Generating disulfides enzymatically: Reaction products and electron acceptors of the endoplasmic reticulum thiol oxidase Ero1p


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Ero1p is a key enzyme in the disulfide bond formation pathway in eukaryotic cells in both aerobic and anaerobic environments. It was previously demonstrated that Ero1p can transfer electrons from thiols to molecular oxygen. However, the fate of electrons under anaerobic conditions and the final fate of electrons under aerobic conditions remained obscure. To address these fundamental issues in the Ero1p mechanism, we studied the transfer of electrons from recombinant yeast Ero1p to various electron acceptors. Under aerobic conditions, reduction of molecular oxygen by Ero1p yielded stoichiometric hydrogen peroxide. Remarkably, we found that reduced Ero1p can transfer electrons to a variety of small and macromolecular electron acceptors in addition to molecular oxygen. In particular, Ero1p can catalyze reduction of exogenous FAD in solution. Free FAD is not required for the catalysis of dithiol oxidation by Ero1p, but it is sufficient to drive disulfide bond formation under anaerobic conditions. These findings provide insight into mechanisms for regenerating oxidized Ero1p and maintaining disulfide bond formation under anaerobic conditions in the endoplasmic reticulum.

It is likely that the basic reaction scheme described above for the QSOX/ALR family applies to Ero1p as well. First, the similar arrangements of the functional groups in the active sites of Ero1p and Erv2p (4) suggest that the two thiol oxidase families may share features of their catalytic mechanisms in common. Second, depletion of oxygen in the solution, which should disable the oxidative half-reaction in the above scheme, traps the Ero1p-bound FAD in its reduced state (4). Third, as shown herein, stoichiometric transfer of electrons from thiol substrates to Ero1p occurs under such conditions.

In this study, we focus primarily on the oxidative half-reaction (Eq. 2) to address two questions regarding the mechanism of Ero1p. The first concerns the product of the reaction when molecular oxygen is the direct oxidative substrate of Ero1p. A two-electron reduction of molecular oxygen is expected to yield hydrogen peroxide, and the potential for Ero1p-mediated thiol oxidation to contribute to the pool of reactive oxygen species in the cell has been appreciated (12). However, only low levels of hydrogen peroxide were detected in vitro as a result of Ero1p activity (3). Because any peroxide produced can potentially go on to react with electron donors present in the solution, we sought assay conditions to maximize peroxide recovery or detection.

The second question we addressed is whether electron acceptors other than oxygen can reactoxidize Ero1p. It was reported that the addition of free FAD, in excess of the FAD bound by the purified enzyme, was essential for RNaseA refolding in vitro catalyzed by Ero1p and protein disulfide isomerase (3). One hypothesis explaining this observation was that Ero1p has a “sensor” site for exogenous FAD, which may function as an allosteric effector to modulate disulfide bond formation according to the cellular metabolic status (3). To investigate the possibility of another role for FAD in the Ero1p-mediated reaction, we tested whether free FAD could be reduced directly by FADH₂ of the Ero1p holoenzyme as an alternative to the reduction of molecular oxygen at the enzyme active site. We then went on to determine whether nonflavin compounds could also serve as electron acceptors under anaerobic conditions.

In live yeast cells, the ero1-1 mutation causes temperature sensitivity under either aerobic or anaerobic conditions (8, 13). This observation suggests that Ero1p is required for disulfide bond formation in both the presence and the absence of oxygen and that the enzyme has the ability to transfer electrons to acceptors other than oxygen in vivo. In this article, we demonstrate that various routes for regeneration of an oxidized Ero1p

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Abbreviations: ER, endoplasmic reticulum; CTAB, cetyltrimethylammonium bromide; Trx₄₉₉, reduced Trx1.

