



Research Paper

A single-cysteine mutant and chimeras of essential *Leishmania* Erv can complement the loss of Erv1 but not of Mia40 in yeast

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ARTICLE INFO

Keywords:

ALR
CHCHD4
Erv
Intermembrane space
Leishmania
Mia40
Mitochondria
Oxidative protein folding

ABSTRACT

Mia40/CHCHD4 and Erv1/ALR are essential for oxidative protein folding in the mitochondrial intermembrane space of yeast and mammals. In contrast, many protists, including important apicomplexan and kinetoplastid parasites, lack Mia40. Furthermore, the Erv homolog of the model parasite *Leishmania tarentolae* (*LtErv*) was shown to be incompatible with *Saccharomyces cerevisiae* Mia40 (*ScMia40*). Here we addressed structure-function relationships of *ScErv1* and *LtErv* as well as their compatibility with the oxidative protein folding system in yeast using chimeric, truncated, and mutant Erv constructs. Chimeras between the N-terminal arm of *ScErv1* and a variety of truncated *LtErv* constructs were able to rescue yeast cells that lack *ScErv1*. Yeast cells were also viable when only a single cysteine residue was replaced in *LtErv*^{C17S}. Thus, the presence and position of the C-terminal arm and the kinetoplastid-specific second (KISS) domain of *LtErv* did not interfere with its functionality in the yeast system, whereas a relatively conserved cysteine residue before the flavodomain rendered *LtErv* incompatible with *ScMia40*. The question whether parasite Erv homologs might also exert the function of Mia40 was addressed in another set of complementation assays. However, neither the KISS domain nor other truncated or mutant *LtErv* constructs were able to rescue yeast cells that lack *ScMia40*. The general relevance of Erv and its candidate substrate small Tim1 was analyzed for the related parasite *L. infantum*. Repeated unsuccessful knockout attempts suggest that both genes are essential in this human pathogen and underline the potential of mitochondrial protein import pathways for future intervention strategies.

1. Introduction

The vast majority of mitochondrial proteins are synthesized at cytosolic ribosomes and have to be imported into one of the four mitochondrial compartments [1]. One of these compartments, the intermembrane space (IMS) between the inner and outer mitochondrial membrane, harbors proteins that contribute to respiration, programmed cell death, the assembly of protein complexes, and the transfer of ions, metabolites and proteins [2–6]. In addition to a functional classification, IMS proteins can be also grouped according to their protein import pathway (reviewed in [2,7]): One group of

proteins, hereinafter referred to as class I proteins, contains a positively charged N-terminal matrix-targeting signal that is followed by a hydrophobic transmembrane segment. This bipartite pre-sequence is recognized by the TIM23 complex, which inserts class I proteins into the inner mitochondrial membrane where they often undergo limited proteolysis. For example, yeast cytochrome *b*₂ is imported via this pathway and its soluble domain is subsequently released into the IMS by the inner membrane protease Imp1 [8–10]. Another group of proteins from yeast and mammals, hereinafter referred to as class 2 proteins, is oxidatively trapped in the IMS by a system that consists of Mia40 and Erv1 in yeast or the respective mammalian homologs CHCHD4 and ALR

Abbreviations: GSH, reduced glutathione; IMS, mitochondrial intermembrane space; *Li*, *Leishmania infantum*; *Lt*, *Leishmania tarentolae*; *Pf*, *Plasmodium falciparum*; *Sc*, *Saccharomyces cerevisiae*

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<https://doi.org/10.1016/j.redox.2017.12.010>

Received 25 August 2017; Received in revised form 17 December 2017; Accepted 21 December 2017

Available online 23 December 2017

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[2,11–17]. The oxidoreductase Mia40 recognizes substrates with twin Cx₃C- or twin Cx₉C-motifs as well as proteins with more specialized cysteine patterns such as Erv1 [3,7,17–20]. Reference substrates of this pathway in *Saccharomyces cerevisiae* include the twin Cx₃C proteins ScTim9 and ScTim13, the twin Cx₉C cytochrome c oxidase assembly factor ScPet191, and the copper chaperone for superoxide dismutase ScCcs1 [11,12,15,21–24]. During protein import, the reduced substrates undergo one or two rounds of dithiol-disulfide exchange reactions with the oxidized CPC-motif of Mia40, yielding the trapped oxidized substrate and reduced Mia40 [13,25–31]. Reduced Mia40 is subsequently oxidized by the flavoprotein Erv1, which transfers the electrons to cytochrome c and the respiratory chain [25,30,32–37]. In addition, ScErv1 was also suggested to transfer electrons to molecular oxygen, yielding H₂O₂ as a substrate of cytochrome c peroxidase [36], and to fumarate with the help of the fumarate reductase Osm1 [38].

As outlined above, oxidative protein folding in the IMS has been intensely studied in yeast and mammals. These organisms belong to the same group of eukaryotes termed Opisthokonta [39]. In contrast, numerous protists have no Mia40 homolog, although their genomes encode an Erv homolog as well as candidate substrates with twin Cx₃C- or twin Cx₉C-motifs [16,40–44]. This raises the fundamental question whether several of the major groups of eukaryotes have a replacement for Mia40 or whether their Erv homologs can oxidatively fold substrates without Mia40. Among the protists without a Mia40 homolog are numerous important non-related apicomplexan and kinetoplastid parasites, such as the causative agents of malaria (*Plasmodium falciparum*), sleeping sickness (*Trypanosoma brucei*), and visceral leishmaniasis (*Leishmania infantum*). Depletion of *T. brucei* Erv (*TbErv*) by RNA interference was shown to result in growth arrest and mitochondrial swelling [44]. Furthermore, down regulation of *TbErv* in combination with quantitative proteomics revealed the down regulation of 13 proteins with twin Cx₃C- or twin Cx₉C-motifs and led to the identification of 25 candidate substrates in the IMS of *T. brucei* [45]. The depletion studies strongly support a crucial role of *TbErv* in oxidative protein folding. However, it remains to be analyzed whether Erv is generally essential for kinetoplastid parasites and how oxidative protein folding can occur without Mia40. We previously showed that the IMS of the kinetoplastid model parasite *Leishmania tarentolae* harbors an Erv homolog (*LtErv*) as well as a twin Cx₃C protein termed small Tim1 (*LtsTim1*). Both proteins as well as the twin Cx₃C protein ScTim9 from yeast were imported into isolated mitochondria from *L. tarentolae* and yeast, suggesting that the import signals and the oxidative folding pathway have been functionally conserved during the course of evolution [42]. Heterologous *LtErv* was also imported into the IMS of yeast mitochondria *in vivo*. However, neither *LtErv* nor an Erv homolog from *P. falciparum* (*PfErv*) could functionally replace essential ScErv1 in yeast [43]. Here we (i) addressed the physiological relevance of Erv and small Tim1 for survival of the deadly human pathogen *L. infantum*, (ii) determined structure-function relationships of ScErv1 and *LtErv*, and (iii) analyzed the compatibility of *LtErv* with the oxidative protein folding system in yeast.

2. Materials and methods

2.1. *L. infantum* cell culture

L. infantum promastigotes (strain MHOM MA67ITMAP263) were cultured at 25 °C in RPMI 1640 Glutamax supplemented with 10% (v/v) inactivated fetal bovine serum (iFBS), 50 U/mL penicillin, 50 mg/mL streptomycin (Gibco), and 25 mM HEPES sodium salt, pH 7.4 (Sigma). Parasites were synchronized by three or four daily passages of 1 × 10⁶ cells/mL and subsequently seeded at 10⁶ cells/mL (day 0). Afterwards, growth curves for wild-type and mutant strains were recorded for up to 7 days by determining the cell density in a hemocytometer.

2.2. Generation of *L. infantum* knockout and rescue constructs

Fragments of the 5′ and 3′ UTRs of *L. infantum* ERV (*LIERV*) for homologous recombination were PCR-amplified from *L. infantum* genomic DNA using primer pairs P1/P2 and P3/P4 (Table S1). Both PCR products were cloned into the *HindIII/SpeI* and *BamHI/XbaI* restriction sites of plasmids pGL345 and pGL726, which carry coding sequences for hygromycin B phosphotransferase (*HygR*) or phleomycin hydrolase (*PhleoR*), respectively. Deletion constructs for *L. infantum* small TIM1 (*LsTIM1*) were generated likewise. Fragments of the 5′ and 3′ UTRs of *LsTIM1* were PCR-amplified with primers P5/P6 and P7/P8 (Table S1) and cloned into the *HindIII/SpeI* and *BamHI/XbaI* restriction sites of pGL345 and pGL726. Before transfection of *L. infantum*, the replacement cassettes were excised from the plasmids by digestion with *HindIII/BglII* and purified from agarose gels. Rescue plasmid pTEX-NEO-*LIERV* was assembled by cloning the *LIERV* ORF, which was PCR-amplified with primers P9/P10 (Table S1), into the *BamHI/HindIII* restriction sites of pTEX-NEO. This trypanosomatid expression vector carries a coding sequence for the neomycin phosphotransferase (*NeoR*) [46]. To generate the pTEX-NEO-*LsTIM1* rescue plasmid, the *LsTIM1* ORF was PCR-amplified with primer pair P11/P12 (Table S1) and cloned into the *BamHI/XhoI* sites of pTEX-NEO.

