Cysteines 208 and 241 in Ero1α are required for maximal catalytic turnover

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A R T I C L E   I N F O

Article history:
Received 10 October 2015
Received in revised form
11 November 2015
Accepted 12 November 2015
Available online 14 November 2015

Keywords:
Ero1α
PDI
Disulfide bond
Redox homeostasis
Endoplasmic reticulum

A B S T R A C T

Endoplasmic reticulum (ER) oxidoreductin 1α (Ero1α) is a disulfide producer in the ER of mammalian cells. Besides four catalytic cysteines (Cys94, Cys99, Cys394, Cys397), Ero1α harbors four regulatory cysteines (Cys104, Cys131, Cys208, Cys241). These cysteines mediate the formation of inhibitory intramolecular disulfide bonds, which adapt the activation state of the enzyme to the redox environment in the ER through feedback signaling. Accordingly, disulfide production by Ero1α is accelerated by reducing conditions, which minimize the formation of inhibitory disulfides, or by mutations of regulatory cysteines. Here we report that reductive stimulation enhances Ero1α activity more potently than the mutation of cysteines. Specifically, mutation of Cys208/Cys241 does not mechanistically mimic reductive stimulation, as it lowers the turnover rate of Ero1α in presence of a reducing agent. The Cys208/Cys241 pair therefore fulfills a function during catalysis that reaches beyond negative regulation. In agreement, we identify a reciprocal crosstalk between the stabilities of the Cys208–Cys241 disulfide and the inhibitory disulfide bonds involving Cys104 and Cys131, which also controls the recruitment of the H2O2 scavenger GPx8 to Ero1α. Two possible mechanisms by which thiol-disulfide exchange at the Cys208/Cys241 pair stimulates the catalytic turnover under reducing conditions are discussed.

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1. Introduction

The synthesis of extracellular proteins is initiated at the endoplasmic reticulum (ER) where translating ribosomes translocate the nascent polypeptides into the ER lumen. Many of these polypeptides subsequently acquire critical covalent linkages between cysteine residues (termed disulfide bonds) through thiol-disulfide exchange reactions. Among the different systems for disulfide bond formation in the ER, ER oxidoreductin 1 (Ero1)-catalyzed oxidation of the active-site cysteine pair in protein disulfide isomerase (PDI) constitutes the best-conserved pathway [1,2]. Ero1 in vertebrates exists in two isoforms, Ero1α and Ero1β, whereas Ero1α is ubiquitously expressed and being viewed as the major source of disulfides in humans [3].

The flavoprotein Ero1α is an oxidase that couples the reduction of molecular oxygen (O2) to disulfide-bond formation [3]. In the reductive phase of the Ero1α catalytic cycle, the flavin adenine dinucleotide (FAD) cofactor in Ero1α is reduced to FADH2 as a result of disulfide transfer from Ero1α to reduced PDI (PDIred). The reaction is catalyzed by an outer active-site cysteine pair (Cys54/Cys59), which shuttles two electrons from PDIred via an inner active-site cysteine pair (Cys394/Cys397) to FAD. This process is tightly regulated by the reversible formation of two inhibitory disulfides (Cys94–Cys131 and Cys99–Cys104) [4–6]. In the oxidative phase, FAD is regenerated by the transfer of two electrons onto O2, which leads to the formation of one molecule of hydrogen peroxide (H2O2) [7]. Recent evidence indicates that H2O2 is instantaneously reduced to H2O by the Ero1α-associated glutathione peroxidase family enzymes GPx7 or GPx8, which also introduce the resulting second disulfide into PDI [8,9]. We showed that access of O2 to the buried FAD molecule is negatively regulated by the Cys208–Cys241 disulfide [10].

The thiol-disulfide statuses of all inhibitory disulfide bonds in Ero1α are governed by canonical PDI or other PDI family members [10,11]. Accordingly, regulatory cysteines fine-tune the activation state of Ero1α in a redox environment-dependent manner by blocking either the reductive (Cys104 and Cys131) or the oxidative (Cys208 and Cys241) phase of the catalytic cycle through the formation of feedback-regulated inhibitory disulfides in response to oxidizing conditions. Conversely, a reducing ER environment promotes...
Ero1α activation through PDI-catalyzed reduction of these disulfides, as could for instance be important in response to physiological hypoxia or peak concentrations of reduced glutathione (GSH) or redox-active vitamins.

