Lifetime imaging of a fluorescent protein sensor reveals surprising stability of ER thiol redox

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Interfering with disulfide bond formation impedes protein folding and promotes endoplasmic reticulum (ER) stress. Due to limitations in measurement techniques, the relationships of altered thiol redox and ER stress have been difficult to assess. We report that fluorescent lifetime measurements circumvented the crippling dimness of an ER-tuned fluorescent redox-responsive probe (roGFPiE), faithfully tracking the activity of the major ER-localized protein disulfide isomerase, PDI. In vivo lifetime imaging by time-correlated single-photon counting (TCSPC) recorded subtle changes in ER redox poise induced by exposure of mammalian cells to a reducing environment but revealed an unanticipated stability of redox to fluctuations in unfolded protein load. By contrast, TCSPC of roGFPiE uncovered a hitherto unsuspected reductive shift in the mammalian ER upon loss of luminal calcium, whether induced by pharmacological inhibition of calcium reuptake into the ER or by physiological activation of release channels. These findings recommend fluorescent lifetime imaging as a sensitive method to track ER redox homeostasis in mammalian cells.

Introduction

Whereas the position of disulfides in folded proteins is ruled largely by thermodynamic considerations (Anfinsen, 1973), the rate at which disulfides form is influenced by the concentration and activity of reduced and oxidized protein thiol oxidoreductases in the ER lumen. The mammalian ER contains at least 20 such protein disulfide isomerase (PDI)–like oxidoreductases, which have in common the presence of one or more thioredoxin domains through which they accept and donate disulfides to contingent proteins (Ellgaard and Ruddock, 2005; Hatahet and Ruddock, 2009). Several factors affect the redox poise of this heterogeneous collection of oxidoreductases: Upstream are enzymes that function as ER oxidases; they accept electrons from the reduced oxidoreductases and transfer them to small molecule acceptors. ERO1, conserved from yeast to mammals, oxidizes PDI family members by reducing molecular oxygen to hydrogen peroxide (Tu et al., 2000; Gross et al., 2006) and the ER localized peroxiredoxin 4 (PRDX4) serves a backup, oxidizing PDIIs by reducing hydrogen peroxide to water (Tavender et al., 2008; Zito et al., 2010b). Other enzymatic (Kodali and Thorpe, 2010; Nguyen et al., 2011) and nonenzymatic processes (Ruddock, 2012) may also contribute to disulfide bond formation in the ER lumen. The aforementioned oxidizing features of the ER are opposed by the ingress of reduced cysteine residues in newly synthesized secreted proteins and by the pool of reduced glutathione (Hwang et al., 1992; Cuozzo and Kaiser, 1999; Chakravarthi et al., 2006).

ER oxidation is vigorously defended. The genes encoding ERO1 and several PDIs are under positive transcriptional control by the unfolded protein response (Frand and Kaiser, 1998; Pagani et al., 2000; Travers et al., 2000) and allosteric mechanisms are in place to regulate the enzymes (Sevier et al., 2007; Appenzeller-Herzog et al., 2008; Baker et al., 2008). These regulatory features likely account for the recovery of disulfides within minutes of reversing a reductive pulse (Braakman et al., 1992; Appenzeller-Herzog et al., 2010). Mice tolerate severe loss-of-function mutations to the two genes encoding ERO1

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Abbreviations used in this paper: DTT, dithiothreitol; FLIM, fluorescence lifetime imaging; PDI, protein disulfide isomerase; roGFP, redox-sensitive green fluorescent protein; SERCA, smooth endoplasmic reticulum calcium; TCSPC, time-correlated single-photon counting; UPR, ER unfolded protein response.

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isoforms, Ero1L and Ero1Lb (Zito et al., 2010a; Chin et al., 2011), further attesting to the redundancy of pathways for disulfide bond formation. Against this background of robustness are studies reporting strong deviation of redox poise under conditions of unfolded protein stress in the ER. Pharmacological and genetic manipulations that adversely affect protein folding homeostasis in the yeast ER are reported to promote a more reducing environment (Merkosmer et al., 2008), whereas in mammals, enforced expression of a difficult to fold secreted protein leads to hyperoxidation, which contributes to failure of secretion (Malhotra et al., 2008).

These studies suggest that deviation from healthy thiol redox poise may contribute to impaired protein folding in the overloaded ER, but offer contradictory indications as to the direction of the deviation. Resolving this conundrum requires tools to accurately measure ER redox poise; a nontrivial problem. It is possible to detect changes in ER redox poise by following the distribution of disulfides and dithiols in sentinel ER resident proteins. Bulk biochemical methods to track the redox status of individual ER proteins exist, but these end-point assays are subject to post-lysis experimental artifacts, lack spatial and temporal resolution, and provide limited information on intra-sample variation. These limitations have stimulated the development of redox-sensitive intravital optical probes that function in the ER lumen. A particularly promising approach has been to exploit the optical perturbation introduced by the presence of a disulfide on the surface of the green fluorescent protein β-barrel (Björnberg et al., 2006; Cannon and Remington, 2008; Meyer and Dick, 2010). Introduction of cysteine residues on the surface of such redox-sensitive GFP (roGFP) results in a fluorescent protein whose excitation properties differ between the dithiol and the disulfide state: The ratio of emitted fluorescence when excited at 390 nm versus 475 nm reports on the ratio of disulfide versus dithiol in the population of probe molecules (Hanson et al., 2004).