2.3. *L. infantum* transfection and selection

Transfections were performed in an Amaxa Nucleofector (Lonza) using program U-033 as described previously [47]. Briefly, 5 × 10⁷ logarithmic phase promastigotes were suspended in Human T Cell Nucleofector (Lonza) and transfected with 2–10 µg DNA. Parasites were allowed to recover in 10 mL of culture medium without selection drug for 24 h before centrifugation and plating on medium M199 agar plates containing 10% (v/v) iFBS, 50 U/mL penicillin, 50 mg/mL streptomycin, 0.1 mM adenine, 0.023 mM hemin, 25 mM HEPES sodium salt, pH 7.4 (all from Sigma), and the selective drug(s). Hygromycin (Invitrogen) was used at 15 µg/mL, G418 (Sigma) at 15 µg/mL, and phleomycin (Sigma) at 17.5 µg/mL. Colonies were picked after 1–2 weeks and transferred to liquid medium.

2.4. Generation and site-directed mutagenesis of yeast constructs

Full length *LTERV*, *LTERV*^{C17S}, *LTERV*^{C63S}, *LTERV*^{C300S} and *LTERV*^{C300S/C304S} in plasmid pQE30 as well as *SCERV1* and *PFERV* in pYX232 were cloned previously [43]. Additional mutant and chimeric pYX232-*LTERV* constructs were generated as follows. *LTERV*^{C92S/C109S} in pQE30 was generated by two rounds of site-directed mutagenesis using primer pairs P26/P27 and P28/P29 (Table S2) and pQE30-*LTERV* as a template. *LTERV*, *LTERV*^{C17S}, *LTERV*^{C63S}, *LTERV*^{C300S}, *LTERV*^{C300S/C304S} and *LTERV*^{C92S/C109S} were PCR-amplified using primer pair P30/P31 (Table S2) with the according pQE30 constructs as templates. The truncated constructs *LTERV*^{C-term} and *LTERV*^{N-term} were PCR-amplified using primer pairs P32/P31 and P30/P33 (Table S2) with pQE30-*LTERV* as a template. *Chimera 1–5* were amplified by overlap extension PCR according to a standard protocol [48] using sense primer P34, overlap primer pair P35/P36 or P37/P38, and one of the antisense primers P39–P41 (Table S2) with pYX232-*SCERV1* and pYX232-*LTERV* as templates. Fusion constructs encoding the N-terminal mitochondrial targeting signal and transmembrane segment of ScMia40 (*Mia40N*) *MIA40N-PFERV*, *MIA40N-LTERV*, *MIA40N-LTERV*^{N-term}, *MIA40N-LTERV*^{C-term}, *MIA40N-LTERV*^{C17S} and *MIA40N-Chimera 1–6* were also amplified by overlap extension PCR using the sense primer P48, one of the overlap primers P51–54, and one of the antisense primers P31/P33/P40/P41/P50. All PCR products were subsequently cloned into the *EcoRI/XhoI* restriction sites of the yeast expression vector pYX232. *Chimera 6* was generated by site-directed mutagenesis using primer pair P42/P43 (Table S2). The truncated Mia40 variant *SCMIA40*^{short} (encoding residues 1–70 and 284–403) was previously cloned into the

single-copy plasmid pRS314 [49] and was a kind gift by Johannes Herrmann. All constructs were confirmed by commercial sequencing (GATC Biotech).

2.5. Yeast cell culture and complementation assays

S. cerevisiae cells were grown at 30 °C in 1% yeast extract, 2% peptone (YP) medium or in synthetic (S) minimal medium supplemented with or without the amino acids lysine, leucine, histidine, and tryptophan as well as the bases adenine and uracil [50]. For growth analysis, the medium contained either 2% glucose (YPD, SD), 3% galactose (SGal) or 3% glycerol (SG). Plasmid shuffling experiments were performed as described previously [43] with a YPH501-derived $\Delta erv1$ strain (genotype *ura3-52*, *lys2-801*^{amber}, *ade2-101*^{ocre}, *trp1- Δ 63*, *his3- Δ 200*, *leu2- Δ 1*, *erv1::His3*) that harbors a *SCERV1* copy under control of the *MET25* promoter on the *URA3*-containing plasmid pRS426 [51]. Briefly, cells were transformed with the *TRP1*-containing plasmids pYX232-*SCERV1*, pYX232-*LTERV*, pYX232-*Chimera 1–6*, pYX232-*LTERV*^{C-term}, pYX232-*LTERV*^{N-term}, pYX232-*LTERV*^{C17S}, pYX232-*LTERV*^{C63S}, pYX232-*LTERV*^{C92S/C109S}, pYX232-*LTERV*^{C300S}, pYX232-*LTERV*^{C300S/C304S} and the respective *MIA40N* constructs according to a standard protocol [52]. After transfection and selection on SD agar plates without histidine, uracil and tryptophan, yeast strains were grown overnight in uracil-containing liquid medium. Overnight cultures were diluted in sterile water to an optical density at 600 nm (OD₆₀₀) of 0.1 and further diluted 1:10, 1:100 and 1:1000. Drops (10 μ L) of the overnight cultures and each serial dilution were spotted on SD agar plates without histidine and tryptophan containing 50 mg/L uracil and 1 g/L 5-fluoroorotic acid (FOA) for negative selection and plasmid shuffling [53]. The same dilutions were spotted on SD agar plates without uracil, histidine and tryptophan as a control. Analogous complementation assays for plasmids pYX232-*PFERV*, pYX232-*LTERV*, pYX232-*LTERV*^{C-term}, pYX232-*LTERV*^{N-term}, pYX232-*LTERV*^{C17S}, the respective *MIA40N* constructs and pRS314-*SCMIA40*^{short} were performed with a $\Delta mia40$ strain, which has the same genetic background with a *mia40::S.pombeHis5* instead of the *erv1::His3* replacement. This strain harbors a copy of a short functional version of *SCMIA40* under the control of its endogenous promoter on the *URA3*-containing plasmid pRS316 [49]. All plates were incubated at 30 °C or the indicated temperature for up to one week. Single colonies or, in the absence of colonies, swabs of background cells from FOA plates were subsequently incubated overnight or for up to 3 days in 1 mL FOA- and uracil-containing SD liquid medium (either without histidine and tryptophan in the case of $\Delta erv1$ cells or without tryptophan when handling $\Delta mia40$ cells). If cell growth was detected, liquid cultures were diluted as described above and 10 μ L drops of each dilution were spotted again on FOA- and uracil-containing SD agar plates without histidine and tryptophan as well as SD agar control plates without FOA, uracil, histidine and tryptophan in order to validate the genotype.

2.6. Cell fractionation and western blot analysis

Clones from plasmid shuffling experiments that only grew on uracil-containing medium were incubated in 50 mL SD liquid medium without histidine and tryptophan until they had reached an OD₆₀₀ of 1.0–1.8. Ten OD volumes of cells were harvested by centrifugation (1750 \times g, 10 min, 4 °C) and disrupted as described previously [54]. Briefly, yeast cells were resuspended in 300 μ L ice-cold buffer containing 0.6 M sorbitol, 80 mM KCl, 20 mM MOPS, pH 7.4 that was freshly supplemented with 1 mM phenylmethylsulfonyl fluoride (PMSF) before 200 μ L glass beads (\varnothing 0.5 mm, Sigma-Aldrich) were added. The mixture was vortexed six times for 30 s before glass beads and cell debris were removed by centrifugation (1000 \times g, 3 min, 4 °C). The supernatant was centrifuged again (20,000 \times g, 15 min, 4 °C) to separate the mitochondria-containing pellet fraction from the cytosol-containing supernatant fraction. The pellet fraction was directly boiled in 2 \times SDS-

PAGE Laemmli-buffer. The supernatant fraction was precipitated by the addition of 5–6 volumes of ice-cold acetone to 150 μ L supernatant and incubated overnight at – 20 °C. The mixture was subsequently centrifuged (15,000 \times g, 15 min, 4 °C) and the precipitate was air dried for 15 min at room temperature before boiling in 2 \times SDS-PAGE Laemmli-buffer. For western blot analysis, 20 μ L of the pellet sample and 10 μ L of the supernatant sample were separated by SDS-PAGE on a 15% gel followed by wet blotting and immunodecoration with antibodies against *LtErv* (1:500 [42]), *ScErv1* (1:2000 [13]), *ScZwf1* (1:10000 [55]), and *ScTim50* (1:250 [56]). Samples for western blot analysis of *L. infantum* promastigotes were prepared as follows. Parasites were harvested by centrifugation (3000 \times g, 10 min, 4 °C) and washed twice in ice-cold PBS. Parasites were suspended at 6.25 \times 10⁸ cells/mL in 4% (w/v) SDS, 50 mM Tris-HCl, pH 7.4 and stored at – 70 °C until further use. Before separation by SDS-PAGE, 8 μ L extracts were mixed with 12 μ L PBS and 5 μ L 5 \times SDS-PAGE Laemmli-buffer and heated for 10 min at 65 °C.

