UBC9-dependent Association between Calnexin and Protein Tyrosine Phosphatase 1B (PTP1B) at the Endoplasmic Reticulum*

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Background: PTP1B is an enzyme localized to the cytoplasmic face of the ER.

Results: Calnexin binds UBC9, is SUMOylated and forms complexes with PTP1B at the ER membrane.

Conclusion: This work reveals a previously unrecognized role for calnexin in the retention of PTP1B at the ER membrane.

Significance: SUMOylation machinery and UBC9 link two ER proteins from divergent pathways.

Calnexin is a type I integral endoplasmic reticulum (ER) membrane protein, molecular chaperone, and a component of the translocon. We discovered a novel interaction between the calnexin cytoplasmic domain and UBC9, a SUMOylation E2 ligase, which modified the calnexin cytoplasmic domain by the addition of SUMO. We demonstrated that calnexin interaction with the SUMOylation machinery modulates an interaction with protein tyrosine phosphatase 1B (PTP1B), an ER-associated protein tyrosine phosphatase involved in the negative regulation of insulin and leptin signaling. We showed that calnexin and PTP1B form UBC9-dependent complexes, revealing a previously unrecognized contribution of calnexin to the retention of PTP1B at the ER membrane. This work shows that the SUMOylation machinery links two ER proteins from divergent pathways to potentially affect cellular protein quality control and energy metabolism.

The endoplasmic reticulum (ER) is an intracellular organelle responsible for many critical cellular functions including the synthesis, folding, post-translational modification, and transport of proteins; the synthesis of lipids and steroids; assembly and trafficking of membranes; Ca²⁺ signaling; regulation of gene expression; energy metabolism; signaling to other cellular organelles; and stress responses (1–3). To support these versatile functions the ER takes advantage of many membrane-associated proteins including molecular chaperones, folding enzymes, oxidoreductases, and various transporters and signaling proteins (1–3). For example, calnexin is an integral ER membrane protein and molecular chaperone involved in the folding and quality control of membrane associated and secreted proteins, including many cell surface receptors regulated by tyrosine phosphorylation (4). Calnexin spans the ER membrane with a large portion of the C-terminal domain (C-tail) residing in the cytoplasm where it may play a role in substrate folding (5, 6). The C-tail undergoes distinct post-translational modifications including phosphorylation and palmitoylation (7–12). The C-tail also affects FGF-dependent STAT3 signaling (13). Palmitoylation of the C-tail of calnexin mediates its association with the ribosome-translocon complex (13) and affects calnexin association with the mitochondria associated region of the ER (10). Together, these post-translational modifications indicate that the C-tail of calnexin may play a functional role in linking ER luminal events with cytoplasmic signaling.

Protein tyrosine phosphatase 1B (PTP1B) is a member of the protein tyrosine phosphatase family of enzymes localized to the cytoplasmic face of the ER (14, 15). PTP1B acts as a critical regulator of the insulin and leptin receptor-dependent pathways (16, 17), plays a role in cancer and modulation of the immune system, and may affect ER stress signaling pathways (17–19). PTP1B is believed to be anchored to the ER membrane via a 35 amino acid C-terminal targeting sequence (ER targeting domain) whereby the protein localizes to the cytoplasmic face of the ER membrane and the nuclear envelope (14). Despite numerous reports on the ER localization of PTP1B, to our knowledge, a contribution of ER proteins to PTP1B association with the ER membrane has received no attention.

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† The abbreviations used are: ER, endoplasmic reticulum; BIFC, bimolecular fluorescence complementation; FLT3, Fms-like tyrosine kinase 3; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; PTP1B, protein tyrosine phosphatase 1; UBC9, ubiquitin-conjugating enzyme 9; CNX, calnexin; SUMO, small ubiquitin-like modifier.
In this study, while testing for a functional role of the calnexin C-tail, we identified several C-tail interacting proteins including UBC9, the SUMOylation E2 ligase, and showed that calnexin can be SUMOylated both in vitro and in vivo. We discovered that calnexin and PTP1B form UBC9-dependent complexes, revealing an unexpected role of calnexin in holding PTP1B to the ER membrane. This work indicated that PTP1B association with the ER membrane is not only restricted to the 35 amino acid C-terminal PTP1B "ER targeting" sequence but also association involves calnexin, an ER molecular chaperone and component of the translocon. This work shows that the SUMOylation machinery and UBC9 link two ER proteins from divergent pathways, calnexin and PTP1B, in a common pathway that may affect cellular protein quality control and energy metabolism.

