions specific to SLC30A9 appear to be exceptionally well-conserved as we can detect SLC30A9 homologous sequences containing the invariant ExxHxxxD...(I/V/L)(I/V/L)xED in every clade except for the yeast and other fungi (Table 1, Figures 1C, 2, see additional trees in Supplementary Figures S2, S3). This includes archaea, proteobacteria, cyanobacteria, firmicutes, and actinobacteria, as well as every animal species tested. Organisms on which SLC30A9 is present typically contain a single SLC30A9 homolog, with the exception of *Nostoc punctiforme* (2 copies), *Actinobacteria bacterium* (2 copies), and *Micrarchaeota archaeon* (3 copies). The SLC30A superfamily increases in complexity in eukaryotes, with ten members (including SLC30A9) in mammals, six members in *Ciona intestinalis* and *Drosophila melanogaster*, and 4 members in *Hydra vulgaris*. However, of important note, the numerous duplications of SLC30As are not paralleled by increasing numbers of SLC30A9 homologs (Figures 1C, 2). This distinction points toward a unique evolutionary trajectory for SLC30A9 compared with other SLC30s, potentially because of a unique molecular function for this transporter.

As further evidence of SLC30A9's unique evolution, the human SLC30A9 is more closely related to bacterial SLC30A9 than to other human SLC30As based both on sequence identity and on clustering through sequencebased phylogenetic tree construction (Figures 1D, 2). The SLC30A9 sequences appear to be an ancient clade and have remained a single-copy orthologous group during eukaryotic and bacterial evolution. This monophyletic relationship of the ancient SLC30A9 subtree is supported by strong branch support (100% bootstrap support). The phylogenetic tree also shows a clade of SLC30A9 sequences from archaea that may not be monophyletic with the rest of the SLC30A9 clade, making their membership in this family uncertain. Together these findings show that SLC30A9 is an ancient, deeply conserved protein that likely arose via ancient duplication followed by neofunctionalization, meaning that it should be considered functionally distinct from the rest of the eukaryotic SLC30A family. As such, it appears that the ancestor of the SLC30A family was present early in prokaryote evolution. Indeed, SLC30A9 has remained relatively static, not undergoing further duplications. In contrast, the remaining SLC30A family members have gone through multiple gene duplications and deletions, leading to the SLC30A1–8 and 10 paralogs in eukaryotes.

Yeast and other fungi do not appear to have SLC30A9. MMT1/2 yeast transporters, which have been suggested to be the yeast analog of SLC30A9, do not have the ExxHxxxD...(I/V/L)(I/V/L)ED substitution, and thus likely have a different function. Nonetheless, there appears to be mutual exclusivity between SLC30A9 and MMT1/2 expression, as organisms containing SLC30A9 do not seem to have MMT1/2 (Table 1, Figure 2), suggesting that

MMT1/2 may replace SLC30A9 in those species, although our sequence identity and phylogeny construction analyses suggest that MMT1/2 would not have originated from substitutions in SLC30A9. Although we did not further pursue this avenue of inquiry, mechanisms by which yeast species compensated for the loss of such a deeply conserved protein is of great interest. MMT1/2 are found in the yeast mitochondria. This, together with the correlation between the presence of SLC30A9 and MCU in many species (Table 1) strongly suggested that SLC30A9 would be present in the mitochondria.

SLC30A9 coevolution with the mitochondrial components

Since we found that SLC30A9 is an evolutionarily distinct and ancient protein, we next applied another evolutionary method to generate a hypothesis about its function and localization. To gain an insight into the cellular localization of SLC30A9 we performed an ERC analysis between SLC30A9 and a battery of other organellar transporters. Because some SLC30A transporters have been shown or proposed to use a proton gradient as a driving force for zinc transport [49–51], our rationale for this evolutionary approach was as follows: if SLC30A9 uses an ionic gradient to pump zinc, then its evolution could be influenced by the same evolutionary pressures acting on the transporter that establishes the ionic gradient. Hence, the two transporters will have correlated evolutionary rates, which will be uncovered by ERC. We can, therefore, use well established ion transporters, whose localization is known, as baits to search for SLC30As that have coevolved with them, and, as such, probably located within the same organelle.

The analysis was performed using a publicly available ERC portal (https://csb.pitt.edu/erc_analysis/) [52]. This approach utilizes gene-specific evolutionary rates from 33 mammalian species and their ancestral branches calculated for more than 19 000 protein-coding genes. A gene pair's ERC value is calculated as the correlation coefficient between their rates in those species. Analysis of SLC30A9 focused on several organellar transporters including the vacuolar/vesicular/lysosomal H⁺ pump (multimolecular complex comprising structurally unrelated proteins that are coded by the ATP6 family of genes [53,54]), the endolysosomal ClC transporters [55,56], and the components of the mitochondrial oxidative phosphorylation chain [57], which either establish or utilize the mitochondrial H⁺ gradient (see Materials and Methods and Supplementary Figure S2 for the list). We calculated ERC values between SLC30A9 and each of the genes annotated to those batteries of transporters. MCU coevolution with MICU1 [58,59] was used as a positive control for coevolution of functionally linked proteins (R value 0.584, *P* = 0.002).