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protein consists of GST, a thrombin cleavage site, a His6-tag, and the cysteine-rich domain of \textit{Saccharomyces cerevisiae} Ero1p were produced as described in ref. 4, except that inducible expression was done at 15°C for 48 h. The fusion fragment is 45,294 Da, taking into account the loss of two protons for every disulfide bond formed. Unless otherwise noted, the data shown here were obtained by using this version of Ero1p. A second construct, spanning amino acid residues 10–424, was also produced and tested. The relevant segment of the Ero1p coding sequence was inserted into the vector. The molecular mass measured by mass spectrometry was calculated to be 14.4 Da at 446 nm for free FAD (16). The cofactor/enzyme ratio was determined by measuring absorbance in 6 M guanidine/20 mM NaPO4 (pH 6.5) at 446 nm (e = 11.3 M\(^{-1}\)cm\(^{-1}\)) and 280 nm (calculated e = 79.1 M\(^{-1}\)cm\(^{-1}\)) (17), which includes the contribution of the FAD moiety at this wavelength: e = 21.3 M\(^{-1}\)cm\(^{-1}\)). The concentration of protein was thus measured to be 14.4 \(\mu\text{M}\), and the concentration of FAD was measured to be 13.9 \(\mu\text{M}\), a ratio of 1.04.

An expression vector for \textit{E. coli} thioredoxin I (Trxl) was constructed by amplifying the Trxl coding sequence from bacterial lysate and inserting it between the NdeI and BamHI sites in the pET-28 vector (Novagen). Protein was expressed in BL21(DE3)pLysS (Novagen) bacterial cells grown at 37°C to 0.7 OD units and induced with 0.5 mM isopropyl \(\beta\)-D-thiogalactopyranoside. After 3 h, cells were harvested, lysed by sonication in 50 mM NaPO4 (pH 8.0)/300 mM NaCl/10 mM imidazole, and centrifuged to remove insoluble material. Triton X-100 was added to a concentration of 1% to the supernatant, which was then applied to a Ni-NTA column. The column was washed first with 50 mM NaPO4 (pH 8.0)/300 mM NaCl/20 mM imidazole/20% glycerol to remove the Triton X-100, and washed again with the same solution lacking glycerol. Protein was eluted with a step gradient of increasing imidazole. DTT was added to the eluted Trxl to a concentration of 100 mM, and the protein was incubated for 1 h at room temperature. The reduced Trxl (0.5–0.7 \(\mu\text{M}\)) was applied to a PD-10 column equilibrated with buffer A. Fractions were collected and tested qualitatively for thiol content by using Ellman’s assay (18). A clear separation was seen between the peak corresponding to reduced Trxl (Trxl\text{red}) and that of DTT. Trxl\text{red} peak fractions were pooled, and the reported Trxl\text{red} concentrations were determined quantitatively by Ellman’s assay. All oxygen consumption and spectroscopic
assays were performed within 6 h of measuring the thiol concentration of Trx\textsubscript{red}.

Bovine liver catalase and horse heart cytochrome \( c \) were purchased from Sigma. The concentration of cytochrome \( c \) was determined by subtracting the absorbance at 550 nm of oxidized protein from that of dithionite-reduced protein (\( \Delta \epsilon = 21.1 \text{mM}^{-1}\text{cm}^{-1} \)). \( S. \text{cerevisiae} \) cytochrome \( b5 \) was amplified by PCR from yeast genomic DNA, inserted between the NdeI and BamHI sites in the pET-15 vector (Novagen), and expressed and purified by using the procedure for Trx\(i \). \textit{Pseudomonas aeruginosa} azurin was a gift of Israel Pecht (The Weizmann Institute of Science). Its concentration was determined at 630 nm (\( \epsilon = 5.6 \text{mM}^{-1}\text{cm}^{-1} \)).

**Anaerobic Titration.** Ero1p (7.4 \( \mu \text{M} \)) in buffer A was deoxygenated in the main space of an anaerobic cuvette by repeated cycles of vacuum and flushing with nitrogen (19). A solution of Trx\textsubscript{red} (0.65 mM in buffer A) was deoxygenated by cycles of degassing and flushing with nitrogen for \( > 1 \) h and then used to fill a 50-\( \mu \text{l} \) gas-tight syringe (19). The syringe was loaded onto a ground-glass flotation chamber under a gentle stream of nitrogen gas. The apparatus was designed to prevent air from contaminating the solution. The syringe was then placed within a 5-mI anaerobic chamber under a gentle stream of nitrogen gas. The syringe was then placed within a 5-mI anaerobic chamber under a gentle stream of nitrogen gas. The syringe was then placed within a 5-mI anaerobic chamber under a gentle stream of nitrogen gas.

