ERdj4 Protein Is a Soluble Endoplasmic Reticulum (ER) DnaJ Family Protein That Interacts with ER-associated Degradation Machinery*§

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Background: ERdj4 is an endoplasmic reticulum chaperone co-factor that assists with destruction of misfolded secretory proteins.

Results: ERdj4 is a mobile soluble luminal protein that is recruited to the misfolded secretory protein quality control machinery.

Conclusion: The role of ERdj4 in secretory protein destruction is specified by association with the degradation machinery.

Significance: These studies provide a new insight into how different ERdj family members regulate the fates of their clients.

Protein localization within cells regulates accessibility for interactions with co-factors and substrates. The endoplasmic reticulum (ER) BiP co-factor ERdj4 is up-regulated by ER stress and has been implicated in ER-associated degradation (ERAD) of multiple unfolded secretory proteins. Several other ERdj family members tend to interact selectively with nascent proteins, presumably because those ERdj proteins associate with the Sec61 translocon that facilitates entry of nascent proteins into the ER. How ERdj4 selects and targets terminally misfolded proteins for destruction remains poorly understood. In this study, we determined properties of ERdj4 that might aid in this function. ERdj4 was reported to retain its signal sequence and to be resistant to mild detergent extraction, suggesting that it was an integral membrane protein. However, live cell photobleaching analyses of GFP-tagged ERdj4 revealed that the protein exhibits diffusion coefficients uncommonly high for an ER integral membrane protein and more similar to the mobility of a soluble luminal protein. Biochemical characterization established that the ERdj4 signal sequence is cleaved to yield a soluble protein. Importantly, we found that both endogenous and overexpressed ERdj4 associate with the integral membrane protein, Derlin-1. Our findings now directly link ERdj4 to the ERAD machinery and suggest a model in which ERdj4 could help recruit clients from throughout the ER to ERAD sites.

Within the environment of a cell, the ability of a protein to perform its function depends on the “availability” of the protein. The probability of encountering and interacting with substrates reflects a combination of subcellular localization, abundance, and the mobilities of both the protein and its substrates. For example, a freely and rapidly diffusing protein can readily encounter a substrate, even if the substrate is relatively immobile. In contrast, if both protein and substrate diffuse slowly and are expressed at low levels, then encounters will be rare events. The latter situation could represent an inefficient reaction or a cellular regulatory mechanism. In this study, we describe how ERdj4 availability is regulated.

In the endoplasmic reticulum (ER), nascent proteins fold and assemble with the assistance of an array of chaperones (1, 2). The ER Hsp70 chaperone, BiP, participates in both protein folding and quality control of unfoldable proteins (3, 4). Similar to all of the Hsp70 family members, the ability of BiP to perform both of these roles depends on its interactions with co-factors, including the DnaJ-like proteins, ERdjs, that stimulate the ATPase activity of BiP and stabilize its binding to substrates. In many cases, ERdjs can bind directly to unfolded proteins and hand the proteins off to BiP (5, 6). By stimulating the ATPase activity of BiP, ERdjs transition BiP from a low to high affinity substrate binding form (7). In other cases, DnaJ-like proteins can stimulate the ATPase activity of their Hsp70 partner without interacting directly with substrates (8, 9). The nucleotide states of Hsp70 family members regulate their conformation and substrate affinity. Nucleotide-free and ATP-bound Hsp70 family members have low affinity for substrates, whereas ADP-bound Hsp70s bind substrates with high affinity (10). Similar to all Hsp70s, the intrinsic ATP hydrolysis rate of BiP is inefficient, and ERdjs accelerate the reaction 2–5-fold (5, 11–15). The ER DnaJ-like, termed ERdj, proteins include seven members (ERdj1/Mtj1, ERdj2/Sec63, ERdj3/HEDJ, ERdj4/Mdg1, ERdj5/JPDL, ERdj6/p58IKK, and ERdj7), four (ERdj1, 2, 4, and 7) of which have been reported to be integral membrane proteins.

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5 The abbreviations used are: ER, endoplasmic reticulum; ERAD, endoplasmic reticulum-associated degradation; FLIP, fluorescence loss in photobleaching; FRAP, fluorescence recovery after photobleaching; Prl, bovine prolactin; RFP, monomeric red fluorescent protein; ROI, region of interest; sfGFP, superfolder GFP; SS, signal sequence; VSV G, vesicular stomatitis virus G protein; MDCK, Madin-Darby canine kidney; Tricine, N-[2-hydroxy-1,1-bis(hydroxymethyl)ethyl]glycine.
(3). ERdj1–3 and ERdj6 have been linked to folding of nascent proteins, often at the translocation channel (3, 9, 16). In contrast, ERdj4 and 5 play important roles in the ER-associated degradation (ERAD) of unfolded secretory proteins (17–19). Interestingly, ERdj4 is reported to be membrane-associated (11), whereas ERdj5 is a soluble protein (18).