SLC30A3 and SLC30A4, known to localize in the acidic organelles containing high levels of ATP6, namely synaptic vesicles [16,60] and lysosomes [18], respectively, were used as controls for specificity of the observed effects. Both of these transporters were shown, or proposed, to utilize H⁺ gradient established by ATP6 pumps in order to move zinc. As predicted in this control analysis, SLC30A3 and SLC30A4 show several significant ERC values with the ATP6 components (Figure 3A, top, red points; see original plots of ERC values in Supplementary Figure S4), which is consistent with their cellular localization and H⁺ requirements for transport. In contrast, their ERC scores with the mitochondrial components (blue points) were lower, and, in both cases, ERC was skewed towards the

vacuolar/vesicular/lysosomal H⁺ pump relative to the mitochondrial signal (Figure 3A, bottom).

In contrast, we found that SLC30A9 shows a strong coevolutionary signal with several components of the mitochondrial oxidative phosphorylation chain, which is a very unique pattern when compared with other members of the SLC30A family (Figure 3A, top). Specifically, it has significantly correlated rates with three individual mitochondrial components, belonging to mitochondrial ATP synthase and complex I. In addition, the full distribution of SLC30A9's scores with the mitochondrial complex encoding genes was much higher than its scores with lysosomal components (Figure 3A, bottom).

We next examined the full set of scores for each SLC30A in mammalian genomes to learn with which functions high-scoring genes were associated with it. This study also shows that SLC30A9 is uniquely enriched for high ERC values with genes encoding proteins that are associated with the mitochondria (Figure 3B). First, we searched the human RefSeq gene descriptions of the high-scoring genes (ERC value >0.3) for specific search terms related to calcium, zinc, lysosomes, and mitochondria [61]. SLC30A9's high-scoring genes were strongly associated with the 'mito' search term and with genes annotated as mitochondrial by MitoCarta 2.0 (Figure 3B, upper) [35,36]. Similarly, we studied the entire distribution of genes as ranked by ERC values with SLC30As. That distribution for SLC30A9 was significantly enriched at high ERC scores for genes with mitochondrial annotations (RefSeq gene description and MitoCarta status) (Figure 3B, lower). In fact, out of all SLC30As, only SLC30A9's ERC scores showed such associations with mitochondrial annotations.

We also examined the networks of connections drawn based on ERC scores between SLC30As and components of mitochondrial oxidative phosphorylation complexes (Figure 4A, upper) and organellar ion transporters, including the vesicular Cl⁻/proton transporters (Figure 4A, lower). See also Supplementary Figure S5 for more detail. It is clear that SLC30A9 uniquely co-evolves with many mitochondrial components, involving several connections to them, when compared with other SLC30As, as well as other organellar markers. The models in Figure 4B summarize the findings. Based on the totality of these evolutionary studies, we propose that SLC30A9 localizes to the mitochondria. We set out next to test this hypothesis.

SLC30A9 localization and effect on mitochondrial zinc

Although MitoCarta predicts the mitochondrial localization of SLC30A9, it has not been directly demonstrated. We did not find commercially available antibodies that produce a convincing organellar stain. Therefore, we transfected HeLa cells with recombinant cDNA constructs coding for human SLC30A9 with in-frame C- or N-terminal YFP fusions. Figure 5A,B shows that both constructs produced a reticular stain overlapping with the mitochondrial marker TOM20 [62]. The stretch of positively charged amino acids in the extreme N-terminus of human SLC30A9 is indicative of mitochondrial localization signal, which is supported by Mitoprot II prediction [63]. Deleting this sequence (MLPGLAAAAHRCSWSSLCRLRLRCR, aminoacids 1–26) resulted in SLC30A9 mislocalization and significant cell toxicity following transfection, indicating that the mitochondrial localization of SLC30A9 is essential (Figure 5C). Based on this evidence,

we conclude that SLC30A9 localizes in the mitochondria. This is the first direct evidence of SLC30A9 localization to this or any other organelle.

siRNA-driven SLC30A9 deletion in HeLa cells produced significant changes in mitochondrial zinc handling. Mitochondrial zinc content was analyzed using fluorescent divalent-cation sensitive dye Rhod-2,am, which was loaded into the cells for 15 min, followed by a washout. Fluorescence intensity was analyzed using time-lapse fluorescent imaging; the intensity was measured in the regions of interest which were drawn manually. In zinc-loaded cells, the pattern of the fluorescent signals seemed to conform to the mitochondrial shape. When exposed to 300 μ M zinc overnight, SLC30A9 siRNA-transfected cells (knockdown confirmed by qPCR, Figure 6A) accumulated significantly more zinc, which is evident from statistically higher fluorescence in these cells, compared with control-transfected cells (Figure 6B). Next, we measured the dynamics of the mitochondrial zinc dissipation by returning zinc-exposed cells to a nominally zinc-free medium. Under such conditions, SLC30A9 siRNA-transfected cells lost mitochondria fluorescence signal at a significantly slower rate, indicating slower dissipation of zinc (Figure 6C). These data suggest that SLC30A9 is a functional mitochondrial zinc transporter, which, under normal conditions, is responsible for zinc extrusion from the mitochondria.