The first generation of roGFPS, although useful in tracking redox in the cytoplasm and mitochondrial matrix, proved too reducing for the oxidizing environment of the ER. This challenge was ingeniously met by introducing a single amino acid insertion into the roGFP β-barrel adjacent to cysteine 147, in what were named “roGFP insertion X” (roGFPIX, where X is the inserted amino acid). Destabilization of the optically perturbing 147–204 disulfide in roGFPIX resulted in a less reducing redox environment (Merksamer et al., 2008), whereas in mammals, enforced expression of a difficult to fold secreted protein leads to hyperoxidation, which contributes to failure of secretion (Mallota et al., 2008).

Fluorescent spectroscopy of roGFPIX purified from Escherichia coli showed the expected double peaked excitation spectrum with a prominent peak at 475 nm, after reduction with dithiothreitol (DTT), and its attenuation by oxidation with lipoic acid. The emission spectrum was unaltered by the redox state of the protein (Fig. 1 A). Thus, ratiometric measurements of roGFPIX faithfully reported on the redox status of the probe in vitro, as previously described (Lohman and Remington, 2008).

Thiol redox reactions tend to kinetic isolation (Kemp et al., 2008). Therefore, an in vivo thiol redox reporter should be coupled to the prevalent enzymatic system of the organelle. We tested the response of the sensor in the presence of PDI, an abundant chaperone mediating client protein oxidation and reduction in the ER. Ratiometric measurements tracked time-dependent changes in roGFPIX redox status, as trace amounts of the fully reduced protein in vitro were exposed to an excess of a redox buffer of oxidized and reduced glutathione. Alone, roGFPIX equilibrated slowly with the glutathione redox buffer, reflecting kinetic isolation, however PDI markedly accelerated probe oxidation (Fig. 1 B) with half maximal activation attained at ~20 µM PDI (Fig. 1 C).

The tight coupling of roGFPIX to changes in PDI redox status justified its use as an in vivo probe. When targeted to the ER of mammalian cells, ERroGFPIX colocalized with the ER chaperone BiP and the disulfide isomerase PDI (Fig. 1 D). COS7 cells, with superior spatial resolution, were used to assess colocalization, but similar observations were made in HEK 293T cells (see Fig. 3), which were used in subsequent experiments. Non-reducing immunoblot confirmed the presence of a mixed population of reduced and oxidized ERroGFPIX, which were completely reduced in cells exposed to DTT (Fig. 1 E).

Despite these favorable features, in vivo ERroGFPIX proved unreliable as a ratiometric ER redox probe: Whereas the excitation spectrum of Sepharose beads coated with roGFPIX tracked its redox status in vitro (Fig. S1 A), nonspecific autofluorescence competed with the probe’s feeble signal in vivo, severely compromising the detection of changes in ER redox in cells using the same microscope setup (Fig. S1, B–D). This motivated the development of an alternative readout that could be tracked more readily in vivo.

Fluorescence lifetime imaging detects changes in ER protein thiol redox

The lifetime of the compromising autofluorescence (3.0 ns) was found to be significantly longer than that reported for GFP (2.4 ms; Esposito et al., 2005; Wallrabe Periasamy, 2005) and was unresponsive to DTT (Fig. S2 A). Therefore, we hypothesized...
that this parameter may provide a basis for deconvoluting the specific signal.

Proximity of the C147/C204 disulfide to the modified 2,3-didehydrotyrosine residue of roGFPiE (PDB: 3CD1) suggested that it might influence the lifetime of the fluorophore. Indeed, fluorescence lifetimes of roGFPiE varied from a low of 1242 ± 62 picoseconds, for the fully reduced probe, to a high of 2311 ± 56 ps for the fully oxidized form (Fig. 2 A). Fluorescence lifetimes correlated with changes in excitation ratio in a titration with reduced and oxidized lipoic acid (Fig. 2 B).

In its oxidized form, the fluorescence lifetime of roGFPiE approached that of conventional EGFP and refGFP (a thiol-lacking revertant mutant roGFP; Fig. 2 C). Together these observations suggested that the fluorophore was perturbed in the reduced state of roGFPiE. Interestingly, the fluorescence lifetimes of roGFP1, the C147/C204-containing parent of roGFPiE that lacks the E147a insertion, was long in both its reduced and oxidized configuration (Fig. 2 C), indicating that the perturbation promoted by reduction of the disulfide requires the insertion at 147a. The identity of the inserted residue seemed less important, as reduced roGFPiE, roGFPiL, and roGFPiR had similar short fluorescence lifetimes. However, in their oxidized configuration roGFPiR had shorter fluorescence lifetimes than roGFPiE (1807 ± 28 and 2150 ± 46 vs. 2311 ± 56 ps, respectively), endowing the latter with a superior dynamic range (Fig. 2, A and C). Fluorescence lifetime of roGFPiE was unaffected by the presence of iodine (Fig. 2 D), suggesting that the lifetime alterations in roGFPiX variants does not result from exposure of the fluorophore to collisional quenching and predicting a measure of indifference to the ionic environment.