Here, we report that reducing conditions, which activate Ero1α through the removal of inhibitory disulfides, more potently stimulate Ero1α activity than the mutation of all regulatory cysteines. Thus, the presence of the Cys208/Cys241 pair is required for maximal catalytic turnover under reducing conditions. Our data indicate a new mechanism of Ero1α regulation in which thiol-disulfide exchange at Cys208–Cys241 affects the stability of the Cys94–Cys131 inhibitory disulfide through allosteric and/or intermolecular communication.

2. Materials and methods

2.1. Fluorescence excitation spectrum analysis

Cells stably transfected with HyPerER were subjected to fluorescence excitation spectrum analysis as described before [12].

2.2. Dithiothreitol (DTT) washout assays

The cellular glutathione disulfide:total glutathione (GSSG:GS(τ)) ratio after DTT washout was measured using a 5,5′-dithiobis(2-nitrobenzoic acid)/glutathione reductase recycling assay as previously described [13].

2.3. Statistics

Data sets were analyzed for statistical significance using Student’s t test (two-tailed distribution; heteroscedastic).

2.4. Cell culture and transient transfections

The culturing of HeLa cells [14] and FlipIn TRex293 cells for doxycycline (1 μg/ml, Sigma)-inducible expression of Ero1 variants [4] has been described. The following FlipIn TRex293 cell lines have been published previously: Ero1α [4], Ero1α-AA [6], Ero1α-C208S/C241S, Ero1α-AA:HyPerER [8], Ero1α-C208S/C241S:HyPerER [10] and Ero1α-AASS:HyPerER [10]. The Ero1α−WT:HyPerER cell line was created as before [8] (with the HyPerER vector kindly provided by Miklos Geiszt, Semmelweis University, Hungary).

Transient transfections of HeLa cells were carried out using Turbofect (Thermo Scientific). Transient transfections of FlipIn TRex293 cells were carried out using Metafectene Pro (Biontex).
2.5. Oxygen and NADPH consumption assay

Oxygen consumption was measured as previously described [15]. Experimental procedure of NADPH consumption assay has been described before [16,17].

2.6. Determination of the redox equilibrium constant (\(K_{eq}\))

\(K_{eq}\) values of Cys\(^{94}\)-Cys\(^{131}\) in Ero1\(\alpha\)-WT and Ero1\(\alpha\)-C208S/C241S were determined as follows. Ero1\(\alpha\)-WT or Ero1\(\alpha\)-C208S/C241S (2 \(\mu\)M) was incubated for 30 min at 30 \(^\circ\)C in degassed 50 mM Tris/HCl (pH 7.5) buffer containing 300 mM NaCl, 0.2 mM GSSG and various concentrations of GSH (0.2–4 mM). After incubation, 1 mM NEM was added to avoid subsequent redox reactions. All the samples were separated by non-reducing SDS-PAGE and stained with Coomassie brilliant blue (CBB). The values of fraction of OX2 state of Ero1\(\alpha\) were estimated by LAS-3000 image reader, and plotted against [GSH]/[GSSG] ratios. \(K_{eq}\) values were determined by fitting the data according to equation:

\[ R = \frac{(\text{GSH}^2/\text{GSSG})}{(\text{GSH}^2/\text{GSSG})} + (\text{GSH}^2/\text{GSSG}) \]

where \(R\) is fraction of OX2 state of Ero1\(\alpha\) at equilibrium.

2.7. Bimolecular fluorescence complementation (BiFC) assay

HeLa cells were transfected and analyzed as previously described [10]. CRTss+EYFP1 + mature Ero1\(\alpha\) in pcDNA3.1 (dubbed EYFP1–Ero1\(\alpha\)-WT) and CRTss+EYFP2+luminal domain GPx8 in pcDNA3.1 were kindly provided by Lloyd Ruddock [18]. EYFP1–Ero1\(\alpha\)-AA was generated by site-directed mutagenesis.