**Oxygen Consumption Assays.** Oxygen consumption was measured by using an Oxygraph Clark-type oxygen electrode (Hansatech Instruments, Pentney King’s Lynn, U.K.). FAD, FMN, and riboflavin concentrations were determined by absorbance at 446 nm in 6 M guanidine/20 mM NaPO\( 4 \) (pH 6.5) by using extinction coefficients of 11.3, 12.5, and 12.2 \( \text{mM}^{-1}\text{cm}^{-1} \), respectively. The oxygen electrode chamber was covered with aluminum foil during reactions to avoid photoreduction of the flavins. DTT solutions were prepared by weight and then corrected by using Ellman’s assay. All experiments were performed in buffer A unless otherwise noted. All components of each reaction, except Ero1p, were mixed fresh in a total volume of 1.0 ml. Oxygen levels were monitored until a linear baseline was established, and the reaction was initiated by injection of Ero1p, typically in a volume of 20 \( \mu \text{l} \), to yield a final enzyme concentration of 1 \( \mu \text{M} \). For determination of the apparent \( K_m \) for oxygen by using Trx\textsubscript{red} as the reducing substrate, the Trx\textsubscript{red} solution was stirred in the oxygen electrode chamber under a gentle stream of nitrogen until the oxygen concentration dropped to 30 \( \mu \text{M} \) (\( \sim 15 \) min) before Ero1p was injected.

**Hydrogen Peroxide Assays.** Hydrogen peroxide production was detected by monitoring oxygen evolution upon addition of catalase. After the thiol oxidation reaction was complete, 10 \( \mu \text{l} \) of a 2-mg/ml stock of catalase was injected into the reaction vessel of the oxygen electrode. In parallel, hydrogen peroxide production was followed kinetically by using the PeroXOQuant quantitative peroxide assay kit (Pierce). \( \text{H}_2\text{O}_2 \) standards were made by using an \( \sim 30\% \text{ H}_2\text{O}_2 \) stock solution (10.8 M). The stock solution concentration was standardized by absorbance at 240 nm of a 1:1,000 dilution by using an extinction coefficient of 43.6 \( \text{M}^{-1}\text{cm}^{-1} \). All PeroXOQuant measurements, including the standard curve, were made in buffer A lacking EDTA. The standard curve was derived from the following \( \text{H}_2\text{O}_2 \) concentrations: 0, 1, 5, and 10–100 \( \mu \text{M} \) in intervals of 10 \( \mu \text{M} \). The plot of absorbance vs. \( \text{H}_2\text{O}_2 \) concentration was linear in this range. For this assay, Trx\textsubscript{red} was prepared as described above except that after reduction, the protein was exchanged into buffer A lacking EDTA and used within 2 h. Ero1p was dialyzed against buffer A lacking EDTA. The enzymatic reaction was performed in a total volume of 750 \( \mu \text{l} \) while being stirred in a glass vial. The concentration of Trx\textsubscript{red} was 100 \( \mu \text{M} \), and the enzyme concentration was 1 \( \mu \text{M} \).

**Results**

**Enzyme and Substrates.** For enzymatic assay of Ero1p, we used forms of the \( S. \text{cerevisiae} \) enzyme that could be efficiently expressed as recombinant proteins, comprising Ero1p residues 56–424 (shorter construct) or 10–424 (longer construct). The full-length yeast enzyme is 563 aa in length, but the carboxy terminus of our fragments corresponds to that of other Ero1p orthologs (i.e., human), which lack the \( \sim 130\)-aa extension found in yeast Ero1p (2). Enzyme was produced in the Origami B(DE3)pLysS \( E. \text{coli} \) bacterial strain as detailed in an earlier article describing the high-resolution structure of Ero1p (4) and in \textit{Materials and Methods}. Previous biochemical experiments on full-length \( S. \text{cerevisiae} \) Ero1p purified from yeast were conducted by using enzyme preparations that bound the FAD cofactor substoichiometrically (13). In contrast, our preparations of Ero1p contained equimolar amounts of protein and FAD. The extinction coefficient for FAD bound to Ero1p is 12.5 \( \text{mM}^{-1}\text{cm}^{-1} \) at 454 nm (Fig. 2).

Enzyme activity was directly monitored by oxygen consumption or reduction of the substrate of the second half-reaction (Eq. 2). Ero1p can oxidize a variety of small and macromolecular substrates (data not shown). In all cases, a lag phase is observed in the progress curves. Because the lag can be eliminated by preincubation of the enzyme with substrate but not by incubation with reaction products (data not shown), we attribute the lag to reductive activation of the enzyme. We found that Ero1p has a higher turnover number for \( E. \text{coli} \) Trx\textsubscript{red} than for DTT at equivalent substrate concentrations. For assays in which the reducing substrate would be stoichiometrically limiting, we therefore used Trx\textsubscript{red}. Although Trx\textsubscript{red} is a nonphysiological substrate, its reaction rate was suitable, and the protein could be produced easily in large quantities.