Given that roGFPiE had the greatest dynamic range of fluorescence lifetime in vitro, it was chosen for further in vivo studies. Fluorescence lifetime imaging microscopy (FLIM) of mammalian cells expressing ErroGFPiE, stained with rabbit polyclonal anti-GFP, mouse monoclonal anti-PDI, and chick polyclonal anti-BiP (as ER markers). The nucleus was visualized by Hoechst staining (in blue in the merged right-most panel). (E) Immunoblot of ErroGFPiE expressed in untreated or DTT-treated (5 mM DTT for 10 min) HEK 293T cells, resolved by nonreducing (NR) and reducing (R) SDS-PAGE. Shown are representative experiments reproduced three times (A–D) or twice (E).
genetically deficient in ER disulfide oxidases, which readily defend ER thiol redox at baseline but exhibit a subtle kinetic delay in disulfide formation after recovery from a pulse of DTT (Fig. 4 D; Zito et al., 2012).

Stability of ER protein thiol redox poise in the face of changing levels of unfolded protein stress

Diverse pathophysiological processes perturb protein folding homeostasis in the ER lumen (Walter and Ron, 2011). However, the effect of the resulting stress on ER thiol redox remains poorly characterized. In budding yeast, an ER-localized roGFP2 had been reported to undergo reduction on exposure to tunicamycin, an inhibitor of N-linked glycosylation that causes unfolded protein stress (Merksamer et al., 2008). However, given the reducing midpoint potential of roGFP2, \(-0.272\) V (Hanson et al., 2004), it is suited to detect catastrophic deviations in redox state, but cannot track physiological variations around the organelles’ midpoint lifetime, as expected (Fig. S2 A). The autofluorescent signal of untransfected cells had a patchy distribution and although foci of high intensity autofluorescence were noted, pixels containing such signal could be rejected based on their unusually long lifetime and lack of response to DTT (Fig. S2 A). Together, these observations confirmed that roGFPiE FLIM tracked protein thiol redox in the ER of living cells and could be unmixed from polluting background autofluorescence.

Despite the dimness of the ERroGFPiE signal, fluorescence lifetime imaging had the sensitivity and time resolution needed to detect subtle and rapid changes in ER redox poise in wild-type cells exposed to DTT or to the oxidizing agent 2,2′-dipyridyl disulfide (DPS; Fig. 4, A–C). Unlike the ratiometric method used to detect redox changes in roGFPiL in vivo, which suffered from the confounding effects of a creeping baseline (van Lith et al., 2011), a steady baseline of fluorescence lifetime was observed in cells expressing ERroGFPiE (Fig. 4 C). FLIM of roGFPiE also readily revealed a defect in disulfide formation in cells genetically deficient in ER disulfide oxidases, which readily defend ER thiol redox at baseline but exhibit a subtle kinetic delay in disulfide formation after recovery from a pulse of DTT (Fig. 4 D; Zito et al., 2012).
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ERroGFPiE (Fig. 5 A), despite strong induction of ER stress, reflected in the induction and nuclear localization of the ER unfolded protein responsive marker CHOP (Fig. 5 B). ERroGFPiE remained unresponsive for up to 12 h (Fig. 5 C), a point at which death became noticeable in the stressed cells.

These observations indicate that prolonged unfolded protein stress does not affect the steady-state thiol redox poise of the potential. Therefore, ERroGFPiE-expressing 293T cells were exposed to conditions that promote unfolded protein stress in the ER: azetidine (a proline analogue that perturbs protein structure), thapsigargin (an inhibitor of the smooth endoplasmic reticulum calcium [SERCA] uptake pump that inhibits protein folding by depleting ER calcium stores), DTT, or tunicamycin. 3-h exposure to tunicamycin or azetidine had no measureable effect on ERroGFPiE (Fig. 5 A), despite strong induction of ER stress, reflected in the induction and nuclear localization of the ER unfolded protein responsive marker CHOP (Fig. 5 B). ERroGFPiE remained unresponsive for up to 12 h (Fig. 5 C), a point at which death became noticeable in the stressed cells.