Subcompartment localization may help define substrate or functional specificity for ERdj proteins, similar to the way that lectin chaperones, membrane-associated calnexin, and soluble calreticulin interact with different sets of substrates (20). ERdj4 levels are exceptionally low in homeostatic cells (15) but rise significantly when the ER unfolded protein response is activated (11). Together, these findings suggest that ERdj4 would normally encounter substrates at relatively slow rates because of both the low mobility caused by membrane association and its low abundance. Unfolded protein response-mediated up-regulation could compensate for the slow membrane diffusion and increase encounters by simply increasing ERdj4 levels. Dong et al. (17) reported that overexpression of ERdj4, indeed, increases turnover of mutants of the surfactant protein SP-C.

In this study, we investigated regulation of ERdj4 availability in live cells through photobleaching analysis. We found that ERdj4 is mobile throughout the ER and, surprisingly, that it is not an integral membrane protein. However, ERdj4 does associate with at least one ER integral membrane protein, the ERAD component Derlin-1. This finding suggests a mechanism for how differences in ERdj substrate specificity are achieved. Just as ERdj1–3 preferably interact with nascent proteins, at least in part because of association with the ER translocon, ERdj4 interactions with clients slated for ERAD could be specified through direct association with ERAD machinery.

**EXPERIMENTAL PROCEDURES**

**Plasmid Construction and Transfection**—To construct ERdj4-GFP and ERdj4sfGFP, mouse ERdj4 cDNA (11) was PCR-amplified with primers: 5′-GATCGTACGGCCACCATGGCTACTCCACAGTCAG (forward) and 5′-GATCACCTACTCCACAGTCAG (reverse) that include a Kozak sequence (bold type) and NheI and AgeI sites (underlined) for ease of subcloning into a monomeric GFP (21) or sfGFP (22, 23) vector based on the Clontech N1-GFP backbone. VSV G-GFP (24) was mutated to increase ER retention without interfering with its low abundance. Unfolded protein response-mediated up-regulation could compensate for the slow membrane diffusion and increase encounters by simply increasing ERdj4 levels. Dong et al. (17) reported that overexpression of ERdj4, indeed, increases turnover of mutants of the surfactant protein SP-C.

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**Immunofluorescence and Imaging of Live and Fixed Cells**—For immunofluorescence, the cells were fixed with freshly diluted 3.7% formaldehyde in PBS for 15 min at room temperature, permeabilized with PBS with 0.1% Triton X-100. Blocking was performed in 10% fetal bovine serum in 1× PBS. Subsequently, the cells were labeled with primary antibodies, followed by Alexa 555-conjugated anti-rabbit or anti-mouse IgG secondary antibodies (Invitrogen). Live cells were imaged in phenol red-free RPMI supplemented with 10 mM Hepes, GlutaMAX, and 10% fetal bovine serum. Live cells were imaged on a 37 °C environmentally controlled chamber of a confocal microscope system (Duoscan; Carl Zeiss MicroImaging, Inc.) with a 63×/1.4 NA oil objective and a 489-nm 100-milliwatt laser. Composite figures were prepared using Photoshop CS4 and CS5 and Illustrator CS4 software (Adobe Systems, San Jose, CA).

**Photobleaching Analysis**—Fluorescence recovery after photobleaching (FRAP) and fluorescence loss in photobleaching (FLIP) were performed by photobleaching a small region of interest (ROI) and monitoring either fluorescence recovery or loss with repeated bleaching in the same ROI over time, as described previously (29, 30). Fluorescence intensity plots and effective diffusion coefficient (D_{eff}) measurements were calculated as described previously (29, 30). To create the fluorescence recovery curves, the fluorescence intensities were transformed into a 0–100% scale and were plotted using Kaleidagraph 3.5 (Synergy Software). The p values were calculated using a Student’s two-tailed t test in Excel (Microsoft) or Prism 5.0c (Graphpad Software Inc., La Jolla, CA).