**Reductive Titration of Ero1p with Trx\textsubscript{red}**. Fig. 3 shows a titration of Ero1p with Trx\textsubscript{red} under anaerobic conditions. Substantially
complete reduction of the enzyme-bound flavin required approximately four electrons, indicating that at least two molecules of Trxred can be oxidized by Ero1p in this static experiment. Similar results were obtained previously for egg white sulfhydryl oxidase, which required ~2 eq of DTT (four electrons) to completely reduce the flavin (20). In both cases, the additional two electrons are taken up by disulfides. The Ero1p enzyme used in this study (shorter construct) contains five disulfide bonds in total, so it is possible that additional electrons may be transferred from Trxred to Ero1p after the bound flavin is reduced, although these later events would be largely silent spectroscopically. There is a noticeable lag in Fig. 3 Inset, consistent with the involvement of additional redox centers beside the flavin in the early stages of the experiment. During the titration, a rise in long-wavelength absorbance, exhibiting distinct shoulders at 580 and 630 nm, is observed. These features indicate the generation of the blue neutral flavin radical. Much higher levels of this semiquinone species are observed in another flavin-linked sulfhydryl oxidase, the human augmenter of liver regeneration, upon the addition of DTT (21).

Hydrogen Peroxide Is a Product of Ero1p-Catalyzed Thiol Oxidation. Previous studies reported only low levels of hydrogen peroxide formation during catalysis by Ero1p (3), although typical FAD-binding oxidases produce hydrogen peroxide (22). Because the generation of reactive oxygen species during oxidative folding in the ER is of considerable potential significance (12, 14, 15), we re-investigated this issue by using two independent methods. Trxred was chosen as the thiol substrate because it is rapidly and completely oxidized at low concentrations. Fig. 4A illustrates oxygen consumption during Ero1p-mediated oxidation of 50 μM Trxred. After a drop of 49 μM oxygen, the addition of catalase restored 23 μM oxygen to the solution, indicating that 46 μM hydrogen peroxide [stoichiometry 2H2O2 ↔ H2O + O2; (reviewed in ref. 23)] had been present. Thus, the generation of one disulfide yielded 0.94 molecules of hydrogen peroxide.

As an independent measure of peroxide formation, aliquots from a Trxred oxidation reaction were mixed with a colorimetric reagent for the quantitation of hydrogen peroxide (Fig. 4B). Because EDTA interferes with the development of the Fe(III)-xylenol orange chromagen, it was eliminated from the reaction. The absence of EDTA leads to a decreased peroxide yield (~0.4 molecules per disulfide formed). A correspondingly lower recovery of hydrogen peroxide was obtained when EDTA was omitted from the experiment in Fig. 4A (data not shown). Although the peroxide yield is reduced, the lag phase in Fig. 4B mirrors that seen in the oxygen consumption curve obtained in Fig. 4A. This lag reflects the reductive activation of Ero1p; it is not seen when the Erv2p enzyme is assayed by using DTT as the thiol substrate, in which case hydrogen peroxide is produced with single-exponential kinetics (data not shown).

*K*m Value for Ero1p Reduction of Oxygen. Once we determined that hydrogen peroxide is the product of oxygen reduction during Ero1p-mediated thiol oxidation, we sought to determine the *K*m value for oxygen. The use of 12.5 mM DTT such that the thiol substrate concentration remained effectively unchanged upon oxygen depletion yielded an approximate *K*m value of 4 μM for oxygen (Fig. 5). Similar values were obtained for the long and short Ero1p constructs. Because the rate of the reaction is initially determined by the nature of the reducing substrate, the apparent *K*m value for oxygen will be different for different reductants. Using the longer Ero1p construct, we performed a similar experiment with 300 μM Trxred as the substrate of the reductive half-reaction and an initial oxygen concentration of around 30 μM such that the Trxred would be depleted by 10% during the course of the reaction. Under these conditions, the *K*m value of Ero1p for oxygen was found to be ~10 μM (data not shown). These low *K*m values suggest that Ero1p can function efficiently at low oxygen tension.