Discussion

In the course of the present studies, we used ERC, a bioinformatics assay, to predict the localization of a poorly understood ion transporter. While ERC has been described and tested before [23,25–30], our approach utilized a novel idea that tracking evolutionary histories of functionally linked proteins may inform their functional context, such as localization. Therefore, using this approach we were able to predict protein localization in the absence of obvious clearly defined functional domains. We suggest that this approach can have broad applications in diverse biological fields.

Little is known about the SLC30A9 function, although it has recently been linked to Covid-19 because of strong co-purification of SLC30A9 with SARS-CoV-2 proteins (Supplementary Data in [64]). Furthermore, SLC30A9 has recently been identified as a master regulator of Parkinson's pathogenesis [65]. Given the central role of mitochondria in Parkinson's pathology [66] and the emerging role of zinc in this process [67], the evidence of SLC30A9 in the mitochondria suggests a novel contributing factor to Parkinson's disease.

While SLC30A9 is considered to be a member of the zinc family of transporters, the evidence of its involvement in zinc handling is scarce. Interestingly, polymorphisms in SLC30A9 in African populations have been linked to environmental zinc availability [32], and dysregulation of zinc handling has been detected in fibroblasts from human patients a novel autosomal recessive cerebro-renal syndrome, which has been linked to mutations in *SLC30A9* [31]. Our data suggest that under normal conditions, SLC30A9 is a mitochondrial zinc exporter. While this seems to be counteractive to the known function of other members of the SLC30A family, namely, moving zinc *into* organelles, it is important to note that the mitochondrial proton gradient operates in the opposite direction in mitochondria, when

compared with other membranous organelles such as lysosomes and the Golgi apparatus. Given that some zinc transporters have been shown or proposed to be proton-driven, the 'reversed' mitochondrial proton gradient may explain the outward direction of SLC30A9-driven cation transport detected in our system and attributed to SLC30A9 activity.

Mitochondrial zinc transport mechanisms are poorly understood. Several components of the mitochondrial respiratory chain require zinc [20,21]. Furthermore, proteostasis, protein insertion, enzymatic activities and other functions of the mitochondria are regulated by zinc [68–71]. With this in mind, the SLC30A9 coevolution with the mitochondrial oxidative phosphorylation chain components is informative. However, excess zinc has adverse impacts on the mitochondrial respiratory chain and has been implicated in several pathologies including Alzheimer's disease and stroke [11,72–76]. Importantly, a mechanism of zinc efflux from the mitochondria had, heretofore, not been identified. Ascribing this role to SLC30A9 is likely to have far-reaching implications in a more complete understanding of zinc management by the mitochondria.

At the moment, the mechanistic reasons for coevolution between SLC30A9 and mitochondrial oxidative phosphorylation chain components are not completely clear. ERC suggests functional, but not necessarily physical interaction between these proteins. The coevolution may reflect the shared function of zinc flux and utilization in the mitochondria, as discussed above. Furthermore, SLC30A9 appears to specifically coevolve with the components of the peripheral stalk of the mitochondrial ATP synthase (Figure 4B) suggesting a shared role in proton transport, as the membrane components of the peripheral stalk were suggested to contribute to the formation of the proton channel [77]. Further investigation would help uncover the mechanism behind the coevolution between SLC30A9 and mitochondrial respiratory chain components and further validate ERC as a cutting-edge tool of molecular discovery and characterization.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

This work was supported by the NIH grant 1R21NS111944 to K.K. and E.A., and by NIH grant 1R01HG009299 to M.C. and N.L.C.

Data Availability

All supporting data are included within the main article and its Supplementary Files.