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Figure 3. Redox-dependent changes to fluorescence lifetime of ER-localized roGFPiE in live cells. (A) Fluorescent intensity–based (in grayscale) and lifetime images (color-coded to the scale of the histogram on the right) of HEK 293T cells expressing wild-type ERroGFPiE (C147-E147α-C204) and its ER-localized mutant derivatives, with or without 2 mM DTT (10 min). A histogram of the distribution of lifetimes in the population of cells is provided (right), noting the mean ± SD lifetime. The asterisk in panel 6 points to a region of intense autofluorescence with similar spectral properties as roGFPiE, but recognizable by its relatively long fluorescence lifetime. (B) Nonreducing (NR) and reducing (R) immunoblot of the GFP in the cells shown in A. The dithiol (RED) and disulfide (OX) forms are indicated. (C) Schema showing the mean fluorescence lifetime of the indicated GFP proteins in vivo and in vitro, as measured in the experiments shown in Fig. 2, A and C, panel A here; and in Fig. S2 B. roGFPiE and ERroGFPiE are in bold font. Redox state is indicated by S-S (disulfide) and SH (dithiol). Treatments with DTT, the oxidizing agent 2,2'-dipyridyl disulfide (DPS), or lipoic acid (LA) are noted. A and B are representative experiments reproduced twice.
Figure 4. Faithful tracking of dynamic changes in ER redox by ERroGFPiE. (A) Fluorescent intensity–based (in grayscale) and lifetime images (color-coded to the scale of the histogram on the right) of HEK 293T cells expressing ERroGFPiE exposed sequentially at 10-min intervals to escalating concentrations of the reducing agent DTT (concentration noted in the inset). A histogram of the distribution of lifetimes in the population of cells is provided (right), noting the mean ± SD lifetime. (B) Plot of the relationship between DTT concentration and probe lifetime derived from the experiment described in A. (C) A trace of time-dependent changes in ERroGFPiE fluorescence lifetime (FLT in picoseconds) in live HEK 293T cells, before, during, and after brief exposure to 2 mM DTT or 1 mM of the oxidizing agent DPS. Fluorescent emission intensity (in black and white) and lifetime (color-coded) images of cells before (UNT), during the DTT pulse, and after its washout (WO) are shown in the bottom panel. (D) As in C, comparison of lifetime changes after a reductive pulse of DTT applied to wildtype (WT) and compound Ero1L−, Ero1L−, and Prdx4-deficient mutant (MUT) mouse embryonic fibroblasts. Shown are representative experiments reproduced twice.
ER. To determine if prolonged unfolded protein stress affects the dynamics of disulfide formation, we compared the recovery of oxidized ErGFPIE after a reductive pulse of DTT in untreated and 3-h tunicamycin-pretreated cells. Both populations rapidly recover their steady-state oxidative poise, a process that was mildly accelerated by tunicamycin pretreatment (Fig. 5 D), presumably due to ERO1 induction. Lowering the burden of unfolded proteins by treating cells with protein synthesis inhibitors was likewise with no measurable effect on ER redox. Neither cycloheximide, which arrests protein synthesis during elongation, freezing nascent chains in the translocon, nor puromycin, which releases the nascent chains from the translocon, affected roGFPIE redox (Fig. 5 E).

Next we turned our attention to the role of the unfolded protein response in maintaining the stability of ER redox. ER redox was defended equally well in mutant mouse cells lacking all IRE1 activity and in mutant cells rescued with an IRE1 transgene (Fig. 6 A; Calfon et al., 2002). Unfortunately, spectral overlap with roGFPIE precluded the use of IRE1 inhibitors (which fluoresce strongly in that range) and confined our analysis to the consequences of long-term disruption of the IRE1 pathway. However, as in mammals the IRE1 pathway plays a crucial role in the long-term process of building ER capacity (Cross et al., 2012); this comparison is likely relevant to any role IRE1 might have in maintaining redox equilibrium.

PERK-mediated translational repression regulates the moment-to-moment flux of proteins into the ER and is thus best studied by acute pharmacological inhibition, which precludes long-term adaptation in cultured cells (Harding et al., 2012). Neither PERK inhibition with the selective kinase inhibitor GSK2606414 (Axten et al., 2012; Harding et al., 2012) nor tunicamycin affected ErGFPIE's redox poise over 15 h of...
observed a correlation between depletion of lumenal calcium and a more reducing ER (Fig. 6 B). This observation was consistent with the idea that ER redox is defended across a broad range of levels of unfolded protein load and breaks down only when the defensive mechanism of translational repression is acutely disabled. It was of interest therefore to note that within minutes of exposure to thapsigargin, ERroGFPiE fluorescence lifetime decreased to a new baseline, corresponding to an apparent reductive shift of 24 mV from −0.209 V to −0.233 V (Fig. 5 A). These observations suggested an unanticipated relationship between perturbed intracellular calcium homeostasis and ER protein thiol redox.

Thapsigargin irreversibly inhibits the SERCA pump; in keeping with this, depletion of ER calcium and the shortening of ERroGFPiE fluorescence lifetime persisted upon thapsigargin washout (Fig. S3, A and B). HEK 293T cells exhibit a wide range of sensitivity to the ER calcium store–depleting effects of thapsigargin, necessitating the use of relatively high concentrations of this SERCA inhibitor (Fig. S3, C and D). To determine if the correlation between depletion of lumenal calcium and a more reducing ER extended to reversible, physiological calcium signaling, we exploited the responsiveness of pancreatic acinar AR42j cells to the gut hormone cholecystokinin (Zhao et al., 1990). As expected, cholecystokinin reversibly depleted lumenal calcium stores, as monitored by the fluorescence lifetime of the FRET-based probe D1ER cameleon (Fig. 7, A and B, left; Palmer et al., 2004). The temporal profile of ER calcium store depletion correlated well with the cholecystokinin-mediated shortening of ERroGFPiE fluorescence lifetime and with its

Calcium store depletion promotes a more reducing ER

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However, the feeble fluorescent signal of these roGFPiX probes compromises their utility in vivo, where normal background autofluorescence overlaps their emission spectrum and confounds the intensity-based ratiometric excitation measurement used to estimate the redox state of the probe. Our findings indicate that the insertion of residue 147a, which destabilizes the disulfide and renders the probe less reducing (and thus suited to the ER environment), fortuitously subordinates the fluorescence lifetime of the fluorophore to the redox state of the adjacent cysteines. The dithiol has a considerably shorter fluorescence lifetime than the disulfide, thus the redox state of the probe can be estimated by measuring its fluorescence lifetime in vivo.