2.7. Yeast DNA extraction and PCR analysis

Plasmid DNA was extracted from yeast as described previously [57]. For PCR analysis, 1 μ L of extracted DNA was mixed with 1 μ M of primer pairs P44/P45, P44/P46, and P47/P45, 400 μ M dNTP mix (Thermo Scientific) and 2.5 U *Taq* DNA polymerase (NEB) in a final volume of 50 μ L to amplify *SCERV1*, *Chimera 3*, and *Chimera 6*, respectively. Initial denaturation was carried out at 94 °C for 2 min, followed by 30 cycles of denaturation at 94 °C for 45 s, annealing at 60 °C for 45 s, and elongation at 68 °C for 45 s. The last cycle was followed by a final elongation step at 68 °C for 90 s.

2.8. Yeast growth assays on agar plates and in liquid medium

For growth assays on agar plates, yeast strains were incubated in 1 mL liquid SD medium without histidine and tryptophan and cultured in the logarithmic growth phase for up to five days. Drop dilutions assays in sterile water were subsequently performed as described above. Dilutions were spotted on SD, SG or SGal agar plates without histidine and tryptophan. All plates were incubated at 30 °C for up to 12 days. Growth assays in liquid medium were performed in a Tecan Infinite F200 Pro plate reader using transparent flat bottom 96-well-plates (Greiner). Briefly, 200 μ L of liquid cultures in SD, SG or SGal medium were diluted to an absorbance of 0.1 in the according medium and were grown at 30 °C for 48 h. Plates were shaken at 183 rpm for 550 s before each hourly measurement.

2.9. Protein import into isolated mitochondria

Radiolabeled Tim13 precursor protein was synthesized *in vitro* in a TNT-coupled reticulocyte lysate system according to the manufacturer's standard protocol (Promega). Mitochondria were isolated as described [13] and quantified based on their protein content using a Bradford protein assay (BioRad). The import of Tim13 precursor protein into isolated mitochondria was performed in import buffer [56] containing 0.1 mg/mL BSA, 5 mM NADH, 2.5 mM ATP and 2 mM 2-mercaptoethanol. The import reaction contained 15% of lysate and 100 μ g of mitochondria in 100 μ L import buffer. The import reaction was incubated for 10 min at 25 °C and then stopped by diluting the reaction tenfold with ice-cold SH-buffer (0.6 M sorbitol, 20 mM HEPES-KOH, pH 7.4) with 50 μ g/mL proteinase K. Proteinase K treatment was performed for 20 min and then stopped by addition of 2 mM PMSF. Following isolation of mitochondria, the import of precursor protein was analyzed by SDS-PAGE and autoradiography.

2.10. Mia40 redox mobility shift assays

Mitochondria (1 mg/mL) were incubated for 20 min at 25 °C in SH-

buffer with or without 5 mM reduced glutathione (GSH). Following incubation, the samples were treated for 20 min at 25 °C with 40 mM iodoacetamide in 50 mM Tris (pH 8.0), 0.6 M sorbitol to prevent artificial oxidation. The redox state of re-isolated mitochondria was analyzed by non-reducing SDS-PAGE and western blotting with antibodies against ScMia40. The buffers used were first degassed for 10 min using a Vario vacuum pump (Vacuubrand) and then flushed with nitrogen. All sample cups were also flushed with nitrogen.

2.11. Ni-NTA pull-down of His-tagged ScMia40

Pull-down assays were performed with a yeast strain that encodes a His-tagged version of ScMia40 on the plasmid pYX142 replacing wild type ScMia40 [27]. This strain was transformed with the plasmid pYX232-*Chimera 2* or the plasmid pYX232-*SCERV1*. Mitochondria were isolated from cells that were grown in selective medium without tryptophan and leucine containing 2% lactate and 0.5% galactose. For the Ni-NTA-agarose pull-down, mitochondria were solubilized in buffer containing 0.5% (w/v) Triton X-100 in buffer A (20 mM Tris, 80 mM KCl, 30 mM imidazole, 1 mM PMSF, pH 7.4) for 30 min at 4 °C. After a clarifying spin, solubilized material was incubated with 25 µL of Ni-NTA-agarose beads for 1 h. Following removal of the supernatant, beads were washed three times with buffer A containing 0.05% Triton X-100 and bound proteins were eluted with Laemmli buffer containing 300 mM imidazole. Total and elution fractions were then analyzed by SDS-PAGE and western blotting.

3. Results

3.1. ERV and small TIM1 cannot be deleted in *L. infantum*

Kinetoplastid parasites lack Mia40, which raises the question whether they can survive in the absence of Erv or putative substrates. In order to test the relevance of Erv in *Leishmania*, we tried to disrupt the encoding gene in the human pathogen *L. infantum* (locus LinJ.15.1020) by homologous recombination (Fig. 1A). Clonal heterozygous *erv*^{+/-} strains were readily obtained after transfection of *L. infantum* promastigotes with plasmid pGL345-*LIERV* and selection with hygromycin as confirmed by PCR analysis (Fig. 1B). Growth curve analysis for liquid cultures of wild-type (wt) and *erv*^{+/-} strains revealed no morphological abnormalities or drastic growth defects (Fig. 1C), and both strains had similar LiErv protein levels (Fig. S1A). The second *LIERV* allele could not be deleted after subsequent transfection with plasmid pGL726-*LIERV* and selection with phleomycin, even after the introduction of an additional gene copy on plasmid pTEX-NEO-*LIERV* (Fig. 1B). The additional *LIERV* copy was functional as reflected by a drastic increase in the LiErv protein concentration (Fig. S1A), which might explain the short lag phase and decreased maximum cell density for liquid cultures of *erv*^{+/-} strains that carried the episomal *LIERV* copy when grown in the presence of the selection antibiotic G418 (Fig. 1C).

To exclude systematic methodological causes for the negative results regarding the generation of homozygous *erv*^{-/-} knockout parasites, we performed parallel experiments for *LisTIM1* (locus LinJ.25.1610) using the same knockout strategy with plasmids pGL345-*LisTIM1* and pGL726-*LisTIM1* (Fig. 1A). *LisTim1*, the homolog of the IMS marker protein small Tim1 from *L. tarentolae*, is a candidate substrate for oxidative protein folding in *Leishmania* mitochondria [42]. Heterozygous hygromycin-resistant *stim*^{+/-} parasites revealed no abnormal morphology, drastic growth defects or altered protein levels compared to the wt strain (Fig. 1C, Fig. S1B). A homozygous *stim*^{-/-} knockout strain was only obtained in the presence of an additional gene copy on plasmid pTEX-NEO-*LisTIM1* (Fig. 1B). Again, complemented strains had a slight growth defect in the presence of G418 (Fig. 1C) and the protein concentration of *LisTim1* was drastically up-regulated (Fig. S1B). The complemented *stim*^{-/-} knockout strain neither lost its plasmid nor *LisTim1* and remained resistant to G418 even after twelve months of

continuous culture in the absence of the selection drug, whereas a wt control strain lost the empty plasmid pTEX-NEO over time (Fig. 1D, Fig. S1B). Thus, small Tim1 is probably essential in promastigotes of *L. infantum*, indicating the importance of the oxidative folding pathway for parasite survival despite the absence of a Mia40 homolog. Repeated negative results for parallel *LIERV* knockout attempts with *LisTIM1* as a control also support the interpretation that Erv is essential in *L. infantum*.

3.2. Chimeric LtErv can complement the loss of ScErv1

LtErv appears to function in *Leishmania* without a Mia40 homolog and, therefore, may require structural features that are incompatible with the oxidative protein folding system in yeast. To analyze which structure-function relationships might be relevant for the inability of *LtErv* to complement the loss of ScErv1 in yeast [43], we generated a set of chimeras. *LtErv* differs from ScErv1 in two major aspects, (i) the C-terminal addition of a kinetoplastida-specific second (KISS) domain and (ii) the position of two redox-active 'distal' or 'shuttle' cysteine residues at the C-terminal instead of the N-terminal arm (Fig. 2A). The shuttle cysteines transfer electrons to the conserved 'proximal' cysteines within the flavodomain [16,43]. To address the relevance of these structural elements, we either replaced the N-terminus before the Erv domain of *LtErv* with the N-terminal arm of ScErv1 (chimera 1–3) or the complete N-terminal half of *LtErv* with ScErv1 (chimera 4 and 5). A potential negative effect of the C-terminal arm or the KISS domain was addressed by systematically removing these structural elements in chimera 2, 3 and 5. Furthermore, to check whether the shuttle CQVYC-motif of *LtErv* is incompatible with the oxidative folding machinery in yeast, we replaced the CRSC-motif at the N-terminal arm of ScErv1 in chimera 6. The chimeric constructs were analyzed in complementation assays on agar plates with 5-fluoroorotic acid (FOA) for negative selection against the episomal copy of *SCERV1* in a haploid Δ *erv1* knockout strain (Fig. 2B). In contrast to the negative control, which was transformed with wild-type *LTERV*, viable yeast strains were obtained from FOA agar plates for all chimeras. The absence of ScErv1 and the presence of the chimeric proteins in yeast mitochondria were confirmed for at least three different clones of each construct-harboring strain by subcellular fractionation and western blot analysis (Fig. 2C, Fig. S2A). Furthermore, we confirmed the genotype for chimera 3 and 6 by analytic PCR (Fig. S2B).