Abbreviations

ERC	evolutionary rate covariation
HBSS	HEPES-based salt solution

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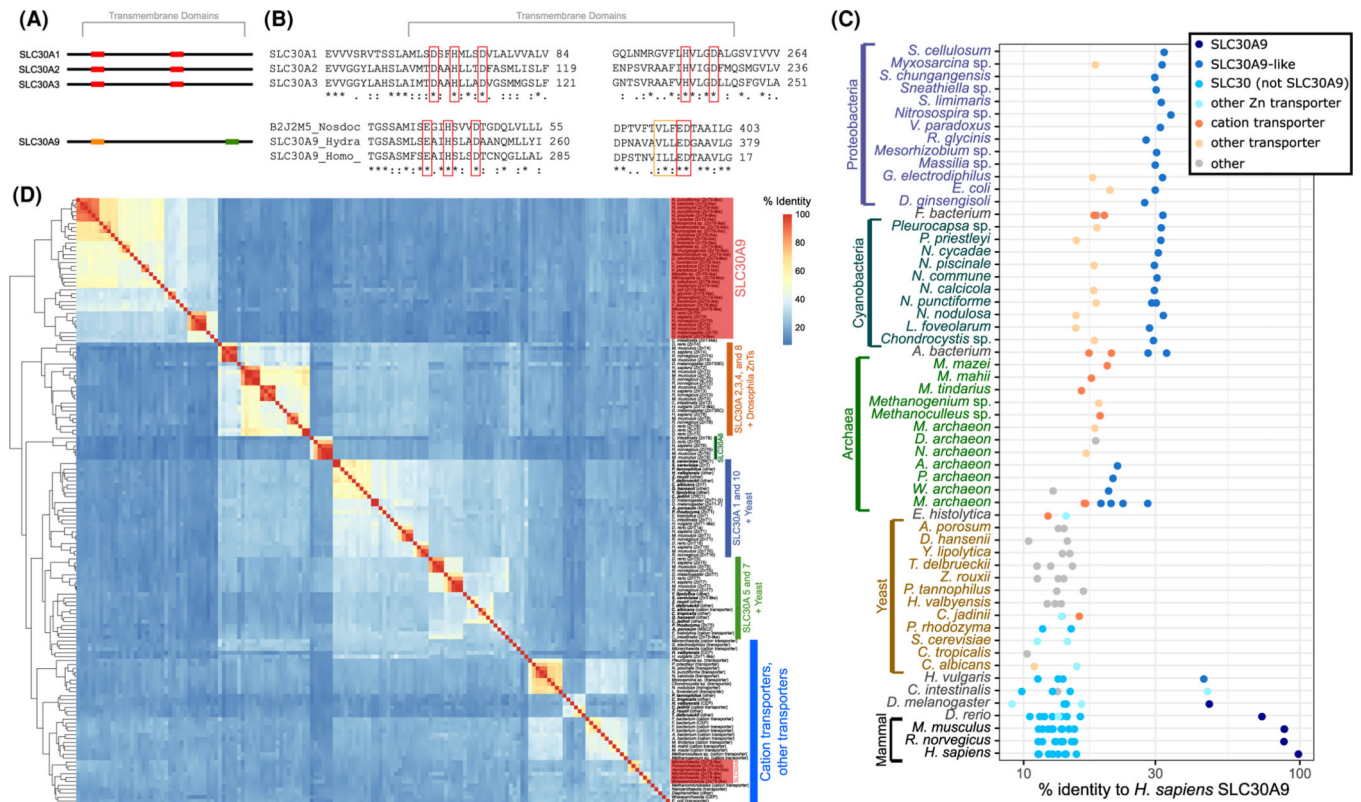


Figure 1. Defining features of SLC30A9.

(A) Cartoon indicating defining features of SLC30A9 and other SLC30As. **(B)** Partial amino acid alignment of select human SLC30A (top) and SLC30A9 from humans and other organisms (bottom), specifically focused on the conserved differences in the putative permeation center depicted in part A. **(C)** Bacterial SLC30A9s show higher percent identity with human SLC30A9 than do other animal SLC30As. This suggests potential divergence of SLC30A9 before animal duplication events give rise to other SLC30As. **(D)** Hierarchical clustering based on percent sequence identity demonstrates that bacterial and animal SLC30A9s cluster together separately from other animal SLC30As, yeast transporters, cation transporters, and other ion transporters, further supporting ancient divergence of SLC30A9 prior to duplication of other SLC30As.

* Bootstrap value at least 85



Figure 2. SLC30A dendrogram suggests ancient divergence of SLC30A9 and other zinc transporters.

The branch leading to all SLC30A9s, both mammalian and bacterial, as well as some other transporters, is supported with high bootstrap confidence. Other SLC30As in animals and some other transporters branch separately.