**ImmunobLOTS and Immunoprecipitations**—Total cell lysates for immunoblotting were prepared by lysing COS-1 or MDCK cells in (1% SDS, 0.1 M Tris, pH 8.0) 6-well plates at 80–90% confluence. The lysates were separated on 12% Tris-Tricine
gels, transferred to nitrocellulose, probed with the indicated antibodies, developed using Pierce enhanced chemiluminescent reagents (Thermo Scientific, Rockford, IL), and exposed to x-ray film. Anti-GFP, horseradish peroxidase-labeled anti-rabbit, and anti-mouse reagents were purchased (Jackson Immunoresearch Laboratories). To isolate protein from cells overexpressing ERdj4, 2 × 10^7 COS-1 cells were transfected with pSGmERdj4 and harvested 24 h later in Nonidet P-40 lysis buffer (50 mM Tris, 150 mM NaCl, 0.5% Nonidet P-40, 0.5% deoxycholate, 1 mM PMSF, and Roche complete protease inhibitor mixture EDTA free) and broken in a Dounce homogenizer, and membranes were pelleted by centrifugation at 500 × g. The isolated membranes, which contained the ER, were lysed in Nonidet P-40 buffer, divided into equal fractions, and immunoprecipitated with the indicated antisera. Rabbit IgG was used as negative control. Immunoprecipitated proteins were separated to SDS-PAGE and immunoblotted with the indicated antibodies.

To isolate endogenous ERdj4 complexes, 1 × 10^8 P3U.1 cells were incubated with the membrane-permeable cross-linking agent dithiobis(succinimidyl propionate) as previously described (5). The cells were incubated in Hepes homogenization buffer (25 mM Hepes, 125 mM KCl, 1 mM PMSF, and Roche complete protease inhibitor mixture EDTA free) and broken in a Dounce homogenizer, and membranes were pelleted by centrifugation at 500 × g. The isolated membranes, which contained the ER, were lysed in Nonidet P-40 buffer, divided into equal aliquots, and immunoprecipitated with the indicated antisera. Rabbit IgG was used as negative control. Immunoprecipitated proteins were separated by SDS-PAGE and immunoblotted.

Membrane Fractionation of Cells Overexpressing Murine ERdj4—Nontransfected P3U.1 cells or COS-1 cells transfected with pSGmERdj4 and harvested at 24 h were fractionated. Membranes fractions were prepared by homogenizing 2.6 × 10^7 cells in 5 ml of Hepes buffer (25 mM Hepes, 125 mM KCl), and a postnuclear supernatant containing ER membranes was obtained by centrifugation at 500 × g for 10 min. The resulting supernatant was divided evenly into six 250-μl aliquots. After higher speed centrifugation (10,000 × g), the pelleted membranes were resuspended in 250 μl of PBS alone or PBS containing the indicated amount of digitonin or 0.1 mM Na2CO3. Samples were rocked at 4 °C for 1 h. In all cases, the samples were then centrifuged at 16,000 × g to repellet the membranes and attached proteins. Cell equivalents of the membrane pellets and supernatants were subjected to electrophoresis and immunoblotting.

Mass Spectrometry and Edman Degradation—Exogenously expressed ERdj4 was immunoprecipitated from COS-1 cells 40 h post-transfection, separated on SDS-polyacrylamide gels and stained with Sypro Ruby. Excised protein was treated with trypsin and subjected to electron spray ionization mass spectrometry (Hartwell Center for Bioinformatics). For N-terminal sequencing of ERdj4, ~6 × 10^7 COS-1 cells transiently transfected with pSGmERdj4 were lysed in Nonidet P-40 lysis buffer and immunoprecipitated with a monoclonal ERdj4 antibody. The immunoprecipitated material was separated by SDS-PAGE, transferred to a PVDF membrane, stained with Commassie Blue, excised, and analyzed by Edman degradation in an ABI 494 protein sequencer (Tufts University Core Facility).

Statistics—To minimize the cell-to-cell variables such as cell cycle stage or contact inhibition, we always selected flat, mononuclear, nonmitotic cells in cultures at between 40 and 70% confluence and expressing low to modest levels of fluorescent proteins for analysis. The data were analyzed using a two-tailed Student’s t test (using Graphpad Prism 5.0c; Graphpad Software La Jolla, CA) to compare the different conditions. The variances of data sets were compared using an F-test (Prism) to establish whether to use equal or nonequal variance t tests. Significance was tested using α = 0.01.