3. Results and discussion

Recent work has demonstrated that Ero1\(\alpha\)-AASS, an Ero1\(\alpha\) variant with all four regulatory cysteines being mutated (C104A, C131A, C208S, C241S), displays higher oxidase activity than Ero1\(\alpha\)-AA where only Cys104 and Cys131 are mutated [10]. A likely mechanism was presented in which PDI\(_{\text{red}}\) facilitates the reaction of FADH\(_2\) with O\(_2\) in Ero1\(\alpha\)-AA by opening the intramolecular Cys208–Cys241 disulfide [10]. Since PDI\(_{\text{red}}\) is depleted through the oxidation by Ero1\(\alpha\)-AA and its regeneration from oxidized PDI (PDI\(_{\text{ox}}\)) is relatively inefficient in the ER at steady state, the reaction rate is limited by the availability of PDI\(_{\text{red}}\). Accordingly, Ero1\(\alpha\)-AASS, which lacks Cys\(^{208}\)–Cys\(^{241}\), generates higher amounts of the reaction products PDI\(_{\text{ox}}\) and H\(_2\)O\(_2\), as it is active irrespectively of the scarcity of PDI\(_{\text{red}}\) [10].

In cells, oxidase activity of wild-type Ero1\(\alpha\) (Ero1\(\alpha\)-WT) or Ero1\(\alpha\)-AA can be stimulated by the addition of the membrane-permeable reductant dithiothreitol (DTT). This manifests in a more prominent detection of the reaction product H\(_2\)O\(_2\) in the ER of Ero1\(\alpha\)-overexpressing cells [8,10,19,20] and can mechanistically be explained by DTT-stimulated reduction of inhibitory disulfides in Ero1\(\alpha\) [4–6,10,21]. In case of Ero1\(\alpha\)-AA, DTT-mediated activation is connected to the enhanced opening of Cys\(^{208}\)–Cys\(^{241}\) by PDI\(_{\text{red}}\).
Based on the data summarized above, Ero1α-AASS might be hyperactive, because it mimics the DTT-activated state of Ero1α-AA. To test this possibility, we examined to what extent DTT influences Ero1α-AASS-catalyzed production of H2O2. We used a cell line that expresses Ero1α-AASS only in presence of doxycycline and stably harbors the H2O2-sensitive fluorescent protein HyPer in the ER (HyPer ER) [10,22]. HyPerER, the fluorescence excitation spectrum of which does not overlap with the one of doxycycline [12], is prone to oxidation not only by H2O2 but also by PDIox [23,24]. In DTT-treated cells, however, the oxidation of HyPerER can basically be ascribed to H2O2 only, as DTT raises the concentration of Ero1α-derived H2O2 but lowers the levels of PDIox [21] (Fig. 1A, left panel). Using this model system in presence of 0.5 mM DTT, we analyzed the fold-increase in HyPerER oxidation in cells treated with doxycycline as compared to untreated cells. We found that expression of Ero1α-AASS did not lead to equal or stronger production of H2O2 compared to Ero1α-AA. Rather, doxycycline-induced HyPerER oxidation was lowered by ~40% in Ero1α-AASS expressing cells (Fig. 1B, bars 6 and 8). Likewise, expression of Ero1α-C208S/C241S trended to induce less prominent HyPerER oxidation than that of Ero1α-AA (Fig. 1B, bars 4 and 8), whereas mutation of Cys104 and Cys131 increased oxidase activity as expected (Fig. 1B, bars 2 and 6, bars 4 and 8).

These findings were incompatible with the notion that Ero1α-AASS mimics the DTT-activated state of Ero1α-AA. Apparently, the Cys208/Cys241 pair is required for full catalytic activity of Ero1α under reducing conditions. Since mutation of these cysteines precludes the thiol-disulfide exchange reactions that engage PDI to the O2-reducing end of Ero1α, the stimulatory effect of PDIred (Fig. 1A, left panel) drops out in these mutants. Thus, although poorly defined yet, the mechanism of PDIred-mediated stimulation of Ero1α-AA cannot solely rely on the removal of the Cys208–Cys241 disulfide.