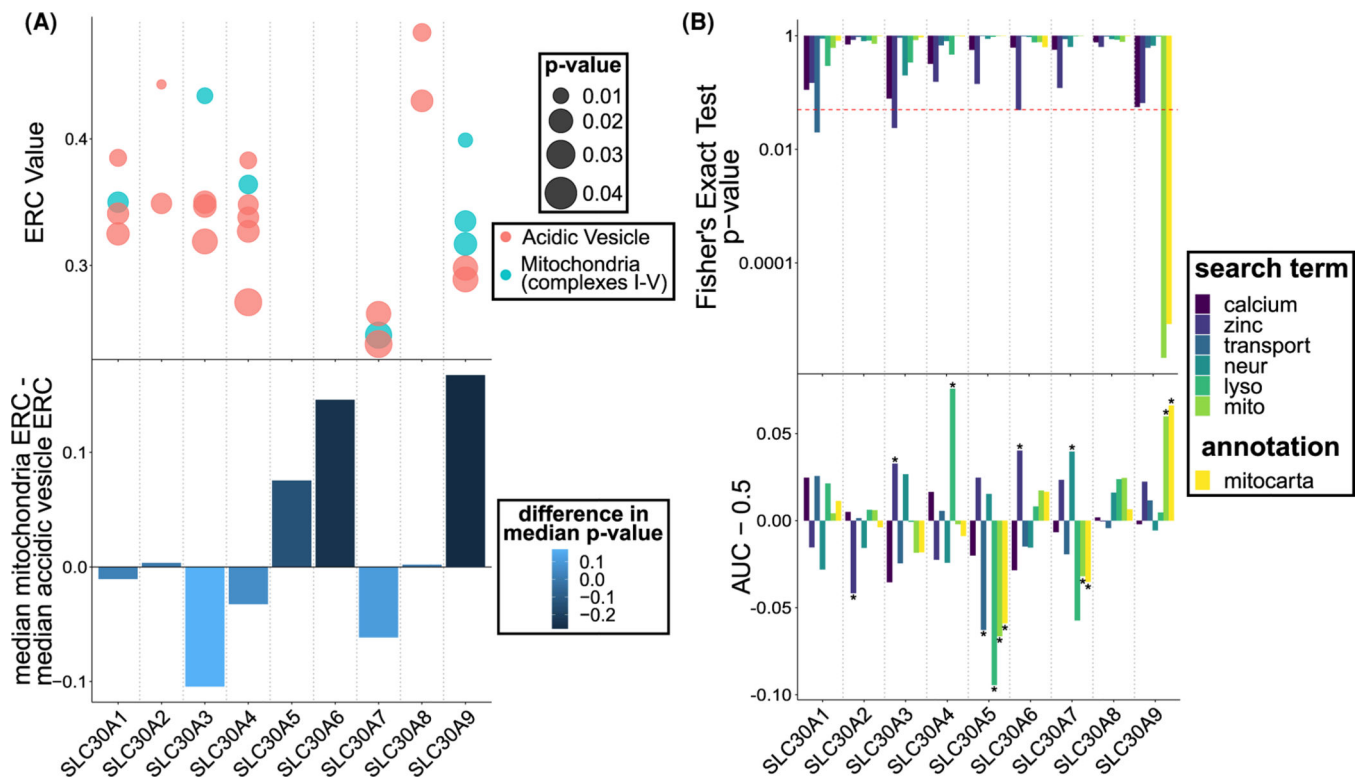


Figure 3. ERC analysis of SLC30 proteins reveals that SLC30A9 (ZnT9) shows unique patterns of correlated evolutionary rates with mitochondrial proteins.

(A) ERC between SLC30A9 and mitochondrial proteins shows higher correlation and lower P -value compared with ERC between SLC30A9 and lysosomal proteins. This contrast is stronger for SLC30A9 than for any other SLC30. Dots represent genes with positive ERC values and P -values less than 0.05. Size of dots are continuously scaled with the P -value. Bars heights show difference in median ERC value between mitochondrial and lysosomal proteins for all genes and bar colors show difference in median P -value between mitochondrial and lysosomal proteins for all genes. (B) SLC30A9 genes with high ERC values are uniquely enriched for mitochondrial function. Search terms represent words identified in gene descriptions to assign them to that term. Also included are Mitocarta annotations. One-tailed Fisher's Exact Tests were performed to test for significant enrichment of search terms and annotations with ERC values above 0.05, and notably only SLC30A9 shows a significant P -value for mitochondria (bar beyond the red dashed line). A Wilcoxon rank-sum test also showed a uniquely significant positive shift of ERC values for mitochondrial terms and SLC30A9 (asterisks show significance). Centered AUC values demonstrate whether the term was enriched (positive values) or depleted (negative values).