Replacing the N-terminus of *LtErv* with the N-terminal arm of ScErv1 allowed *LtErv* to rescue the Δ *erv1* strain in complementation assays (Fig. 2B, Fig. S3A). Yeast cells with chimera 3 grew slowly but were viable, suggesting that the Erv domain of *LtErv* in combination with the N-terminal arm of ScErv1 is sufficient to be functional. Addition of the C-terminal arm and/or the KISS domain improved the growth of chimera 1 and 2, which was still not as good as the growth of the positive control with ScErv1. This suggests that the Erv domain of *LtErv* might not be able to fully replace the function of the endogenous Erv domain (Fig. S3A). Taking into account the improved growth of chimera 1 and 2 compared to chimera 3, it is unlikely that the KISS domain and the C-terminal arm have a deleterious effect on the function of *LtErv* in yeast. Complementation assays for chimera 4 and 5, in which the KISS domain alone or together with the C-terminal arm of *LtErv* were fused to the ScErv1 domain, also support the absence of a dominant-negative effect (Fig. 2B), for example, by inactivating Mia40. Furthermore, the CQVYC-motif of *LtErv* in chimera 6 could functionally replace the CRSC-motif at the N-terminal arm of ScErv1. Thus, the shuttle arm motif of *LtErv* is, at least in part, compatible with yeast Mia40. In summary, attachment of the N-terminal arm of ScErv1 to *LtErv* chimeras suffices to complement and rescue the Δ *erv1* strain. Neither the presence of the KISS domain or the C-terminal arm nor the shuttle cysteine motif of *LtErv* render the protein inactive and cause the incompatibility of *LtErv* with the oxidative folding machinery in yeast mitochondria.

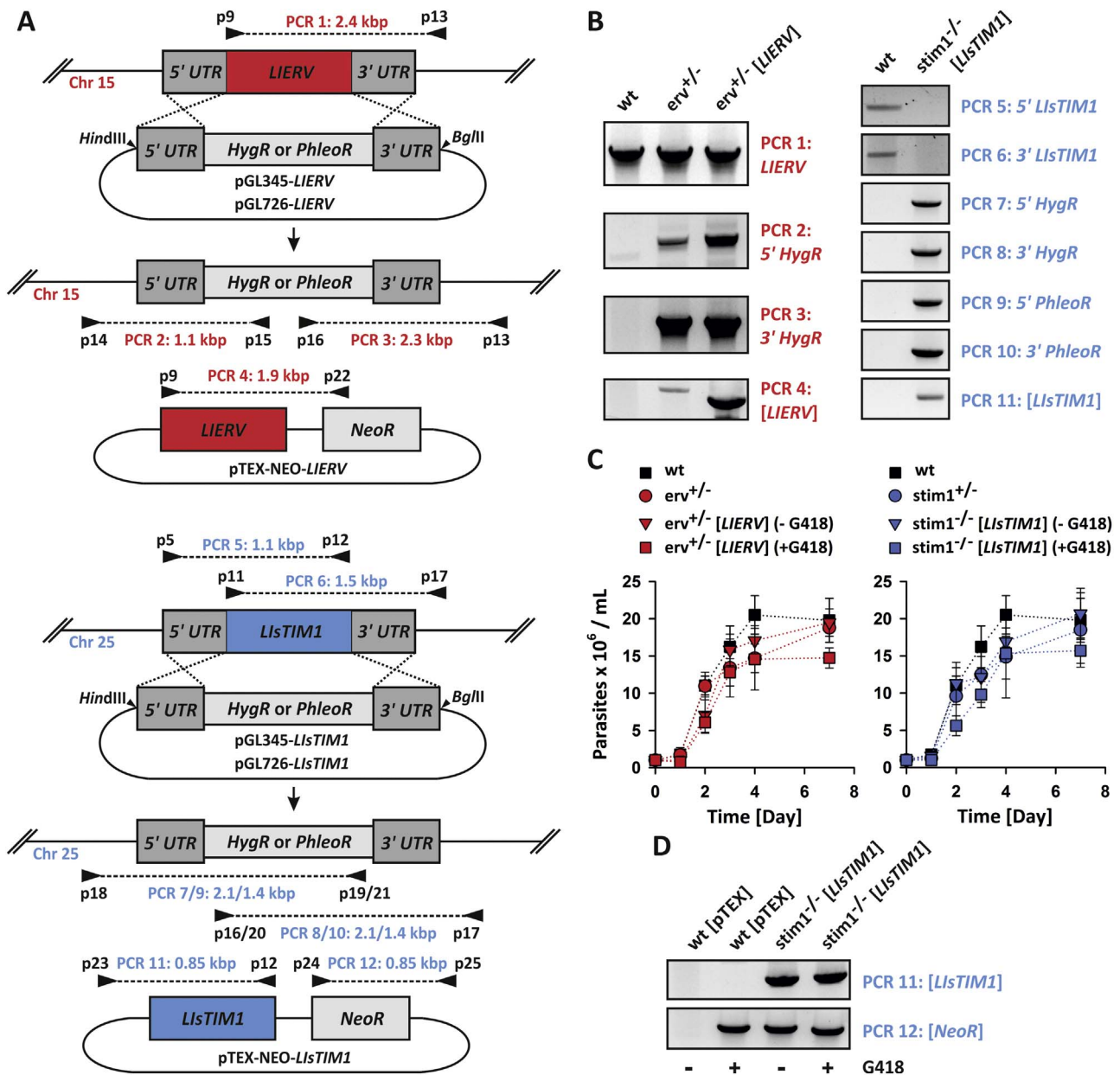


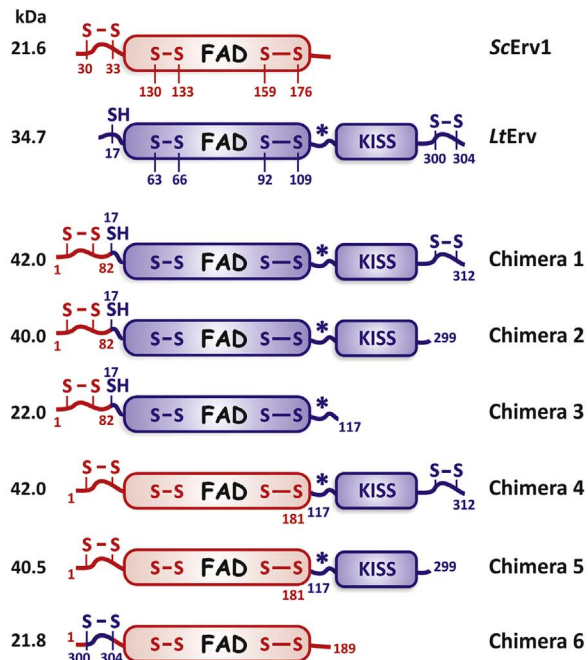
Fig. 1. Knockout attempts for *ERV* and *STIM1* in *Leishmania infantum*. (A) Schematic summary of the *LIERV* and *LIsTIM1* knockout strategy by homologous recombination using excised replacement cassettes from plasmids pGL345 and pGL726. Different selectable marker genes (*HygR* and *PhleoR*) were used in two rounds of transfection to target both alleles in diploid promastigotes. Experiments were performed with and without episomal copies of *LIERV* or *LIsTIM1* on rescue plasmid pTEX-NEO. Dashed lines indicate the sites of homologous recombination at the 5' and 3' untranslated regions (UTRs). Expected product sizes from analytic PCR reactions 1–12 are highlighted. (B) PCR analysis of the indicated parasite lines for (i) the absence or presence of chromosomal *LIERV* (PCR 1) and *LIsTIM1* (PCR 5 and 6), (ii) the 5' and 3' integration of the selectable marker genes (PCR 2, 3 and 7–10), and (iii) the absence or presence of episomal *LIERV* (PCR 4) or *LIsTIM1* (PCR 11). (C) Growth curve analysis for synchronized liquid cultures of the indicated parasite lines. Parasites that were complemented with an episomal copy of *LIERV* or *LIsTIM1* were grown with or without 50 μ g/mL G418 (+/- G418). Values represent the mean and standard deviation of three independent biological replicates. (D) PCR analysis of the indicated parasite lines for the absence or presence of episomal *LIsTIM1* (PCR 11) or the pTEX-NEO selection marker *NeoR* (PCR 12) after twelve months of continuous culture in the absence or presence of 15 μ g/mL G418.