We next examined if these new findings could also be recapitulated in a reconstituted system. Previously, we compared the performance of Ero1α-AA and Ero1α-AASS in oxidizing a one-time bolus of their substrate PDIred [10]. In this setting, PDIox was generated faster in the Ero1α-AASS- compared with the Ero1α-AA-catalyzed reaction. Furthermore, purified Ero1α-AASS produced significantly higher levels of H2O2 compared to Ero1α-AA [10]. In order to mimic the situation in the ER of DTT-treated cells where PDIred is constantly being regenerated, we added an excess of GSH to the reaction (Fig. 2A). This setup matches the commonly used practice to assay the oxidase activity of Ero1α in vitro [5,7,15–17,25]. Consistent with the results presented in Fig. 1B, the O2 concentration dropped less rapidly in presence of 0.5 mM DTT, we analyzed the fold-increase in HyPerER oxidation in cells treated with doxycycline as compared to untreated cells. We found that expression of Ero1α-AASS did not lead to equal or stronger production of H2O2 compared to Ero1α-AA. Rather, doxycycline-induced HyPerER oxidation was lowered by ~40% in Ero1α-AASS expressing cells (Fig. 1B, bars 6 and 8). Likewise, expression of Ero1α-C208S/C241S trended to induce less prominent HyPerER oxidation than that of Ero1α-AA (Fig. 1B, bars 4 and 8), whereas mutation of Cys104 and Cys131 increased oxidase activity as expected (Fig. 1B, bars 2 and 6, bars 4 and 8).

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Oxidase activity of Ero1α in the ER of live cells can also be assessed using a DTT washout approach that is coupled to the time-resolved quantification of the cellular GSSG:GS\textsubscript{tot} ratio [8,10,13]. Because Ero1α is activated by DTT (Fig. 1A), the levels of PDI\textsubscript{red} drop and GSSG builds up rapidly in response to DTT removal through the Ero1α–PDI–GSH cascade (Fig. 3A). Furthermore, the peroxidase-catalyzed reduction of H\textsubscript{2}O\textsubscript{2} contributes to the buildup of GSSG [8] (not depicted in the figure). Ectopic over-expression of Ero1α is evidently reflected in the time-course of GSSG:GS\textsubscript{tot} after DTT washout. Thus, in cells that over-express Ero1α-WT, GSSG:GS\textsubscript{tot} transiently rises to about 500% of its steady-state value in the time-scale of seconds [13,26]. This massive accumulation of GSSG is followed by a decline and the restoration of the steady-state GSSG:GS\textsubscript{tot} [8,10,13].

Consistent with the results in Fig. 1B where over-expressed C208S/C241S mutants displayed lower oxidase activity than their unmutated counterparts in presence of DTT, mutation of the Cys\textsuperscript{208}/Cys\textsuperscript{241} pair decreased the Ero1α-dependent accumulation of GSSG upon DTT washout (Fig. 3B, compare the solid and the dashed lines). Unexpectedly, however, over-expression of Ero1α-AA led to a smaller peak in GSSG:GS\textsubscript{tot} upon DTT washout compared to over-expression of Ero1α-WT (Fig. 3B), although Ero1α-AA is catalytically more active than Ero1α-WT under reductively stimulated conditions (Figs. 1B, 2C and 4D). This may suggest that the decline of GSSG:GS\textsubscript{tot} in Ero1α-AA-expressing cells has already proceeded to significant extent before the assay period. It might also be possible that the constitutive absence of the regulatory Cys\textsuperscript{94}–Cys\textsuperscript{131} disulfide in the C104A/C131A mutant affects the Ero1α activity by altering the stability of the Cys\textsuperscript{208}–Cys\textsuperscript{241} disulfide in an allosteric and/or intermolecular manner (see the following paragraphs for more details).

To address the potential crosstalk between the Cys\textsuperscript{208}/Cys\textsuperscript{241} pair and the Cys\textsuperscript{94}–Cys\textsuperscript{131} regulatory disulfide bond, we measured the reduction potential of Cys\textsuperscript{94}–Cys\textsuperscript{131} in Ero1α-WT and Ero1α-C208S/C241S. As shown in Fig. 4A, purified Ero1α-WT was predominantly in an inactive form (OX2), which harbors the Cys\textsuperscript{94}–Cys\textsuperscript{131} disulfide [4] and migrates faster on a non-reducing SDS gel than the active OX1 form comprising the active-site Cys\textsuperscript{94}–Cys\textsuperscript{99} disulfide. Incubation of Ero1α-WT and Ero1α-C208S/C241S in various GSH/GSSG ratios followed by non-reducing SDS-PAGE revealed that indeed, the Cys\textsuperscript{94}–Cys\textsuperscript{241} pair influences the stability of the Cys\textsuperscript{94}–Cys\textsuperscript{131} disulfide. The C208S/C241S mutations slightly but significantly stabilized the Cys\textsuperscript{94}–Cys\textsuperscript{131} disulfide; the redox
equilibrium constant ($K_a$) of Cys$^{94}$–Cys$^{131}$ in Ero1α–WT is 10.7 ± 0.6 mM and that in Ero1α–C208S/C241S is 12.5 ± 0.7 mM (Figs. 4B and C). It is thus conceivable that the cleavage or absence of Cys$^{208}$–Cys$^{241}$ leads to a higher propensity for the inactive OX2 form. In agreement, Ero1α–C208S/C241S was less active in oxidizing PDI than Ero1α–WT (Fig. 4D).