Both ratiometrics and fluorescence lifetime imaging provide information that is independent of probe concentration. Lifetime measurements offer advantages in deconvoluting signals emanating from cellular autofluorescence, whose lifetime differs considerably from roGFPiE (as exemplified by Fig. 3 A, panel 6, and Fig. S2 A) and thus provides a measurement gated on the fluorescent properties of the probe. The large number of events acquired by photon counting provides reliable data at weak excitation intensities. This is revealed by the narrowness of the distribution of lifetimes measured in vivo under diverse physiological conditions and favors the detection of subtle differences in the state of the probe, while avoiding photo-toxicity and photo-oxidation.

The binary mode of TCSPC eliminates nonlinear amplification of weak signals, a problem affecting analogue light-detecting

restoration upon washout (Fig. 7, A and B, right). By contrast, ER calcium stores were notably preserved in cells experiencing ER stress and only began to decline when viability was compromised at late time points (Fig. S4), suggesting that stability of ER calcium may contribute to the stability of roGFPiE in stressed cells.

Thapsigargin has reciprocal effects on lumenal ER calcium (decrease) and cytosolic calcium (increase; Brayden et al., 1989). To deconvolute their influence on fluorescence lifetimes of ERroGFPiE, we buffered cytosolic calcium with the cell-permeant calcium chelator BAPTA-AM. In the presence of BAPTA-AM, the ratiometric cytosolic calcium probe, INDO1-AM, reported nearly complete buffering of the thapsigargin-induced transient increase of cytosolic calcium (Fig. 7 C). ER calcium was depleted by thapsigargin, as expected (Fig. 7 D, left) but, BAPTA-AM did not reverse thapsigargin’s effect on the redox state of ERroGFPiE (Fig. 7 D, right), implicating changes in lumenal calcium. Importantly, the calcium-dependent changes in ERroGFPiE fluorescence lifetime observed in vivo were not reproduced by exposing reduced or oxidized roGFPiE to different calcium concentrations in vitro, indicating that at neither oxidation state is roGFPiE’s lifetime directly sensitive to calcium (Fig. 7 E).

Discussion
The development of ratiometric optical redox probes suited to the oxidizing environment of the ER was a major conceptual advance (Lohman and Remington, 2008; van Lith et al., 2011). However, the feeble fluorescent signal of these roGFPiX probes compromises their utility in vivo, where normal background autofluorescence overlaps their emission spectrum and confounds the intensity-based ratiometric excitation measurement used to estimate the redox state of the probe. Our findings indicate that the insertion of residue 147a, which destabilizes the disulfide and renders the probe less reducing (and thus suited to the ER environment), fortuitously subordinates the fluorescence lifetime of the fluorophore to the redox state of the adjacent cysteines. The dithiol has a considerably shorter fluorescence lifetime than the disulfide, thus the redox state of the probe can be estimated by measuring its fluorescence lifetime in vivo.

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The binary mode of TCSPC eliminates nonlinear amplification of weak signals, a problem affecting analogue light-detecting
devices commonly used in microscopy. In addition, immunity to saturation of the counting mode used to generate the intensity image and the single light path used throughout the FLIM image acquisition both contribute to its superior accuracy. The practical advantages of lifetime imaging over ratiometric measurements are nicely demonstrated by the contrast between the stable baseline of the probe obtained during repeated lifetime measurement in vivo (Fig. 4C) and the creeping baseline attributed to photo-oxidation caused by the intense illumination required to generate a signal in intensity-based ratiometric measurements (van Lith et al., 2011).

Ratiometric measurement of fluorescent excitation may be acquired by an assortment of configurations. And although these may produce internally consistent data, they are difficult to standardize. Lifetime measurements, by contrast, yield an absolute value that should be comparable across diverse platforms. Notably, our measurement of EGFP fluorescence lifetime, 2,496 ps, is within 3% of values previously reported (Esposito et al., 2005; Wallrabe and Periasamy, 2005).

The photophysical basis of the dramatic difference in fluorescence lifetime of the disulfide and dithiol forms of roGFPiE remains to be determined. The GFP fluorophore is well protected by the encasing β-barrel (Remington, 2011), and fluorescence of reduced roGFPiE is not quenched by halogen. Thus, it seems unlikely that a structural alteration in the wall of the GFP β-barrel exposes the fluorophore to collisional quenching when the C147/C204 disulfide has been reduced. More likely, the shorter lifetime is intrinsic to the disposition of the fluorophore in relation to other residues of the 147a insertion-containing GFP β-barrel, a perturbation that is reversed by formation of the C147/C204 disulfide.