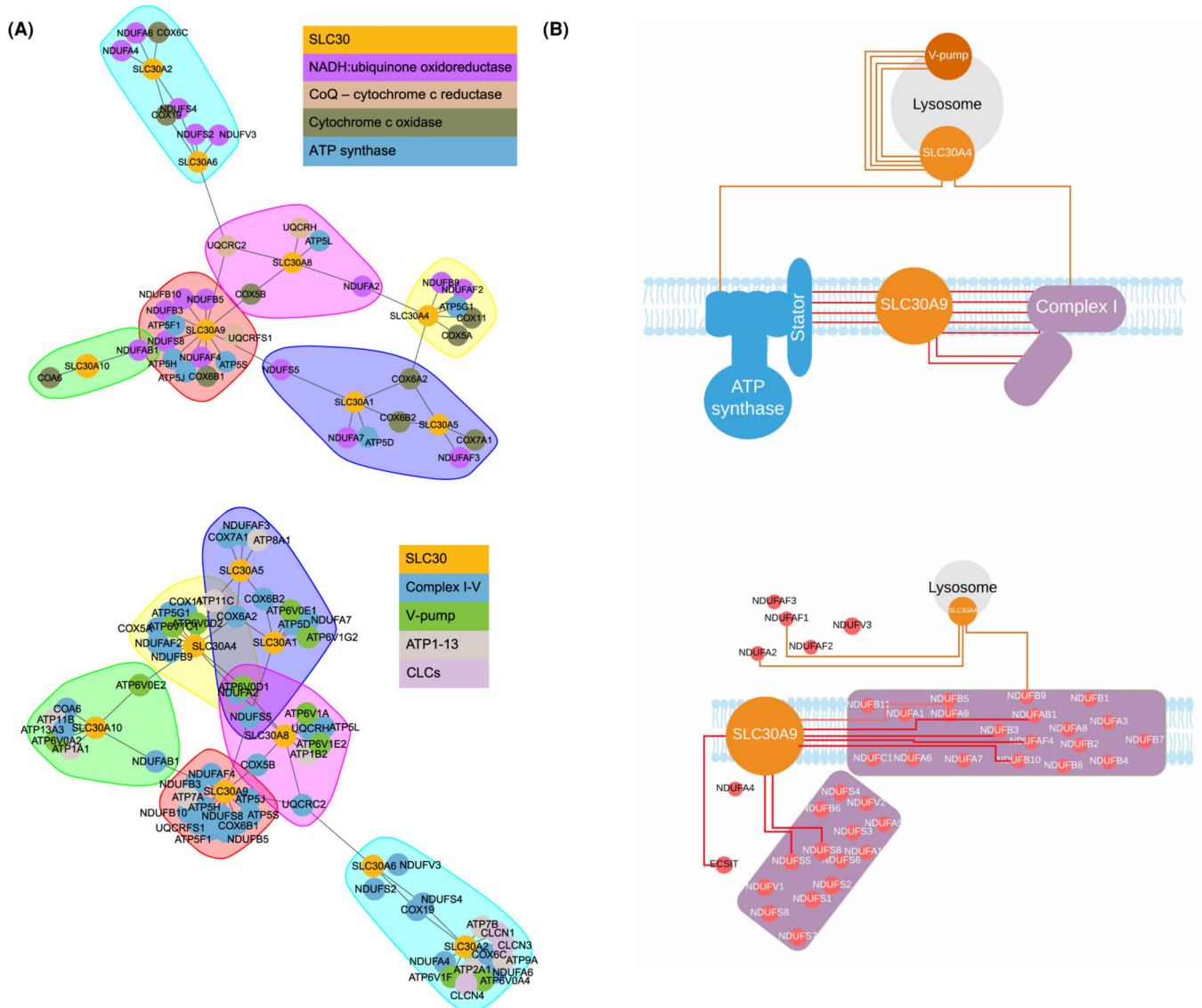


Figure 4. SLC30 evolutionary network clusters and diagrams of SLC30A9 interaction with mitochondrial components.

(A) Clustering of SLC30A transporters with mitochondrial proteins. Clusters highlighted with arbitrary colors indicate communities identified during clustering. Top: SLC30A9 shows robust clustering with numerous mitochondria-related proteins, including NDUF and ATP proteins. Note the cluster including SLC30A9 (in red) with many NADH proteins (purple dots) and ATP synthase proteins (blue dots). Bottom: Clustering using an expanded geneset shows distinctive clustering of SLC30A9 (red cluster) with numerous mitochondria complex I–V proteins (blue circles). **(B)** Wire diagram comparing ERC-positive interactions (at $R > 0.3$) between SLC30A9 and the mitochondrial OxPhos components. SLC30A4 is shown as a control.