RESULTS

The ERdj4-GFP Fusion Is Functional—To better understand how ERdj4 encounters ERAD substrates and participates in the ERAD process, we created a live cell reporter for ERdj4. To visualize ERdj4 behavior in live cells, we generated a fusion of mouse ERdj4 with superfolder GFP (sfGFP) (Fig. 1A). The sfGFP-tagged ERdj4 had a mobility on SDS gels of ~50 kDa, which is consistent with a 27-kDa increase in size caused by the
addition of GFP (Fig. 1B). The presence of a 5-nm sfGFP did not obviously impair ERdj4 functionality, as determined by comparisons with untagged or HA-tagged variants of ERdj4. The sfGFP fusion associated as well as untagged ERdj4 with its Hsp70 partner, BiP, in co-immunoprecipitation experiments (Fig. 1C). In addition, we observed co-localization of ERdj4-sfGFP with the ER marker (ER-RFP) by live cell imaging and with the endogenous ER resident folding enzyme protein disulfide isomerase by immunofluorescence staining (Fig. 1D). Similar ER localization was observed when ERdj4, fused to a much smaller HA tag, was expressed in MDCK and COS-1 cells (supplemental Fig. S1). In addition, ERdj4sfGFP did not co-localize with the Golgi complex marker p115. These results confirmed that the sfGFP tag does not interrupt known characteristics of ERdj4 and argue that mobility studies performed with this construct are likely to be informative. Notably, we consistently observe lower expression levels of the sfGFP fusion compared with the overexpressed untagged protein. This is possibly due to differences in the vectors used to express these two proteins. The untagged construct vector contains a β-globin intron that significantly stabilizes and thus increases mRNA levels (31), whereas the sfGFP fusion does not.

**ERdj4 Signal Sequence Is Cleaved**

**ERdj4 Is More Mobile Than Predicted in Cells**—Using our new live cell reporter of ERdj4 behavior, we asked how mobile ERdj4 was in cells, which would affect its ability to encounter substrates. ERdj1 and 2 are integral membrane proteins that associate with the essentially immobile Sec61 translocation channel (32–34). In these cases, substrate interactions should be relatively restricted to nascent substrates. As a membrane protein, ERdj4 association with ERAD machinery could similarly restrict the mobility of the co-chaperone. In this case, either substrates would have to be brought to ERdj4, or the association of ERdj4 with ERAD components could be triggered by substrate interaction. To investigate ERdj4 availability, we performed photobleaching experiments on live cells expressing the reporter.

First, we asked whether ERdj4sfGFP is at all mobile. Using a technique termed FLIP, a discrete region of the cell was subjected to high intensity laser light to destroy GFP fluorescence. Repeated photobleaching of the same ROI homogeneously depleted fluorescence from the entire ER, demonstrating that GFP-tagged ERdj4 is mobile throughout the ER lumen. Although informative data on mobility are obtained with FLIP, this assay does not quantitate the degree of protein mobility. To quantitate mobility, we performed FRAP (Fig. 2B) and then determined the effective diffusion coefficient (D_{eff}) of the protein. In MDCK cells, the ERdj4sfGFP D_{eff} value (0.6 ± 0.1 μm²/s) was much lower than a freely diffusing fluorescent protein, ER-RFP, but similar to that of the binding partner of ERdj4, BiP (~0.6 μm²/s) (35), as well as other ER chaperones calreticulin and GRP94 (26, 36) (Fig. 2C). We then blocked the synthesis of nascent proteins in the ER to determine whether ERdj4 mobility was sensitive to decreased availability of substrates. Translational inhibition with pactamycin slightly increased the D_{eff} value for ERdj4, although this was not statistically significant (p = 0.1124) compared with that obtained in untreated cells (Fig. 2C). Similar results were also observed in a second cell type, U2OS (Fig. 2D). Although the mobilities of other chaperones, calreticulin and BiP, have been shown to be dependent on substrate levels (26, 35), ERdj4 mobility appears to be relatively insensitive to translational inhibition. This difference in behavior could be due to association of ERdj4 with the ER membrane or an ER membrane protein complex that is relatively stable. Alternatively, it is conceivable that ERdj4 client levels are inherently low at steady state, and/or client interactions are fleetingy transient.

To investigate these possibilities, we exploited a previously described ERdj4 mutant (H54Q) that inhibits the ability of ERdj4 to stimulate BiP ATPase activity (11) and also appears to decrease release of clients from ERdj4 (17). Note that the mutant did not significantly bind correctly folding proteins (17). In keeping with this, both HA- and GFP-tagged wild-type ERdj4 bound much less efficiently than the corresponding QPD mutants to the client NSF1 LC (Fig. 3, A and B). Taken together with the data from Dong et al. (17), ERdj4 interactions with clients appear to be normally highly transient. We then asked whether there are significant levels of endogenous clients in untransfected cells. FRAP analysis of chaperone mobility can report on changes in global client levels (35). FRAP analysis of the QPD mutant revealed significantly slower fluorescence recovery compared with WT cells (Fig. 3, C and D). The decreased recovery was not due to disruption of the ER, because the inert reporter ER-RFP still readily recovered in co-transfected cells (Fig. 3). Thus, ERdj4 is constantly engaging and releasing clients in cells, but this flux is so rapid that only trapping mutants can reveal the extent of the client burden.