We next analyzed the binding of the GPx8 peroxidase to the Cys$^{208}$–Cys$^{241}$ region in Ero1α–WT (Fig. 5; mean ± SD). a.u. arbitrary unit, **p < 0.01.

Fig. 5. (A) Schematic representation of the interaction of the luminal domain of GPx8 with the loop region between Cys$^{208}$ and Cys$^{241}$ in Ero1α (left panel) and the Bi–molecular fluorescence complementation (BiFC) of two YFP half-sites fused to either Ero1α or GPx8 (right panel). (B) 18 h after transfection with the indicated constructs, HeLa cells were trypanosed and analyzed by flow cytometry for BiFC fluorescence ($n$ ≥ 5; mean ± SD). a.u. arbitrary unit, **p < 0.01.

In a second model, the specific architecture of the covalent Cys$^{208}$–Cys$^{241}$–linked Ero1α–PD fast complex (previously termed Ero1α–PD fast [10]) is important, e.g. for the optimal channeling of O2 to the active site. This possibility would be in agreement with a previous working model of the Ero1α catalytic cycle, which accommodated the remarkably high levels of Ero1α–PD fast in the ER [10]. According to this working model, Cys$^{208}$–Cys$^{241}$ can hardly be found in the ER in the reduced dithiol state that resembles Ero1α–AASS, as Ero1α–PD fast is kinetically stabilized. In the same vein, the fact that Ero1α–AA but not Ero1α–AASS associates with the natural binding partner GPx8 through its Cys$^{208}$–Cys$^{241}$ region (likely in the context of Ero1α–PD fast) underscores the non-native behavior of Ero1α–AASS [10].

It is important to note that these two mechanistic explanations are not exclusive to each other. Thus, a slightly modified, “mixed” model is possible where it is the formation of the Ero1α–PD fast complex that stabilizes the inhibitory disulfide bonds involving Cys$^{104}$ and Cys$^{131}$. Conversely, reduction of Cys$^{94}$–Cys$^{131}$ and Cys$^{99}$–Cys$^{104}$ could promote the reformation of Cys$^{208}$–Cys$^{241}$ from Ero1α–PD fast. It will be interesting to more closely characterize the communication between the opposing Cys$^{94}$–Cys$^{131}$/Cys$^{99}$–Cys$^{104}$ end and the Cys$^{208}$–Cys$^{241}$ end of Ero1α. It could be based on allostery or, alternatively, intermolecular catalysis either directly between two Ero1α molecules or indirectly via the redox state of PDI or GSH.

In this work, we present the new finding that Ero1α–AASS, which likely adopts a conformation equivalent to fully reduced Ero1α (disregarding catalytic and structural disulfides), does not reach the maximal catalytic turnover rate. Rather, the highest intrinsic activity is reached in presence of the Cys$^{208}$–Cys$^{241}$ pair and a reducing agent, which ensures unlimited supply of PDIred. We conclude that Cys$^{208}$–Cys$^{241}$ plays an unexpected stimulatory role during catalysis. Cys$^{208}$–Cys$^{241}$–dependent stimulation could be due to optimized O2 penetration into the Ero1α–PD fast complex and/or the destabilization of Cys$^{94}$–Cys$^{131}$ and Cys$^{99}$–Cys$^{104}$ in response to the oxidation of the Cys$^{208}$–Cys$^{241}$ pair. Future experiments designed to elucidate these mechanistic possibilities will further increase our understanding of regulated disulfide-bond formation in the ER.

Acknowledgment
This work was supported by a PhD fellowship by the Boehringer Ingelheim Fonds (to TR), the Swiss National Science Foundation (Ambizione), the University of Basel, and the Freiwillige Akademische Gesellschaft (all to CAH), CREST, JST (to K. I.), Grant-in-Aids for Scientific Research on Innovative Areas from MEXT (to K. I. and M. O.), Takeda Science Foundation (to K. I.) and Grant-in-Aids for JSPS Fellows (to S. K. and M. O.).

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