Autonomy of fluorescence lifetime is an important feature in restricting the responsiveness of roGFPiE to changes in the redox environment. ER redox poise is a convenient heuristic term applied to a nonequilibrium situation; no single sentinel disulfide can serve as a general redox probe for the compartment (Kemp et al., 2008). Nonetheless, the dramatic acceleration by PDI of the equilibration of roGFPiE with a glutathione redox buffer, observed here, suggests that roGFPiE is a good surrogate for clients of PDI. As these include reduced, unfolded, nascent secreted proteins, roGFPiE is likely to report on a physiologically relevant aspect of the redox of PDI substrates.

Applying this refined tool to in vivo measurements in living cells unveiled remarkable stability of the ER redox setpoint in the face of considerable variation in unfolded protein load. Neither tunicamycin nor azetidine, which impede folding and induce high levels of ER stress, affected redox poise. Nor was ER redox measurably affected by shutting off the influx of unfolded proteins into the ER. An extreme imbalance between client protein influx and the capacity of the cellular machinery, imposed by disabling the PERK-mediated translational limb of the unfolded protein response in cells exposed to the glycosylation inhibitor tunicamycin, led to an inappropriately reducing ER. But these extreme circumstances showcase the stability of ER redox under more physiological circumstances.

Our findings seemingly conflict with earlier observations in ER-stressed yeast (Merksamer et al., 2008). These differences may reflect phylogenetic distance, but it may also be worth noting that the yeast study was conducted with roGFP2, which is tuned to the cytosol’s redox potential and would likely miss subtle perturbations like the ones we measured upon calcium release or combined exposure to tunicamycin and PERK kinase inhibitor. Thus, the shift in roGFP2 redox status noted in ER-stressed yeast more likely reflects a different, dramatic process, such as catastrophic permeabilization of the ER or accumulation of cytosolic roGFP2 due a stress-induced translocation defect in the yeast (Rubio et al., 2011).

In mammalian cells the ER unfolded protein response activates a gene expression program that defends against accumulation of reactive oxygen species and whose compromise exposes ER-stressed cells to accumulation of endogenous peroxides (Harding et al., 2003). But the stability of ER thiol redox observed here suggests that the accompanying global increase in reactive species reflects something other than ER hyperoxidation and that the restoration of secretion of difficult-to-fold proteins by electron donors (antioxidants; Malhotra et al., 2008) is unlikely to involve their direct action at the level of the ER. There are other hints that in mammalian cells redox in the ER and cytosol may be quite isolated. Partial loss of ERO1 function, which lowers levels of endogenous peroxides in ER-stressed worms (Marciniak et al., 2004), had no similar protective effect in mouse pancreatic β cells (Zito et al., 2010a). These considerations should prompt a reexamination of the site of action of the pro-oxidant strands of the ER unfolded protein response, mediated by the transcription factor CHOP (McCullough et al., 2001).

This refined method for tracking ER redox revealed that depletion of ER calcium results in a more reduced ERroGFPiE at steady state. This was observed in both thapsigargin-treated cells and in a more physiological setting of regulated ER calcium release, induced by an agonist of the membrane cholecystokinin receptor. Calcium depletion promptly leads to unfolded protein stress in the ER lumen (Prostko et al., 1992), but a role for ER protein misfolding in the shift of roGFPiE to a more reduced poise is rendered unlikely by the observation that tunicamycin and azetidine, which severely perturb ER protein folding homeostasis, have no such effect.

The underlying molecular mechanism linking changes in ER calcium stores to ER redox remain to be worked out. It is nonetheless tempting to speculate on the physiological significance of the shift to a less oxidizing ER after calcium release. Reduction of a luminal disulfide involving cysteines 2496 or 2505 by the ER-localized ERp44 attenuates calcium release by the R1 isoform of the inositol phosphate-3 receptor (Higo et al., 2005), and thiol oxidation has been shown to attenuate the reuptake of calcium into the ER via the SERCA pump (Scherer and Deamer, 1986). Conversely, compromise of ERO1 activity limits activity-dependent calcium release in cardiomyocytes (Chin et al., 2011). This dialectic of thiol redox and calcium flux across the ER membrane is consistent with a rectifying feedback mechanism that opposes the consequences of ER calcium depletion on thiol redox poise and a role for the reducing poise elicited by calcium release as a negative feedback loop to conserve luminal calcium or limit cytosolic calcium signaling.
Materials and methods

Plasmid construction and mutagenesis

The GFP-based probe variants were produced by introducing point mutations into roGFP2 [a gift from S.J. Remington, University of Oregon, Eugene, OR; Lohman and Remington, 2008] in the pGEl vector for bacterial expression variants and in EroGFP2 in PCDNA3.1 His vector (created using roGFP2-pBeG10 as a source) for mammalian ER-targeted variants, directed to the ER by a cleavable signal peptide and Territorial KDEL ER retrieval signal, based on the FLAG-M1 tag in the plasmid pFLAG-CMV1 (Sigma-Aldrich).