3.3. Mutant *LtErv*^{C17S} can complement the loss of *ScErv1*

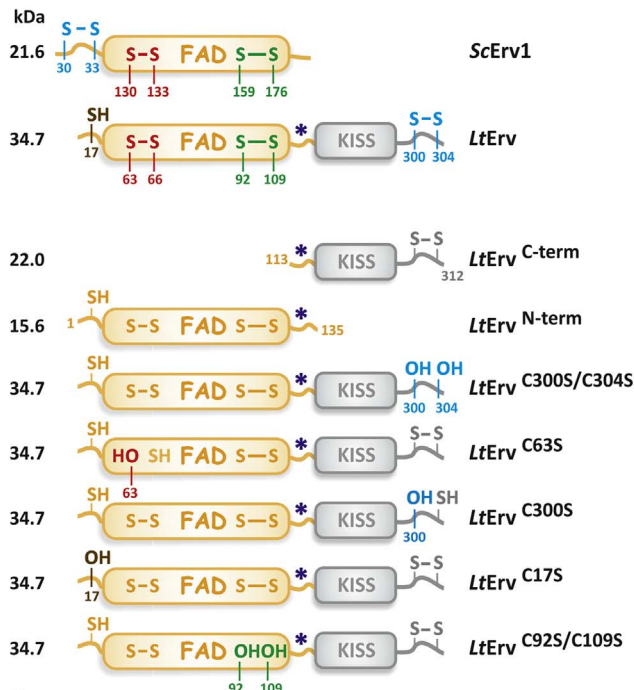
The chimera complementation assays suggested that *LtErv* can, at least in principle, functionally replace *ScErv1*. To further address the structural requirements for a successful complementation, we analyzed additional *LtErv* constructs in which we removed the N-terminal or C-terminal half of the protein (*LtErv*^{C-term} and *LtErv*^{N-term}) or replaced the different cysteine residues (Fig. 2D). Neither *LtErv*^{C-term} nor *LtErv*^{N-term} was able to complement the loss of *ScErv1* (Fig. 2E, Fig. S3A). The latter result indicates that the flavodomain plus the N-terminal sequence of *LtErv* is not sufficient to complement for *ScErv1* in yeast. It also excludes that the inability of full-length *LtErv* to complement is due to a

negative effect of the C-terminal half on its otherwise functional N-terminal half (in accordance with the growth of the truncated chimera in Fig. 2B). Systematic mutation of the distal, proximal, structural, and clamp cysteine residues of *LtErv* revealed that replacement of clamp residue Cys17 results in a functional mutant that was able to complement the loss of *ScErv1* on FOA agar plates (Fig. 2E, Fig. S3A). The absence of *ScErv1* and the presence of *LtErv*^{C17S} in yeast mitochondria were confirmed for different *LTERV*^{C17S} clones by subcellular fractionation and western blot analysis. Clones of all other *LtErv* mutants were either not viable or still contained *ScErv1* (Fig. 2F, Fig. S4). Noteworthy, mutations of the distal cysteine residues in *LtErv*^{C300S/C304S} or of one of the proximal cysteines in *LtErv*^{C63S} appeared to have no effect

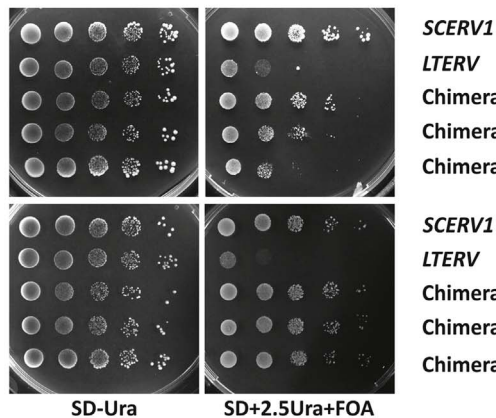
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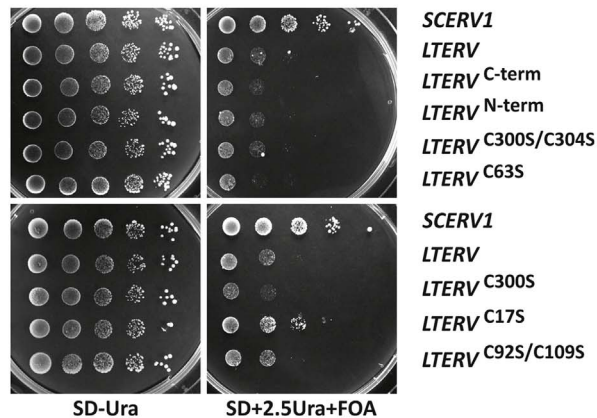
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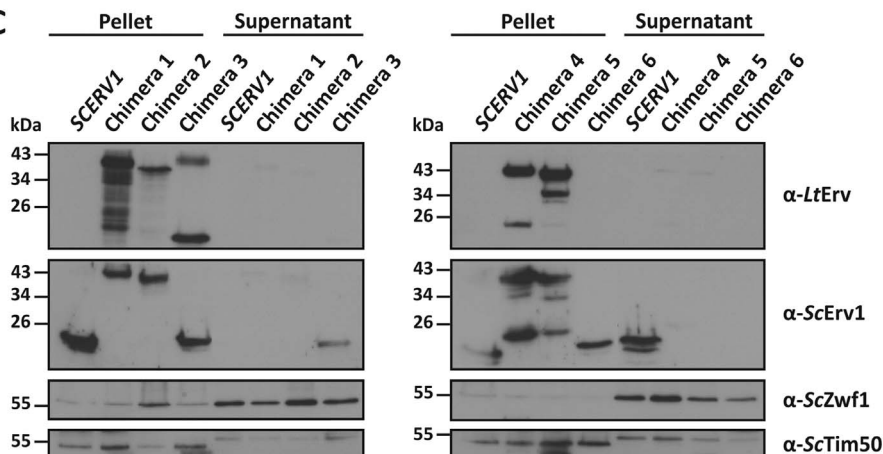
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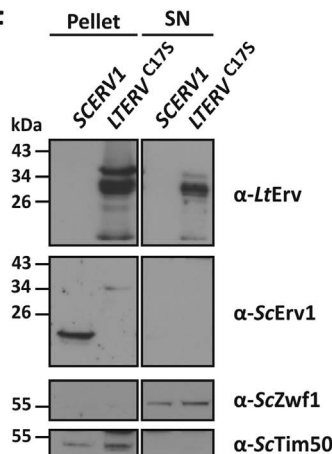
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F



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Fig. 2. Yeast *SCERV1* complementation assays with chimeric and mutant *LTERV*. (A) Schematic summary of the constructs encoding ScErv1 (red), LtErv (purple) and the indicated chimera 1–6. The binding site for the peptide antibody α -LtErv [42] is marked with an asterisk. (B) A Δ erv1 strain with an episomal copy of *SCERV1* on a *URA3*-containing rescue plasmid was subjected to plasmid shuffling in the presence of FOA after transfection with pYX232 plasmids that encode the indicated chimera. Copies of *SCERV1* and *LTERV* on plasmid pYX232 served as positive and negative controls, respectively [43]. Representative drop dilution assays (from left to right) on agar plates with or without FOA are shown for one of four independent biological replicates. (C) Western blot analyses of mitochondria-containing pellet fractions and cytosol-containing supernatant fractions of Δ erv1 strains harboring the indicated chimera. Cytosolic glucose-6-phosphate dehydrogenase (ScZwf1) and mitochondrial ScTim50 served as controls. Expected molecular masses of the chimera are indicated in panel A. The western blots are representative for one clone of each construct (see also Fig. S2A). The absence of ScErv1 and presence of chimera 3 and 6 was also confirmed by analytic PCR (Fig. S2B). (D) Schematic summary of truncation constructs and cysteine mutants of LtErv. (E) Plasmid shuffling experiments with mutant *LTERV* as in panel B. (F) Cell fractionation of a Δ erv1 strain that was complemented with *LTERV*^{C17S} as in panel C (see also Fig. S4). SN, supernatant.

on the mitochondrial localization of the proteins, whereas LtErv^{C17S} was also found in the cytosolic fraction.

In order to test whether the presence of Cys17 might have masked a positive complementation for the other LtErv mutants, we repeated the *SCERV1* complementation experiments with the C17S constructs LtErv^{N-term/C17S}, LtErv^{C17S/C63S}, LtErv^{C17S/C300S}, LtErv^{C17S/C300S/C304S}, and LtErv^{C17S/C92S/C109S}. However, none of these constructs could complement the loss of ScErv1 (Fig. S5).

In summary, Cys17 renders LtErv non-functional in the yeast system, and the presence of this cysteine residue is, at least in part, the cause for the incompatibility between LtErv and the oxidative folding machinery in yeast mitochondria. Replacing either residue Cys17 with serine or residues 1–16 of LtErv with the N-terminus of ScErv1 in chimera 1–3 partially rescues the functionality of LtErv and its compatibility with the yeast system.