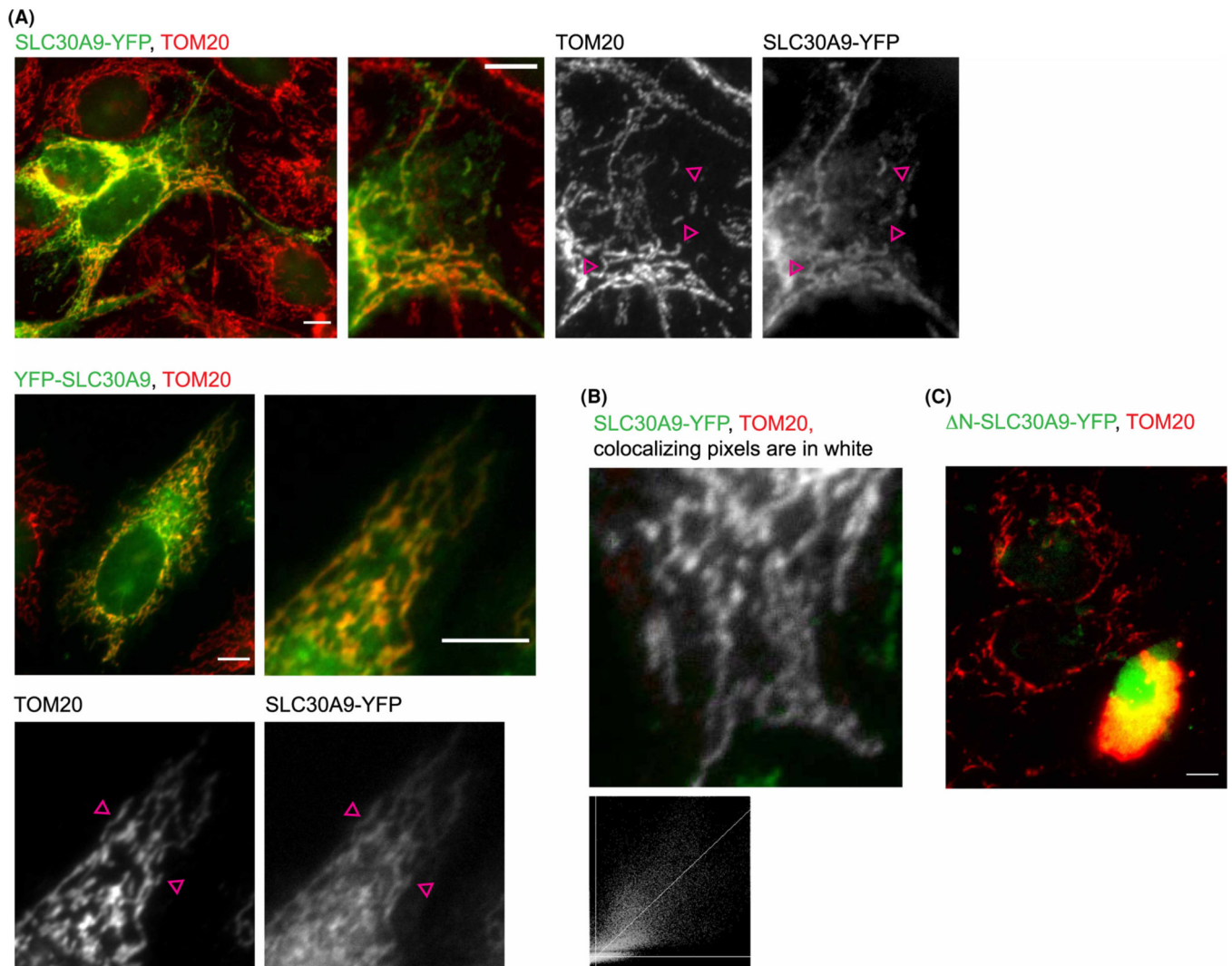


Figure 5. SLC30A9 localization.

(A) Widefield fluorescence image of N-terminal (top) and C-terminal (bottom) YFP fusion of human SLC30A9 (green) expressed in HeLa cells and co-stained with antibodies against TOM20 (red). In addition, zoom-in images are shown for clarity. Pink arrows point to examples of overlapping stains. Scale bars are 10 μm . (B) An example of colocalization analysis of SLC30A9 and TOM20 obtained using the Colocalization threshold Plugin of FIJI. Histogram below shows linear regression of intensities of the overlapping pixels in the red and green channels. (C) Evidence of cell death in cells overexpressing N-terminal deletion of SLC30A9. A representative widefield image of HeLa cells transfected with GFP-tagged N-terminal deletion construct of human SLC30A9 stained as above. Note the swelling and detachment of the transfected cell as well as swelling and compacting at the nucleus of the mitochondria, indicating cell death.