Transfections, immuno blotting, and immunofluorescence

HEK 293T, COS7, and AR42J cells were obtained from the American Type Culture Collection (Manassas, VA). Ire1α-deleted embryonic mouse cells (Ire1α) derived from outbred 9.5-d embryos homozygous for a targeted deletion of the exon encoding the transmembrane domain and lacking all Ire1 activity, and their retrovirally rescued counterpart (Ire1α+), were described previously (Calfon et al., 2002), as were the outbred mouse embryonic fibroblasts derived from 13.5-d embryos with combined homozygosity for mutant alleles of PRDX4 (that delete the signal peptide containing exon), as described previously (Lohman and Remington, 2008). Transfected HEK293T cells from confluent 100-mm plates were washed in phosphate-buffered saline (PBS) with 20 mM N-ethylmaleimide and lysed in 0.5% Triton X-100, 150 mM NaCl, 20 mM Hepes, pH 7.4, 10 mM CaCl2, 20 mM N-ethylmaleimide, and protease inhibitors. Proteins were resolved by 12% SDS-PAGE and blotted with an anti-GFP serum raised in rabbit. CHOP immunoreactivity was detected by immunofluorescence of 2% paraformaldehyde-fixed cells using a primary rabbit polyclonal serum to mouse CHOP (Ire1α) as described previously (Ron and Habener, 1992). Goat anti–rabbit DyLight488- and anti–mouse DyLight532-conjugated IgG were used as secondary antibodies diluted 1:1000 (Jackson ImmunoResearch Laboratories, Inc.).

Protein purification and enzymatic assays

For in vitro assays, human PDI (PDA1 18–508; a gift of C. Thorpe, University of Delaware, Newark, DE) and roGFP variants were expressed in the E. coli BL21 (DE3) strain, purified with Ni-NTA affinity chromatography, dia lyzed into the reaction buffer, reduced by incubation with 20 mM of DTT, and then buffer exchanged on a PD-10 gel filtration column (GE Healthcare; Blais et al., 2010). Fluorescence lifetime imaging of purified roGFP variants was conducted on samples of fluorescent protein immobilized on protein A-Sepharose beads with rabbit polyclonal anti-GFP serum. Redox titrations and redox potential calculations were performed by first exposing bacterially expressed roGFP2 to a reducing concentration of DTT (20 mM), removing the DTT by gel filtration, and then equilibrating the reduced protein with different redox buffers containing defined concentrations of oxidized and reduced glutathione, as described previously (Lohman and Remington, 2008).

For kinetic assays, reduced PDI (5–150 µM) was equilibrated in 100 mM Hepes, pH 7.4, and 150 mM NaCl buffer containing an excess of DTT, and then incubated with oxidized glutathione (4 mM; Sigma-Aldrich) for 1 h at room temperature, then added simultaneously to all the samples in the experiment. The ratio of fluorescence emission at 520 nm after sequential excitation at 395 and 470 nm was measured using EnSpire Multimode Plate Readers (PerkinElmer).

FIM and intensity-based confocal microscopy and image analysis

A laser-scanning confocal system (510 Meta; Carl Zeiss) with a Plan-Achromat 63× oil immersion lens (NA 1.4) was used to acquire intensity-based fluorescence microscopy images of cells fixed with 2% paraformaldehyde PBS. Images were analyzed using ImageJ with JACoP and RatioPlus plugins (National Institutes of Health) to assess colocalization and ratiometrics.

FLIM experiments were performed on a modified version of a previously described laser-scanning multiparametric imaging system (Frank et al., 2007), with a 63× oil immersion lens (NA 1.4) coupled to a microscope incubator, maintaining standard tissue culture conditions (Okolab), using a pulsed (sub-10 ps, 40 MHz) supercontinuum (430–2,000 nm) light source (SC 450; Fianium Ltd.). Desired excitation wavelength (typically 470 nm for FLIM of GFP variants and 440 nm for D1ER cameleon FRET) was tuned using an acousto-optic tunable filter (AOTF-CVIS; AA Opto-Electronic). Desired emission was collected using 510/470 and 470–490 bandpass filters for GFP variants and D1ER cameleon accordingly and detected by a fast photomultiplier tube (PMC-100; Becker & Hickl GmbH). Lifetimes for each pixel were recorded using TCSPC circuitry (SPC-830; Becker & Hickl GmbH), keeping count rates below 1% of the laser repetition rate to prevent pulse pile-up. Images were acquired over 20–60 s, with a typical flow rate of 5 × 104 photons sec⁻¹, while detector speed saturation (pile-up effect) is not observed below 106 photons sec⁻¹ in this instrument. The data were processed using SPClassic (Becker & Hickl GmbH) fitting the time-correlated photon count data obtained for each pixel of the image to a monoexponential decay function, yielding a value for lifetime on the picosecond scale.