The FRAP results for ERdj4 were surprising, because it was previously reported to be an integral membrane protein (11), and the ER integral membrane proteins that have been characterized by this method have D_{eff} values of between ~0.3 and 0.5 μm²/s (37–40). The higher mobility of ERdj4sfGFP raised an important question: was the mobility of ERdj4sfGFP distinguishable from a typical integral membrane protein in the ER? To answer this, FRAP analysis was performed on an ER-retained mutant VSV G-GFP chimera (25, 39), which contains a single transmembrane domain. We observed a striking difference between the mobilities of ERdj4 in both untreated and translationally inhibited cells compared with VSV G-GFP (p = 0.0243 for untreated cells and p > 0.0001 for pactamycin-treated cells) (Fig. 2C). In both cases, ERdj4 diffused at substantially higher rates (0.6 and 0.8 μm²/s versus 0.3 μm²/s) than VSV G. Similarly, in U2OS cells, the single membrane spanning Lull protein diffuses at the much slower D_{eff} of 0.3 μm²/s (40). Furthermore, the experimental variation in observed D_{eff} values was substantially tighter for VSV G-GFP, consistent with a protein constrained in mobility by a high viscosity environment. Together, these data suggested that the mobility of ERdj4 was either atypical for an integral membrane protein or that ERdj4 might not be an integral membrane protein.

Previous data for endogenous ERdj4 suggested the protein tends to co-fractionate in a detergent extraction assay similar to at least one ER membrane protein, calnexin (11). We therefore revisited this point. ERdj4 has been reported to be a nonsecreted, type II resident ER protein that is anchored in the ER membrane via an uncleaved N-terminal signal sequence (11). However, ERdj4 does not possess an NH₂-domain double argi-
nine (RR) motif (41) that is typically required for retention of Type II integral membrane proteins in the ER. A closer inspection of the signal sequence of ERdj4 revealed that it has a predicted hydrophobic \( \alpha \)-helical transmembrane domain of only 11 amino acids (underlined) (\( \text{NH}_2\)-MATPQSVFVFACILM-\text{TELILAS} \), which is shorter than reported for most resident ER integral membrane proteins, which are typically 14–17 hydrophobic amino acids in length (42, 43). In fact, in a survey of 40% of the human integral membrane proteome, no transmembrane domains shorter than 14 hydrophobic amino acids were reported on known ER membrane proteins (43). Presumably, the minimum transmembrane length is constrained by the thickness of the ER lipid bilayer (44). These features raise the possibility that ERdj4 is not stably integrated into the ER membrane as originally proposed.

We investigated the membrane association and signal sequence (SS) processing of both endogenous and transiently transfected ERdj4 by several biochemical methods. First, the cells were extracted with the detergent digitonin, which selectively binds cholesterol. Because the ER membrane has low concentrations of cholesterol, integral membrane and membrane-associated proteins tend to remain associated with pellet fractions. Our control integral membrane protein, calnexin, fractionated primarily in the pellet fraction (Fig. 4). The soluble proteins ERdj3 and BiP (luminal) and p97 (cytoplasmic) were more readily extracted into the supernatant when 0.1% digitonin was used. In contrast, ERdj4 fractionates primarily in the pellet fraction, as had been previously observed (11). Extraction with a membrane-disrupting detergent, deoxycholate, released all of the proteins into the soluble fraction (Fig. 4). To deter-