3.4. LtErv^{C17S} and chimera 1, 2 and 6 have specific carbon source-dependent growth defects

To further characterize the complemented yeast strains, we compared their growth in liquid or on agar minimal medium with glucose (SD), galactose (SGal) or glycerol (SG) as a carbon source (Fig. 3 and Fig. S6). Chimera 1 had a pronounced growth defect on all three carbon sources. This phenotype was observed in liquid medium and on agar plates. In contrast, chimera 2 showed a growth defect in SG liquid medium but not in SGal or SD medium. Chimera 2 also grew normal on SG agar plates. Chimera 3 revealed no striking phenotype, whereas chimera 4 and 5 grew slightly faster on SGal and SG agar plates than the control. This phenotype was absent in liquid medium. Chimera 6 revealed a growth defect on SG agar plates and in SD liquid medium. Cells with LtErv^{C17S} grew normally in SD medium but reached a much lower

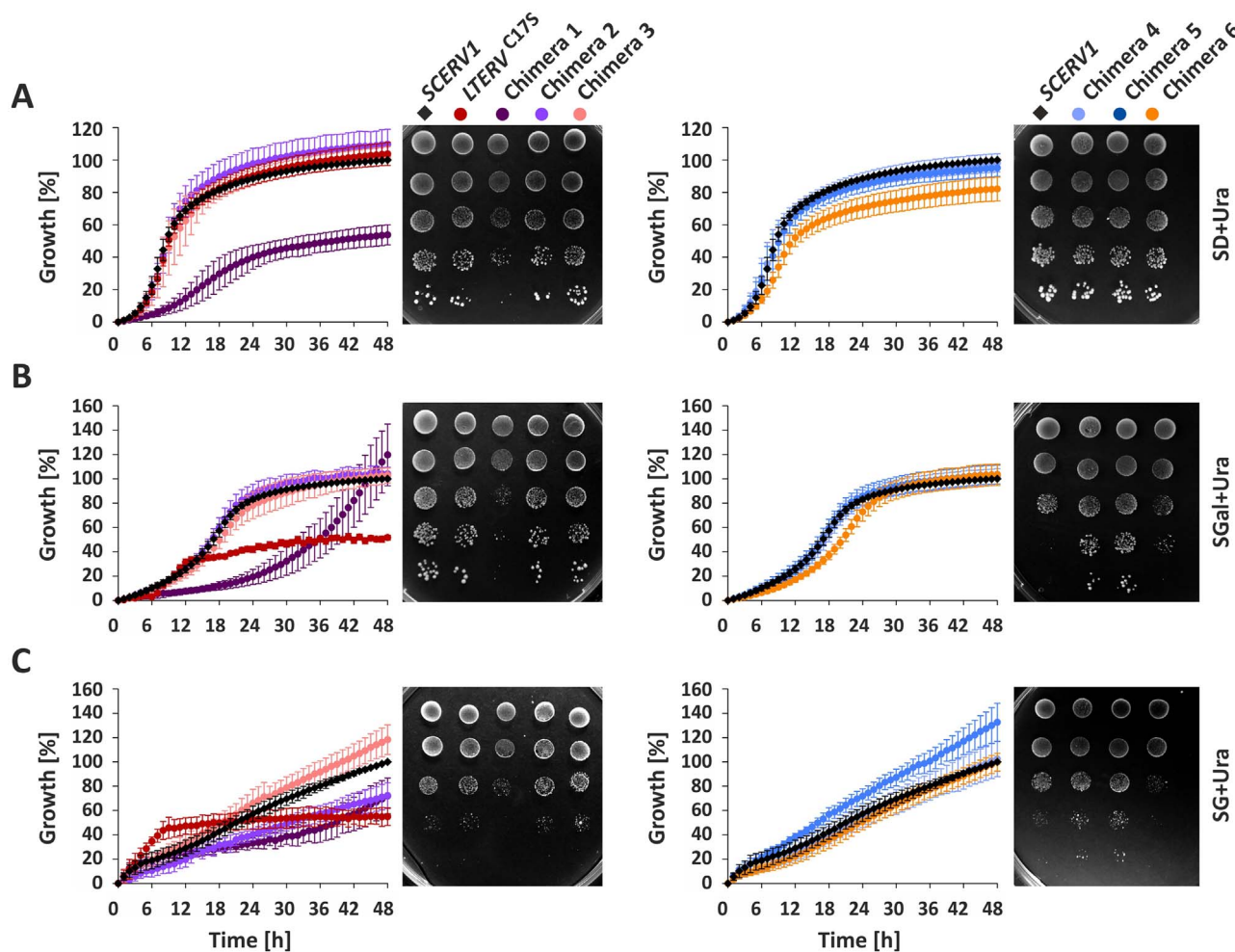


Fig. 3. Growth assays of complemented Δ erv1 strains. Yeast strains harboring Chimera 1–6 or LtErv^{C17S} were analyzed in liquid medium and on agar plates. (A) Growth on minimal medium with 2% glucose as a carbon source (SD). (B) Growth on 3% galactose medium (SGal). (C) Growth on 3% glycerol medium (SG). Growth curves in liquid medium were normalized to the growth of the Δ erv1 strain harboring an episomal copy of *SCERV1* on plasmid pYX232. Values in liquid medium represent the mean and standard deviation of three to five independent biological replicates. Representative drop dilution assays (from top to bottom) on agar plates on day 3 (SD), day 5 (SGal) and day 8 (SG) are shown for one of three to four independent biological replicates. The phenotypes were confirmed for two or three different clones for each construct.

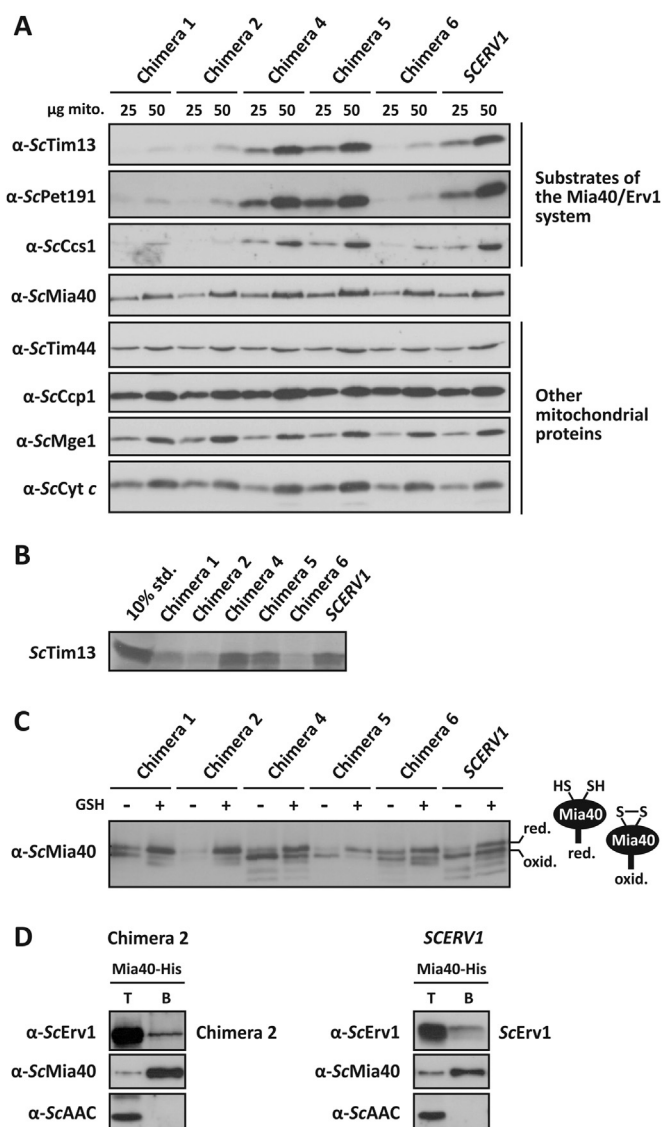


Fig. 4. Functional analysis of *LtErv* chimera in yeast. (A) Western blot analysis of steady-state protein levels in mitochondrial preparations that contain the indicated chimera or ScErv1 as a control. The twin Cx₃C protein ScTim13, the twin Cx₃C cytochrome c oxidase assembly factor ScPet191, and the copper chaperone for superoxide dismutase ScCcs1 represent different substrate classes of the Mia40/Erv1 system [15,21–24]. ScTim44 and the co-chaperone ScMge1 contain a pre-sequence and are imported into the mitochondrial matrix, whereas the IMS proteins cytochrome c and cytochrome c peroxidase ScCcp1 are imported independent of the Mia40/Erv1 system. (B) *In vitro* protein import of [³⁵S]-methionine-labeled ScTim13 into isolated mitochondria harboring the indicated chimera or ScErv1 as a control. Ten percent of the labeled protein served as a loading control (10% std.). (C) Western blot analysis of the redox state of ScMia40 in isolated mitochondria that contain the indicated chimera or ScErv1 as a control. Mitochondria were treated with or without 5 mM GSH before protein separation by non-reducing SDS-PAGE. Red., reduced; oxid., oxidized. (D) Co-isolation of chimera 2 with His-tagged ScMia40. Mitochondrial lysates were prepared from strains containing His-tagged ScMia40 and episomally encoded chimera 2 or ScErv1. The mitochondrial lysates were incubated with Ni-NTA beads and the beads were isolated. Total proteins, T (15%), and bound material, B (100%), were analyzed by SDS-PAGE and immunoblotting with antibodies against ScErv1, ScMia40 and the ATP/ADP-carrier (ScAAC).

final cell density in SGal or SG liquid medium. Noteworthy, the strain grew faster during the initial 12 h in SG medium than the control. The phenotype of *LtErv*^{C17S} was specific for liquid medium and was absent on agar plates. The different phenotypes for *LtErv*^{C17S} on SD and SGal medium are quite surprising taking into account that galactose is converted to glucose-6-phosphate by the Leloir pathway [58], which should not be directly affected by the mitochondria. In summary,

pronounced growth defects on SG medium for chimera 1 and 2 in liquid medium as well as for chimera 6 on agar plates point towards an impaired mitochondrial respiration in these strains.