Fig. 3. Ero1p (7.4 μM in buffer A) was titrated anaerobically with Trxred added from a gas-tight syringe. Inset is a plot of absorbance at 454 nm recorded after each addition. Selected spectra from the titration (curves 1–7, corresponding to 0.0, 0.5, 1.0, 1.5, 2.0, 2.5, and 2.9 eq of Trxred per mol of flavin) are plotted in the main panel.

Fig. 4. Oxygen is reduced by Ero1p activity to form hydrogen peroxide. (A) The reaction was initiated at time 0 by injecting the shorter construct of Ero1p to a final concentration of 1 μM into a solution containing 50 μM Trxred and oxygen consumption was monitored. Approximately half of the oxygen originally consumed during Trxred oxidation was restored with the addition of catalase (arrow), indicating that oxygen has been reduced to H2O2 in the initial reaction. Similar results were obtained for the longer construct. (B) The reaction was initiated at time 0 by injecting the shorter form of Ero1p to a final concentration of 1 μM into a rapidly stirred solution containing 100 μM Trxred. Samples were withdrawn and mixed with the PeroxOquant detection reagent. Absorbance values were converted to H2O2 concentration according to a standard H2O2 calibration curve.
Effect of FAD on the Rates of Reactions Catalyzed by Ero1p. Because of previous reports that the addition of FAD to the enzyme solution increased activity in an RNase refolding assay (3), we tested the effect of FAD on the rate of oxygen consumption. We found that 100 μM FAD did not affect the oxygen consumption rate during Ero1p-catalyzed oxidation of either Trxred (Fig. 6A) or DTT (Fig. 6B). Similar results were obtained for 100 μM FMN and 100 μM riboflavin (Fig. 6). Because riboflavin and FMN are readily photo-reduced (24), all reactions were performed in the dark. Results were qualitatively similar for both Ero1p constructs and are shown for the longer version.

In the presence of Trxred, Ero1p catalyzed the transfer of reducing equivalents to both free FAD and FMN (Fig. 7). Experiments were performed anaerobically and monitored by a decrease in absorbance at 446 nm. The concentration of free flavin in these experiments was at least 20-fold greater than the decrease in absorbance at 446 nm. The concentration of free flavin was depleted to <10% of its initial value. Inset is the first derivative of the smoothed data, which was used to identify the point at which the rate was half maximal.

Ero1p Transfer Electrons to Nonflavin Exogenous Acceptors. In addition to reducing free flavin in solution, Ero1p could also catalyze transfer of electrons to nonflavin acceptors. Although many potential small-molecule electron acceptors had to be excluded from our studies because of high rates of nonenzymatic reduction by the thiol substrates, certain metalloproteins [i.e., horse cytochrome c, yeast cytochrome b5 (data not shown), and bacterial azurin] exhibited sufficiently slow noncatalytic reduction that they could be tested in our system. These proteins were reduced catalytically in the presence of Ero1p with Trxred as the reducing substrate (Fig. 8). Regardless of the electron acceptor, its reduction was detected only after a lag, similar to the kinetics observed for consumption of oxygen, when present (Fig. 6). The abrupt decrease in reaction rate as the concentrations of azurin or cytochrome c approached zero indicates qualitatively that the $K_m$ values for these proteins are very low.

Discussion

In this study, we demonstrate that Ero1p generates hydrogen peroxide when acting on thiol substrates in the presence of oxygen, as expected of a typical flavin-containing oxidase (22). This observation contrasts with a reported stoichiometry of ~0.05 hydrogen peroxide molecules per disulfide bond introduced into reduced RNase (3). Although differences in the Ero1p constructs could conceivably influence the reaction stoichiometry, we note that hydrogen peroxide production is frequently underestimated because it can decompose in the presence of trace metals or can react with thiols (25). In addition, thiols interfere with quantitation of hydrogen peroxide by peroxidase-mediated chromagen formation (26). The low $K_m$ value of Ero1p for oxygen suggests that the reduction of oxygen to hydrogen peroxide is a relevant route for electron transfer during Ero1p-catalyzed disulfide formation, except under severe hypoxic conditions. That reactive and potentially damaging

**Fig. 5.** Determination of the $K_m$ value for oxygen. Ero1p (shorter construct) was injected into a solution containing 12.5 mM DTT in the oxygen electrode chamber. The graph shows the end of the reaction, when the oxygen has been depleted to <10% of its initial value. Inset is the first derivative of the smoothed data, which was used to identify the point at which the rate was half maximal.