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**FIGURE 2. ERdj4sfGFP is a dynamic ER protein.** A, FLIP experiment of an MDCK cell transiently expressing ERdj4sfGFP. The cell was repeatedly photobleached in the ROI (white box), and images are collected between bleaches. If the fluorescent proteins are mobile and can pass through the ROI, all fluorescence will be depleted. B, FRAP of U2OS cell transiently expressing ERdj4sfGFP. The cells were photobleached a single time in the ROI (white box), and recovery of the signal was monitored over the indicated times. Scale bars, 10 \( \mu \text{m} \). C and D, FRAP \( D_{\text{eff}} \) values with ERdj4sfGFP, VSV G-GFP, or ER-RFP transiently transfected in MDCK (left) or U2OS cells (right). Each dot represents the \( D_{\text{eff}} \) value for a single cell and solid lines indicate mean \( D_{\text{eff}} \) values. Translational inhibition was performed with 0.2 \( \mu \text{M} \) pactamycin (Pact) for 1 h to deplete cells of nascent substrates for ERdj4. The mean \( D_{\text{eff}} \) value \( \pm \) S.E. is reported over each column. The dashed line in C was added to emphasize the lack of overlap between the \( D_{\text{eff}} \) of a well established integral membrane protein (VSV G) and the reported transmembrane protein ERdj4 (plus pactamycin (Pact)). The overall higher \( D \) values in U2OS cells likely reflect either decreased ER tortuosity (58) and/or decreased viscosity, because molecular diffusion is dependent on environmental viscosity (59).
mine whether ERdj4 is integrated into the ER membrane, we performed sodium carbonate extraction (45). Sodium carbonate opens microsome membranes into sheets and disrupts many protein-protein interactions, primarily leaving only integral membrane proteins associated in the pellet fraction. After treating microsomes with sodium carbonate, the control membrane protein calnexin was retained in the pellet fraction, and the control soluble proteins, as well as ERdj4, were quantitatively released from the membranes (Fig. 4). Together, these results argue ERdj4 is not an integral membrane protein.

The Signal Sequence of ERdj4 Is Cleaved — Since the original study describing ERdj4 was reported (11), new and improved computational tools have become available for predicting ER targeting SSs and their potential to be cleaved (46). We submitted the mouse and human ERdj4 protein sequences to the SignalP 3.0 server, which predicted that the SS is cleaved with very high probability (0.964 for mouse and 0.953 for human) and that this most likely occurs between amino acids Ala-23 and Ser-24 (Fig. 5A). The small difference in the probability scores is likely due to the fact that the human signal sequence differs from mouse by two conservative amino acids, V17I and E19V, respectively. Although the SignalP 3.0 algorithm is robust, we sought biochemical evidence that the SS is cleaved. We expressed a series of ER-targeted GFP reporters (Fig. 5, B and C) that were predicted to have defined sizes following SS cleavage. Using 12% Tricine-SDS-PAGE, we found that the ERdj4 reporter migrated at a size indistinguishable from the similarly sized Prl SS/H11001 reporter (Fig. 5D). The prolactin SS is an efficiently cleaved SS, and we could demonstrate this with additional control proteins. This confirmed that our SDS-PAGE achieves sufficient resolution to distinguish protein species that differ by 2–13 amino acids (supplemental Fig. S2). Together, these data strongly argue the ERdj4 SS also can be efficiently cleaved.

To test for the cleavage of the predicted SS, we first performed a tryptic digest coupled with mass spectrometry on a transfected ERdj4, because the expression of the endogenous protein is too low to perform this experiment. This analysis
ERdj4 Signal Sequence Is Cleaved

FIGURE 4. ERdj4 is not integrated in the ER membrane. A, fractionation of endogenous ERdj4 from P3U.1 cells. The membranes were prepared from 2.5 × 10⁶ cells and divided evenly into five aliquots. After pelleting by centrifugation, the membranes were resuspended in either PBS or PBS containing 0.1% digitonin, 1% deoxycholate (DOC), or 0.1 M Na₂CO₃. The samples were rocked at 4 °C for 1 h and then centrifuged to pellet the membranes. Pellets (P) and supernatants (S) were subjected to electrophoresis and immunoblotting with the indicated antibodies. B, COS-1 cells were transfected for 24 h with pSGmERdj4. The membranes were processed, separated by gel electrophoresis, and immunoblotted with the indicated antibodies as in A.

detected peptides comprising nearly all of the protein, but importantly, the predicted SS was absent from the sequences identified (Fig. 6A). Although trypsin typically cleaves peptides after lysines or arginines, we note that neither amino acid is present in the predicted SS. Our inability to detect the N-terminal peptide in this analysis is consistent with cleavage of the predicted SS but is inconclusive, because hydrophobic peptides are not always identifiable by this method. To unambiguously determine whether the predicted SS is cleaved, we isolated ERdj4 and subjected it to Edman degradation (Fig. 6B). Consistent with the SignalP 3.0 prediction, the predominant detectable species began at serine 24. There are no other sequences in ERdj4 that resemble a transmembrane domain. Thus, ERdj4, like ERdj3, 5, and 6, is actually a soluble luminal protein.