After filtering out autofluorescence (by excluding pixels with a fluorescence lifetime that was out of range of the roGFP probes, i.e., longer than 2,800 ps), mean fluorescence lifetime of single cells or beads was established. The value obtained represented the redox as sensed by roGFP2 in vivo/in vitro or FRET efficiency of D1ER cameleon (inversely proportional to Ca2+ concentration). Each data point is constituted by the average and SD of measurements from at least 10 cells or beads. Unless indicated otherwise, the images shown are of a representative experiment reproduced at least twice.

Calcium imaging

In vivo assessments of cytosolic Ca2+ concentration were performed on cells loaded with 20 µM Indo 1-AM for 30 min (Sigma-Aldrich), excited at 350 nm and measuring the ratio of emission intensities at 405 vs. 485 nm. The FRET-based, ER-optimized D1ER cameleon (Palmer et al., 2004) was used for ER calcium imaging by FLIM (exploiting fluorescence lifetime shortening effect of FRET on the emission of the cyan fluorescent protein segment of D1ER cameleon). Cytosolic Ca2+ buffering was achieved by exposing cells to the acetoxyethyl ester of 1,2-bis(o-aminophenoxy)ethane-N,N,N',N'-tetracetic acid, BAPTA-AM (50 µM; Sigma-Aldrich).

Online supplemental material

Fig. S1 shows insensitivity of ratiometric fluorescence emission spectroscopy of roGFP2 to changes in ER redox of live cells. Fig. S2 shows that fluorescence lifetime of conventional GFP is not altered by changes to the redox environment. Fig. S3 shows irreversible ER calcium depletion and altered ER redox in livers in mice treated with TGFβ. It is shown that unfolded protein stress does not promote lumenal calcium depletion. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201211155/DC1. Additional data are available in the JCB Data Viewer at http://dx.doi.org/10.1083/jcb.201211155.dv.

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Figure S1. Insensitivity of ratiometric fluorescent emission spectroscopy of roGFPiE to changes in ER redox of live cells. (A) Fluorescent photomicrographs of Sepharose beads coated with roGFPiE (purified from E. coli) and placed in a reducing environment (DTT) or oxidizing environment of lipoic acid (LA), after excitation by laser light at the indicated wavelength. The right-most panels are color-coded images of the ratio of emission at the two excitation wavelengths. (B) ERroGFPiE-expressing HEK 293T cells that were untreated (UNT) or DTT treated for 15 min before fixation and imaging with the setting used in A. (C) Excitation scan (at emission of 520 nm) of untransfected or ERroGFPiE-expressing HEK 293T fixed cells in bulk (ERroGFPiE in cells) and coated Sepharose beads (roGFPiE in vitro), as in A. Shown is a representative experiment reproduced twice. (D) Quantization of the measurements in A and B; shown is the mean ± SD (n = 5). Shown are representative experiments reproduced three times.
Figure S2. **Fluorescent lifetime of conventional GFP is not altered by changes to the redox environment.** (A) Fluorescent intensity-based (in grayscale) and lifetime images (color-coded to the scale of the histogram on the right) of untransfected HEK 293T cells cultured in the absence and presence of 2 mM DTT (10 min). A histogram of the distribution of lifetimes in the population of cells is provided (right), noting the mean ± SD lifetime. (B) Color-coded fluorescence lifetime images of HEK 293T cells expressing cytosolic enhanced GFP (EGFP), or ER-localized fusion of proinsulin or calreticulin with EGFP. Where indicated the cells were exposed to a reducing environment (DTT, 5 mM, 10 min). Shown are representative experiments reproduced twice.
Figure S3. **Irreversible ER calcium depletion and altered ER redox poise in thapsigargin-treated cells.** (A) Time-dependent changes in fluorescence lifetime (FLT, in picoseconds) of the ER calcium reporter D1ER cameleon before, during, and after brief (5 min) exposure of HEK 293T cells to a pulse of 2 µM thapsigargin (TG). Shown is a representative experiment reproduced twice. (B) Time-dependent changes in fluorescence lifetime (FLT, in picoseconds) of the ER redox reporter ERroGFPIE before, during, and after brief exposure of HEK 293T cells to a pulse of thapsigargin as in A. Shown is a representative experiment reproduced twice. (C) Intensity-based (in grayscale, top panels) and fluorescence lifetime (color-coded to the scale shown under the bottom panels) of the ER calcium reporter D1ER cameleon in HEK 293T cells exposed to the indicated concentrations of thapsigargin. (D) Quantization of the data shown in C; shown is the mean ± SD of the D1ER cameleon probe at each concentration of thapsigargin. Shown are representative experiments reproduced twice.

Figure S4. **Unfolded protein stress does not promote lumenal calcium depletion.** (A) Time-dependent changes in fluorescence lifetime (FLT, in picoseconds) of the ER calcium reporter D1ER cameleon after exposure of a cohort of HEK 293T cells to 2 µM thapsigargin (TG), 5 µg/ml tunicamycin, 3 mM azetidine, or 2 mM DTT. Representative color-coded fluorescence lifetime images of the cells are shown below the graph. (B) An extended time course of color-coded fluorescence lifetime images of D1ER cameleon expressing live HEK 293T cells, after exposure to ER stress–inducing agents tunicamycin (5 µg/ml) or azetidine (3 mM). Shown are representative experiments reproduced twice.