3.5. Chimera 1, 2 and 6 have an impaired oxidative folding pathway

Analysis of the substrates of the Mia40/Erv1 system by western blotting revealed reduced steady-state protein levels in mitochondria with chimera 1, 2 and 6 compared to control mitochondria with ScErv1 (Fig. 4A). In contrast, mitochondria with chimera 4 and 5 had normal steady-state substrate concentrations. The results suggest that the CRSC-motif of ScErv1 has been optimized for the interaction with ScMia40 and/or the ScErv1 flavodomain during the course of evolution, as reflected by chimera 6, which functions with a decreased efficiency compared to the control. Furthermore, the N-terminal arm of ScErv1 and the proximal cysteines of the flavodomain of *LtErv* in chimera 1 and 2 are not perfectly compatible.

The altered steady-state concentrations were specific for class II IMS proteins, which are oxidatively trapped [2]. Other mitochondrial proteins were not affected by the chimeras (Fig. 4A). This also includes ScMia40 itself, which has an N-terminal mitochondrial targeting signal and a transmembrane segment that is inserted into the inner mitochondrial membrane by the TIM23 complex before ScMia40 is oxidatively folded by the Mia40/Erv1 system [11,12,18,25,59]. An impaired protein import of the class II reference protein ScTim13 was confirmed using *in vitro* import assays (Fig. 4B). Again, mitochondria with chimera 1, 2 and 6, but not with chimera 4 and 5, imported less ScTim13 compared to mitochondria with ScErv1.

The underlying cause for the impaired protein import of class II proteins was presumably the altered ratio between reduced and oxidized ScMia40, as revealed by redox mobility shift assays and western blot analysis (Fig. 4C). The redox state of ScMia40 from isolated mitochondria with chimera 1, 2 and 6 was shifted to the reduced state compared to the redox state of ScMia40 from mitochondria with ScErv1 or chimera 4 and 5, which were predominantly oxidized. Thus, oxidation of ScMia40 by ScErv1 or chimera 4 and 5 is probably more efficient than the oxidation of ScMia40 by chimera 1, 2 and 6. This interpretation is also supported when isolated mitochondria were challenged with GSH. GSH treatment yielded almost completely reduced ScMia40 for chimera 1 and 2 in contrast to mitochondria with chimera 4 or ScErv1 (Fig. 4C).

Oxidation of ScMia40 by ScErv1 requires the physical interaction of both proteins. We previously showed that *LtErv* cannot replace ScErv1 and that wild type *LtErv* does not interact with ScMia40 [43]. Taking into account that an attachment of the N-terminal arm of ScErv1 to the flavodomain of *LtErv* rescued the deletion of *SCERV1* (Fig. 2), we tested an interaction with ScMia40 for one of these chimera in a Ni-NTA pull-down experiment. Indeed, His-tagged ScMia40 co-precipitated with chimera 2, albeit with reduced efficiency compared to ScErv1 (Fig. 4D). As negative control, a hydrophobic inner membrane protein, the ATP/ADP carrier, was not co-precipitated, indicating the specificity of the pull-down experiment. The experiment supports the hypothesis that the N-terminal arm of ScErv1 of the chimera oxidizes ScMia40 and subsequently transfers the electrons to the flavodomain of *LtErv*. However, the functionality of the N-terminal arm of ScErv1 appears to be impaired by the presence of the flavodomain of *LtErv*, which shifts the steady-state equilibrium between oxidized and reduced ScMia40 towards the reduced form.

In summary, oxidation of ScMia40 by chimera 1, 2 and 6 appears to be less efficient than oxidation by chimera 4 and 5. The subsequent increase of reduced ScMia40 in mitochondria with chimera 1, 2 and 6 most likely impairs the import of substrates of the Mia40/Erv1 system *in vitro* and *in vivo* and results in decreased levels of these proteins in mitochondria.

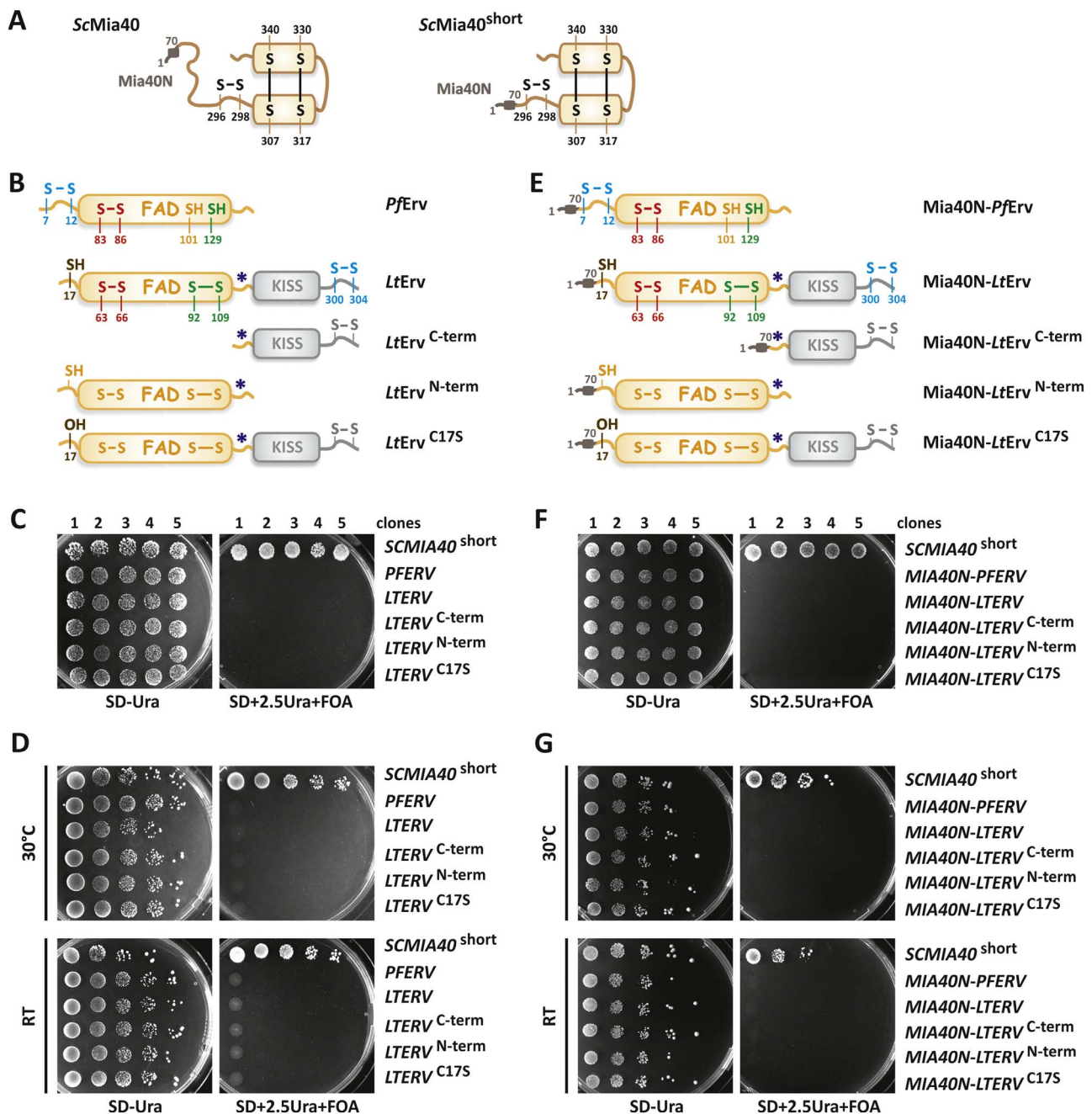


Fig. 5. Yeast *SCMIA40* complementation assays with truncated and mutant *LtErv*. (A) Schematic summary of full length and truncated *ScMia40*^{short} with the N-terminal mitochondrial targeting signal and transmembrane segment (*Mia40N*) and disulfide bonds highlighted. (B) Schematic summary of the constructs encoding *PfErv* as well as full length, truncated or mutant *LtErv*. (C) A Δ *mia40* strain with an episomal copy of *SCMIA40* on a *URA3*-containing rescue plasmid was subjected to plasmid shuffling in the presence of FOA after transfection with pYX232 plasmids that encode the indicated parasite *Erv* constructs. The complementation with *SCMIA40*^{short} served as a positive control. (D) The complementation assays were also repeated in parallel as drop dilution assays at 30 °C and room temperature (RT). Representative drop dilution assays (from left to right) on agar plates with or without FOA are shown for one of three independent biological replicates. (E) Schematic summary of the constructs encoding *Mia40N*-tagged *PfErv* as well as full length, truncated or mutant *LtErv*. (F) and (G) Complementation assays with *Mia40N*-tagged constructs.