ERdj4 Associates with ERAD Component—ERdj4 participates in ERAD-L, a form of ERAD in which misfolded luminal domains of soluble and single membrane-spanning proteins are recognized and retrotranslocated by ERAD machinery (47). The machinery includes integral membrane ER proteins Derlin-1 and Hrd1 (48, 49). We hypothesized that these proteins could be potential binding partners for ERdj4. Immunoprecipitation of overexpressed ERdj4 failed to reveal interactions with Hrd1 (not shown). However, significant amounts of Derlin-1 were observed to co-immunoprecipitate with ERdj4 (Fig. 7A). Because the levels of expression in this type of experiment are so much higher than the endogenous ERdj4 levels, which are quite low in unstressed cells (11), we were concerned that the interaction with Derlin-1 could be an overexpression artifact. To test the physiologic relevance of this interaction, we performed co-IPs in nontransfected PU3.1 cells and then immunooblotted against ERAD components. Once again, anti-ERdj4 co-immunoprecipitated Derlin-1, but not p97 or Hrd1 (Fig. 7B). As a positive control, the ERAD-L components p97, Derlin-1, and Hrd1 all co-immunoprecipitated with each other to varying degrees, consistent with previous reports (50).

DISCUSSION

Together, our findings reveal several new insights into the availability of ERdj4 in the ER lumen. First, ERdj4 is now reclassified as a soluble ERdj protein, and therefore it is not necessarily confined to the ER membrane. In practical terms, this means ERdj4 can diffuse to its substrates, and the diffusion is comparable with other chaperones. Given the much lower level of ERdj4 relative to other ER chaperones, the probability of a substrate encountering ERdj4 is likely to be quite low, regardless of the ERdj4 D value. However, during ER stress, when ERdj4 levels increase significantly (11), encounter probabilities will increase and can enhance ERAD of substrates (17). A co-chaperone that can diffuse more rapidly than integral membrane proteins can more efficiently sample the ER and enhance clearance of ERAD substrates.

The demonstration that the ERdj4 SS is cleaved helps resolve the biochemical problem of reconciling the short hydrophobic helix with the previous conclusion that it was integrated in the ER lipid bilayer. However, cleavage of the SS creates a new problem. How are ERdj4 levels maintained in the ER lumen? ERdj4, similar to ERdj3, lacks a known ER retention motif, such as a C-terminal -KDEL sequence, which can serve as a retrieval sequence that returns secretory pathway proteins from the intermediate compartment of the cis-Golgi complex to the ER by retrograde trafficking (51, 52). Retention of ERdj4 in the ER could occur through at least three general types of mechanisms: 1) binding to a protein that provides an ER retrieval motif similar to the way glucosidase IIα provides a -KDEL sequence for its binding partner glucosidase IIα (53), 2) exclusion from ER exit sites, or 3) the presence of a previously uncharacterized ER retention/retrieval motif (54).

Our data and previous reports provide evidence for mechanism 1 but do not exclude the possibilities of 2 and 3. First, ERdj4 reversibly binds BiP, a KDEL-containing chaperone. The transient interactions with BiP could significantly retard secretion from cells. Second, ERdj4 can interact with at least one component of the ERAD machinery, the ER-localized integral membrane protein Derlin-1. In future studies, it will be important to identify any other binding partners of ERdj4 that might participate in ER retention or function. We cannot exclude the possibility of a novel ER retention motif or an exclusionary mechanism to prevent entry into ER exit sites. These details will be an important focus for future studies.

Why ERdj3, ERdj4, and p58IPK lack the -KDEL sequence found on most resident ER luminal chaperones is an intriguing mystery. This is an evolutionarily conserved phenomenon, because even the yeast luminal ERdj protein Scj1p lacks a -KDEL-related sequence. We speculate that the levels of these
co-chaperones, as regulators of BiP activity, may need to be tightly controlled by the cell. ER chaperones typically have long half-lives, and the ER lumen lacks many of the reversible post-translational modifications of the cytoplasm, such as phosphorylation and ubiquitylation. Thus, there may be few options for regulating co-chaperone activities in the ER. One possibility is constitutive secretion of a protein out of the ER, possibly exiting the cell. If more ERdj molecules are required, the synthesis of these proteins can be up-regulated, as happens for all three of these ERdj proteins during the unfolded protein response (3). If less protein is needed, synthesis could decrease, and either regulated secretion or ERAD could deplete soluble ERdj levels over relatively short times. For example, the half-time for secretion of a small inert soluble protein out of the ER is calculated to be ~40 min (55). Our labs are actively investigating the mechanisms for regulating the levels or activity of ERdj proteins.