**Fig. 6.** Effect of exogenous flavin compounds on the rate of oxygen consumption during Ero1p catalysis of disulfide formation. The experiments shown were performed by using the longer Ero1p construct, but the shorter version gave similar results. At time 0, Ero1p was injected to a final concentration of 1 μM into a solution containing 98 μM Trxred (A) or 10 mM DTT (B) to obtain the curves indicated by filled symbols. Traces measured in the absence of enzyme are indicated by open symbols. The black curves show the kinetics of oxygen consumption without added flavin compounds. The red, orange, and gold curves show the kinetics of oxygen consumption under similar conditions, except that the indicated flavin compounds were present in the reaction (100 μM FAD, 100 μM FMN, and 100 μM riboflavin, respectively).

**Fig. 7.** Reduction of flavins in the absence of oxygen. The reductant was 50 μM Trxred and the electron acceptors were 21 μM FAD or 30 μM FMN. When present, the shorter construct of Ero1p was at a concentration of 1 μM. The longer version of Ero1p also reduces exogenous flavins (data not shown).
redox species are the outcome of catalysis of disulfide bond formation may explain why induction of Ero1p by the ER stress response pathway leads to the production of excess peroxides in Caenorhabditis elegans (14).

In the absence of oxygen, however, we find that flavin compounds can also serve as electron acceptors for Ero1p. This reduction of free flavin by Ero1p cannot be the result of an exchange between bound and free flavin because, under the conditions of the assay, no loss of the stoichiometrically bound FAD cofactor could be detected in dialysis measurements (data not shown). Furthermore, Erv2p, an enzyme with no sequence homology to Ero1p but containing a similar core structure and arrangement of functional groups, also catalyzes reduction of free flavins (data not shown). Precedents for reduction of exogenous flavins by flavoproteins exist, including the reductase homology to Ero1p but containing a similar core structure and FAD cofactor could be detected in dialysis measurements (data not shown). The ability of flavins to function as electron acceptors for Ero1p may be shared by other chromophores and does not necessarily indicate a specific role for free flavins in thiol oxidation by this enzyme. To resolve this issue, two questions need to be addressed in future work. The first question is whether free flavin is indeed present at low micromolar concentrations in the ER (3) and, if so, whether it is a kinetically competitive oxidant of Ero1p under such conditions. The second question is how reduced free flavin might be reoxidized during oxidative folding under anaerobic conditions.

Experiments showing that Ero1p activity in vivo is tied to the amount of free intracellular FAD have been interpreted as evidence that Ero1p activity can be allosterically controlled by FAD (3). However, the importance of intracellular FAD levels is also consistent with a role for FAD as a cofactor and/or possible electron acceptor for Ero1p. An explanation for the synthet lethality of the ero1-1 and fad1-1 mutants (3) may be that the folding and, hence, active levels of the temperature-sensitive Ero1p variant are particularly sensitive to the amount of FAD available during its biosynthesis. The Gly-to-Ser mutation at position 229 encoded by ero1-1 is adjacent to the FAD-binding site and is likely to affect the coupling of folding and cofactor binding.

Our experiments also raise the possibility that nonflavin molecules may be on the pathway for Ero1p catalysis of disulfide bond formation under anaerobic conditions. Two heme-binding proteins and a protein with a copper center all served as efficient electron acceptors from Ero1p under anaerobic conditions in vitro. It was recently shown that ALR, a mammalian homolog of Erv2p found in the mitochondrial intermembrane space, can use cytochrome c as an electron acceptor (21). In this article, we demonstrate that an ER thiol oxidase can also transfer electrons to cytochrome c and to other proteins that function in electron transport chains. One significant difference between Ero1p and ALR, however, is that ALR uses molecular oxygen as an electron acceptor relatively poorly, ~100 times less efficiently than it uses cytochrome c (21), whereas Ero1p can use both oxygen and other electron acceptors efficiently. Although the metalloproteins tested in this study are not known to function in the lumen of the ER, the capacity of Ero1p to use diverse electron acceptors leads us now to consider a variety of redox active molecules in the search for the physiological electron acceptors for Ero1p.

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