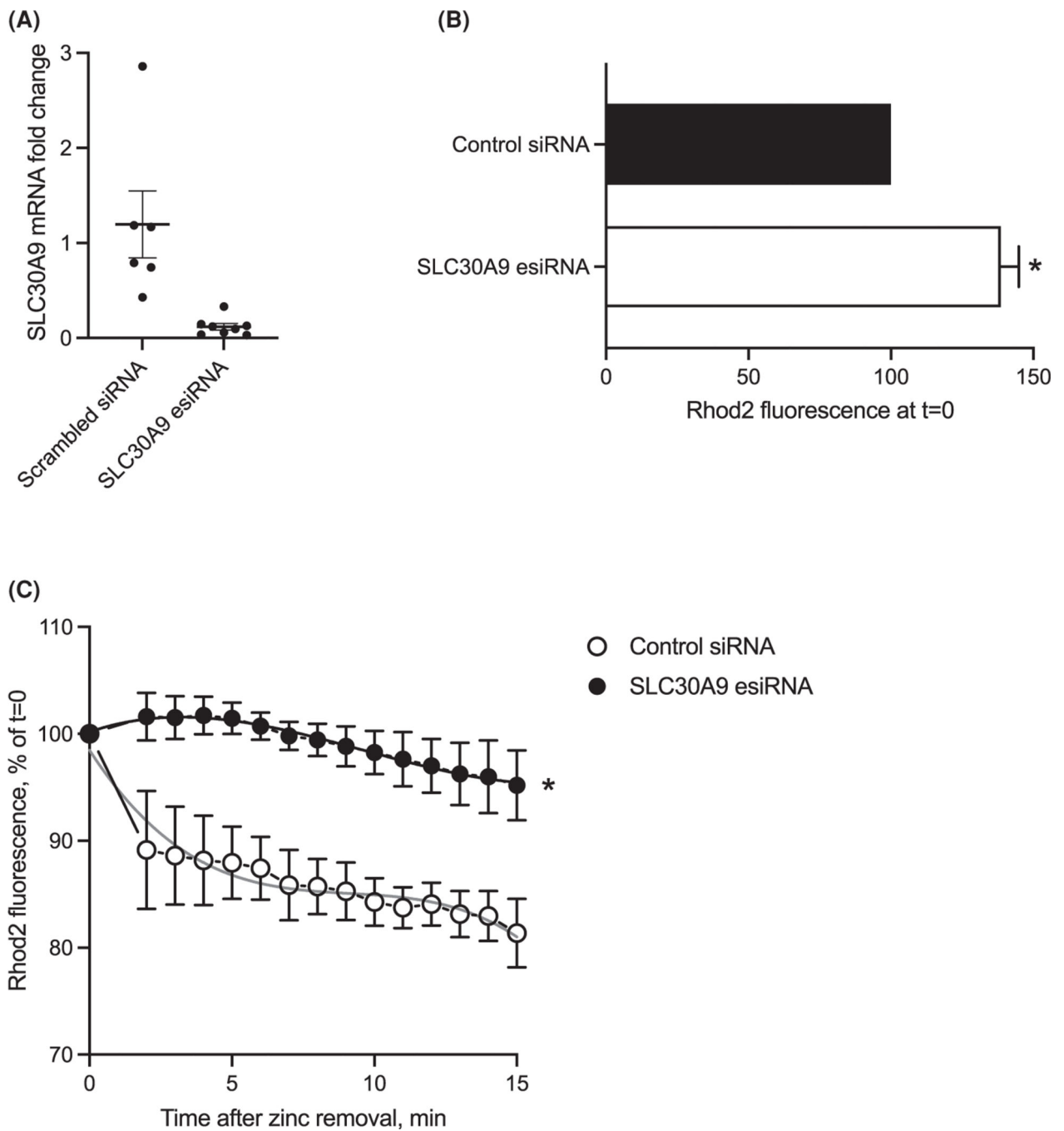


Figure 6. Mitochondrial zinc handling deficits in SLC30A9-knockdown HeLa cells.

(A) qPCR analysis of HeLa cells transfected with SLC30A9 esiRNA and harvested 24 h post-transfection. (B) Mitochondrial zinc content in control and SLC30A9-transfected HeLa cells was evaluated using Rhod-2, am. The cells were exposed to zinc for 24 h. Average fluorescence in each region of interest containing clearly identifiable mitochondria was recorded and normalized to the values in the cells that were transfected with scrambled shRNA. (C) The delayed loss of mitochondrial zinc in SLC30A9-deficient cells exposed to nominally zinc-free medium after 24 h long zinc load. The cells were treated as above; at t

= 0 the zinc-containing medium was replaced with nominally zinc free medium and images were taken every 60 s. The fluorescence intensity was normalized to the values recorded at $t = 0$, which was taken as 100%. In Panels **A** and **B**, data represent three independent trials involving 7–10 cells. Data are shown \pm SEM. Asterisks represent statistical significance at $P < 0.05$, for Student's test in Panel **A**. In Panel **B**, the curves were fitted using a third order polynomial curve (smooth gray lines) and an extra sum-of-squares test was performed to answer whether one curve can adequately fit both datasets. That hypothesis was rejected at $P < 0.0001$ level. Here and below, statistical analysis and data plotting were performed using Prism 9.

Table 1

Expression of SLC30A9, MCU, MICU, MMT and their homologs in various organisms (x shows expression)

Clade	Species	ZNT9	MCU	MICU	MMT
Animals	<i>Homo sapiens</i>	x	x	x	
	<i>Mus musculus</i>	x	x	x	
	<i>Rattus norvegicus</i>	x	x	x	
	<i>Danio rerio</i>	x	x	x	
	<i>Ciona intestinalis</i>	x	x	x	
	<i>Hydra vulgaris</i>	x	x	x	
	<i>Drosophila melanogaster</i>	x	x	x	
Fungi	<i>Yarrowia lipolytica</i>				x
	<i>Saccharomyces cerevisiae</i>				x
	<i>Debaryomyces hansenii</i>				x
	<i>Pachysolen tannophilus</i>				x
	<i>Williopsis saturnus</i>		x		x
	<i>Ustilago maydis</i> 521				x
	<i>Cryptococcus neoformans</i>		x		x
	<i>Laccaria bicolor</i> S238N-H82				x
	<i>Magnaporthe oryzae</i> 70–15		x		x
	<i>Penicillium chrysogenum</i> 54–1255		x		x
<i>Uncinocarpus reesii</i> 1704				x	
Green Algae	<i>Ostreococcus lucimarinus</i> CCE9901	x	x	x	
	<i>Chlamydomonas reinhardtii</i>	x	x	x	
	<i>Volvox carteri f. nagarensis</i>	x	x	x	
Protists	<i>Trichomonas vaginalis</i> G3		x	x	
	<i>Leishmania infantum</i> JPCM5				
	<i>Trypanosoma brucei</i> TREU927		x	x	
	<i>Tetrahymena thermophila</i> SB210		x	x	
	<i>Paramecium tetraurelia</i> strain d4–2		x		
	<i>Phaeodactylum tricorutum</i> CCAP 1055/1	x			
Archaea	<i>Plasmodium chabaudi chabaudi</i>				
	<i>Candidatus Pacearchaeota archaeon</i>	x			
	<i>Candidatus Woesearchaeota archaeon</i>	x			
	<i>Candidatus Aenigmarchaeota archaeon</i> ex4484_52	x			
	<i>Candidatus Diapherotrites archaeon</i>				
	<i>Methanohalophilus mahii</i>				
	<i>Methanolobus tindarius</i>				
<i>Methanosarcina mazei</i>					
Bacteria	<i>Sneathiella chungangensis</i>	x			
	<i>Massilia</i> sp. LC238	x			
	<i>Geopsychrobacter electrodiphilus</i>	x			
	<i>Escherichia coli</i>	x			
	<i>Chondrocystis</i> sp. NIES-4102	x			
	<i>Nosdoc punctiforme</i>	x			
	<i>Leptolyngbya foveolarum</i>	x			

Clade	Species	ZNT9	MCU	MICU	MMT
	<i>Nostoc cycadae</i>	x			

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