The most significant finding of our study comes from the co-immunoprecipitation of ERdj4 with Derlin-1. This result suggests a simple mechanism for specifying whether an individual ERdj regulates events during protein synthesis or as part of quality control of unfoldable proteins. ERdj1 and 2 directly associate with the nascent protein translocation channel (33) (the Sec61 translocon). ERdj3 has recently been reported to interact with the translocation channel protein Sec61 (16). Thus, all three of these ERdj proteins are positioned to specifically and immediately engage nascent proteins as they enter the ER. In contrast, ERdj4 association with the ERAD machinery helps explain why it appears to be an ERAD-specific ERdj (17, 56). Given the mobility of ERdj4-sfGFP, we anticipate that interactions with Derlin-1 are likely to be transient, and we hypothesize that ERdj4 can bind clients throughout the ER and potentially release them to BiP at ERAD retrotranslocon sites. Future biochemistry studies should help resolve the dynamics of ERdj4 association with Derlin-1 and the importance of this interaction in recruiting BiP to ERAD retrotranslocon sites. In our tissue culture cells overexpressing HA- or GFP-tagged ERdj4, we observed the protein throughout the ER. In cells expressing the extremely low endogenous levels (15) of this protein, we

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**FIGURE 5. Cleavage of the ERdj4 SS upon translocation of a GFP reporter.** A, SignalP 3.0 analysis of the mouse ERdj4 N terminus predicts the presence of a cleavable signal peptide. The predicted N, H, and C domains are colored green, purple, and aqua, respectively. The bold red line indicates the predicted cleavage site. B, SS constructs (gray letters) plus first 10 amino acids of mature domain (black letters) fused to GFP-KDEL. Italicized block letters indicate multicloning site linker amino acids. Arrows mark confirmed Prl (bovine prolactin) and predicted (ERdj4) cleavage sites. C, constructs were transiently transfected in MDCK cells and imaged by fluorescence microscopy. The core GFP-KDEL construct lacks a SS and displays a cytoplasmic and nuclear distribution, whereas Prl SS +10 and ERdj4 SS +10 both robustly exhibit an ER localization pattern. Scale bar, 10 μm. D, transiently transfected cells were lysed and immunoblotted with anti-GFP. The Prl SS is efficiently cleaved from the GFP-KDEL, leaving only 15 amino acids fused to GFP-KDEL, which is larger than GFP-KDEL alone on a 12% Tricine gel. ERdj4 SS migrates comparably to Prl SS +10 (see supplemental Fig. S2 for additional immunoblot evidence of SS cleavage).

**FIGURE 6. The ERdj4 SS is cleaved in the ER.** A, mass spectrometry analysis of purified transfected murine ERdj4. ERdj4 was immunoprecipitated from COS-1 cells 40 h post-transfection, separated on SDS-polyacrylamide gels, and excised. Protein was treated with trypsin and subjected to electron spray ionization mass spectrometry. The identified peptides are indicated with underlined, bold letters, and spaces are inserted between sequential peptides that were identified. B, N-terminal sequencing of ERdj4. ERdj4 was immunoprecipitated from lysates of COS-1 cells transiently transfected with ERdj4, purified, and analyzed by Edman degradation. The amino acids detected by sequencing are underlined. The sequence underlined with the solid line was the predominant species, whereas that underlined with the dashed line represents a very minor detected species. The downward facing triangle represents the signal sequence cleavage site.
FIGURE 7. ERdj4 interacts with ERAD component Derlin-1. A, COS-1 cells were transfected with pSGmERdj4 and harvested at 24 h. Cell lysates were divided evenly into four aliquots, each of which was immunoprecipitated with the indicated antibodies. IgG was used as negative control. Immunoprecipitated proteins were subjected to electrophoresis and immunoblotting with the indicated antibodies. B, immunoprecipitation from cells expressing endogenous ERdj4. 1 × 10^6 P3U.1 cells were homogenized. The membranes were pelleted by centrifugation and lysed. The lysates were divided into equal fractions and immunoprecipitated with the indicated antibodies. IgG was used as negative control. Immunoprecipitates were subjected to electrophoresis and immunoblotting with the indicated antibodies. IP, immunoprecipitation.

cannot rule out the possibly discrete localization of ERdj proteins to ER functional subdomains (57). Given the proclivities of the different ERdj proteins to bind distinctly rough ER proteins (the translocon) or not, it is tempting to speculate that ERdj functions may be segregated spatially, as well as biochemically.

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REFERENCES

Lai et al. Supplemental Figure 1
Lai et al. Supplemental Figure 2