3.6. *LtErv* cannot functionally replace yeast *Mia40*

Our results show that the KISS domain does not interfere with the function of *LtErv* in yeast. A potential function of the KISS domain, which lacks a cysteine residue [43], might be the recruitment of protein thiol substrates before they interact with the shuttle disulfide. Thus, the KISS domain and the shuttle arm might functionally replace *Mia40*, which does not only introduce disulfide bonds but also serves as a receptor and binds to a hydrophobic patch of the imported substrate [26,28,29,31,49,60–63]. To test the hypothesis whether protist *Erv* homologs are able to replace *Mia40*, we tried to complement a Δ *mia40*

strain with genes encoding the *Erv* homolog from *P. falciparum* (which lacks a KISS domain), full length *LtErv*, *LtErv*^{C-term}, *LtErv*^{N-term}, and *LtErv*^{C17S} (Fig. 5). Negative selection on FOA agar plates against the episomal copy of a short functional version of *SCMIA40* in the Δ *mia40* knockout strain revealed that none of the constructs could complement the loss of *ScMia40* (Fig. 5C). This was also the case when alternative temperatures were tested (Fig. 5D, Fig. S3B).

In a subsequent set of experiments, we fused the N-terminal mitochondrial targeting signal and transmembrane segment of *ScMia40* to our protist *Erv* constructs in order to localize the proteins to the same location in the inner mitochondrial membrane as endogenous *ScMia40*

(Fig. 5E). However, none of these Mia40N-fusion constructs was able to complement the loss of ScMia40 (Fig. 5F,G). In a last set of experiments, we addressed the hypothesis whether some of the chimeric *LtErv* constructs from Fig. 2 have a Mia40-like activity. We therefore tested chimera 1–6 as well as their fusion constructs with the N-terminal mitochondrial targeting signal and transmembrane segment of ScMia40 in *SCMIA40* complementation assays. Again, none of the twelve constructs could complement the loss of ScMia40 (Fig. S7). In summary, neither *PfErv* nor *LtErv* or structural elements of *LtErv*, such as the KISS domain and the C-terminal arm, can functionally replace ScMia40.

4. Discussion

Our knockout studies for *LIERV* and *LlTIM1* suggest that both genes are essential for parasite survival and underline the potential of mitochondrial protein import pathways for future intervention strategies against important pathogens. An analogous knockout strategy has been previously used to show that a cytosolic trypanredoxin is essential for *L. infantum* survival [64]. Furthermore, RNAi studies in the related kinetoplastid parasite *T. brucei* suggested that *TbErv* is essential [44,45,65]. Nevertheless, we are aware that viable knockout strains with a drastic growth defect can be missed without negative selection and that the recent establishment of CRISPR/Cas9 systems in *Leishmania* [66–68] could be extremely helpful to get a definite answer regarding the essentiality of *Erv* and *sTim1* in *Leishmania* pathogens.

Previous studies on *Erv* homologs from human, yeast, *L. tarentolae*, *T. brucei*, and *Arabidopsis thaliana* revealed that the redox-active shuttle cysteines are present in highly variable motifs, which are sometimes close to the N- or the C-terminus [30,37,43,44,69–73]. Wild-type *Erv* homologs from *A. thaliana* and *L. tarentolae* both carry a C-terminal shuttle arm and were unable to complement the loss of ScErv1 [43,72]. Even though complementation with *LtErv*^{C17S} resulted in a growth defect on FOA agar plates, the viability of the strain clearly shows that the position of the shuttle arm is not a criterion for exclusion for the functionality in yeast. This is also the case regarding the CRSC shuttle arm motif, which was successfully replaced in chimera 6, although at the cost of a drastically impaired mitochondrial protein import. Residue Cys17 of *LtErv* is partially conserved in *Erv* homologs [16,43] and was shown to form a clamp disulfide bond in mammalian ALR [33,37,74]. Notably, heterologous human ALR was found in the yeast cytosol and was unable to functionally replace ScErv1 unless a bipartite pre-sequence was fused to its N-terminus or residues 1–80 of ALR were replaced with residues 1–93 of ScErv1 [75,76]. How residue Cys17 of *LtErv* exactly impairs its function in yeast and whether this neglected residue exerts a physiological function in a subset of *Erv* homologs remains to be studied. Chimera 1 and 4 both contain an N- and C-terminal shuttle arm, but only yeast cells with chimera 1 had a growth defect. Maybe both shuttle arms compete for the proximal cysteines of the flavodomain of chimera 1 or form long-lived disulfide-bonded intermediates in contrast to chimera 4. In summary, growth and import defects for chimera 1, 2 and 6 suggest that the shuttle arm motif and the interactions between the shuttle arm and the flavodomain and/or Mia40 have been optimized for different *Erv* homologs in the course of evolution. Nevertheless, the mitochondrial protein import machinery in yeast is robust enough to allow certain degrees of incompatibility.

Can protist *Erv* homologs act alone and do protists have simplified oxidative folding machineries mediating the IMS protein import as recently suggested [65,77]? This possibility would require an efficient interaction of *Leishmania* class II IMS proteins with *LtErv*, including their direct oxidation by *LtErv*. In this case one might expect that *LtErv* is able to complement the function of ScMia40, because the import signals of IMS proteins are functionally conserved in the course of evolution [42]. Our complementation studies do not provide evidence for a simplified oxidation and folding machinery, because neither the KISS domain nor full length *LtErv*, *LtErv*^{C17S}, *LtErv*^{C-term} or chimera 1–6 could rescue the Δ *mia40* strain. This rather points to the presence of a

Mia40 replacement in protists (compound Z in one of our previously discussed evolutionary scenarios [42]). Such a model suggests that protists without Mia40 have a replacement for this substrate receptor. So far we were unable to identify a stable disulfide-bonded interaction partner between *LtErv* and another protein in *Leishmania* mitochondria using a variety of experimental approaches (Liedgens et al., unpublished). However, just because redox mobility shift assays and pull-down experiments do not reveal a stable interaction, we cannot exclude a transient or non-covalent interaction with a Mia40 replacement. The "sluggish" behavior of ScMia40 with its metastable intermolecular disulfide species is actually rather an exception compared to oxidative protein folding in the periplasm or the endoplasmic reticulum [7,31]. What could be a reasonable replacement of Mia40? A recent study showed that a subpopulation of the yeast peroxidase GPx3 interacts with ScMia40 and the oxidative folding pathway in the IMS [78]. This situation somehow resembles the transient interaction between peroxiredoxin IV and protein disulfide isomerase (PDI) for oxidative protein folding in the endoplasmic reticulum of mammals [79,80]. A trypanredoxin peroxidase (Tb927.9.5750) was also identified in a pull-down assay with *TbErv* [65], and this protein was 1.34-fold up-regulated in the *TbErv* knockdown strain [45]. Whether this is just a false positive hit due to the high abundance of the cytosolic peroxidase or points to a dual localization and alternative function in oxidative protein folding remains to be studied. However, GPx3 and peroxiredoxin IV are both downstream electron acceptors of Mia40 and PDI so that an analogous involvement of the trypanredoxin peroxidase in the IMS of kinetoplastid parasites would still necessitate the existence of an alternative receptor for the incoming protein substrates. Further studies are obviously necessary to identify the receptor for incoming IMS proteins and to decipher the oxidative folding machineries in protist mitochondria.

Acknowledgements

This work was supported by the Deutsche Forschungsgemeinschaft (Grants DE 1431/10-1 and HE 3462/4-1) and the Norte-01–0145-FEDER-000012-Structured program on bioengineered therapies for infectious diseases and tissue regeneration, supported by Norte Portugal Regional Operational Programme (NORTE 2020), under the PORTUGAL 2020 Partnership Agreement, through the European Regional Development Fund (FEDER). The position of M.D. was funded by the Deutsche Forschungsgemeinschaft in the frame of the Heisenberg program (Grant DE 1431/9-1). We thank Johannes Herrmann for the complemented Δ *mia40* strain, the Bukau laboratory for antibody α -ScZwf1, and Andreas Ladurner for his generous support. We are also grateful to Suzan Can and Stefan Wölfl for their support with the Tecan Infinite F200 Pro plate reader measurements.

Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at <http://dx.doi.org/10.1016/j.redox.2017.12.010>.

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