**EXPERIMENTAL PROCEDURES**

**Plasmids and Mutagenesis**—The GFP-SUMO-1 and human UBC9 expression plasmids were used in this study (20). cDNA encoding canine calnexin C-tail (amino acid residues 486–573) was synthesized by PCR-driven amplification using the following primers: forward primer 5'-CAT GCC ATG GCT GGA AAG AAA CAG TCA AG-3' and reverse primer 5'-GCT CCT GTA GCT TCC TCT CCT TGG CTG GCT TCC-3'. The cDNA was cloned into the pBAD His-tag vector using Ncol and Xbal restriction enzymes to generate pBAD-CNXC-C vector. QuikChange Site-directed Mutagenesis (Stratagene) was used to mutate Lys506 to an Ala (consensus site VKEE) residue (forward primer; 5'-CCT CAG CCA GAT GTG GCG GAG GAG GAA GAA GAA G-3' and reverse primer 5'-CCT TGT TTT CTT CCT CCT CCT CCG GCA CAT CTG GCT GAG G-3'). To create Lys534 (potential consensus site QKSD) or Lys554 (consensus site PKAE) to Ala residue the following DNA primers were used: for L534A forward primer 5'-GGA GGA TG-3 and reverse primer 5'-GGA GGA TG-3 and reverse primer complement; L554A forward primer 5'-GGA CGA TAG GAA ACC TGC GGC AGA AGG A GA GGA GGA TG-3' and reverse primer complement. Recombinant protein was expressed and purified as described for pBAD-CNXC-C. To create pEGFP-CNXC and pEGFP-PTP1B plasmids, cDNA encoding calnexin or PTP1B was cloned from Mouse Brain Quick-Clone cDNA (Clontech) using the following set of primers: calnexin forward primer 5'-TAT ACT CGA GAT GGA AGG GAA GTT GTG AC-3' and reverse primer 5'-TAT AGA ATT CCC CCT CTG CCT GGC TTT CTT GCG-3' and subcloned in frame with GFP into pEGFP-C1 and pEGFP-N1 vectors, respectively.

**Yeast-2-hybrid and Dual Membrane System Yeast-2-hybrid**—The mouse brain Matchmaker cDNA library in pACT2 (Clontech, 638841) was transformed into the yeast strain AH109. The library was screened for interacting proteins with the C-terminal cytoplasmic region (C-tail) (Cys484-Glu571) of mouse calnexin. Comparison of amino acid sequence of mouse and human calnexin C-tail is shown in the Fig. 1F. cDNA encoding the C-tail of calnexin was obtained by PCR-driven amplification of mouse calnexin cDNA using the following primers: forward primer 5'-GGA ATT CTA TAT GTG TCC TGG AAA GAA AC-3' and reverse primer 5'-AAG GTT CTG CAG TCA CTC TCT TCG TGG CT-3'. The PCR product was cloned in frame with the GAL4 DNA-binding domain at Ndel and PstI restriction sites to generate the clone pGBK7-CNXC. Screening of the mouse library was carried out as recommended by the manufacturer (Clontech). Canine C-tail (Cys484-Glu571) was also used as bait to screen a human leukemia (Jurkat T cell) Matchmaker cDNA library (Clontech). Plasmid DNA from positive clones was isolated and used as a template for amplification of the insert in the library vector by the 3' AD (activating domain) sequencing primer 5'-AGA TGG TGC TAC AGT CAC AG-3', and the T7 yeast-2-hybrid sequencing primer, 5'-TAA TAC GAC TCA CTA TAG GG-3'.

The dual membrane system was used to assay yeast two-hybrid interactions (Dualsystems Biotech). The cDNA encoding full-length calnexin and PTP1B were subcloned into pBT3- SUC and pPR3 vector, respectively, and co-transformed into the yeast strain NY521. pOst1 and pPR3 were used as a positive and negative control in the system. Yeast transformation, growth conditions, and assays for β-galactosidase activity were performed according to the manufacturer's instructions (Dualsystems Biotech).

**Bimolecular Fluorescence Complementation (BiFC) Analysis**—YN (amino acid 1–154 of YFP) and YC (amino acid 155–238 of YFP) fragments of YFP were amplified by PCR using the pEYFP-C1 vector as a template. The following primers were used: YN forward primer 5'-TAT AAC CCG TCA TGG TGA GCA AGG GCC AAC GTC TAT ATC ATG GCC-3' and reverse primer 5'-TAT ATG TAC AAG GCC ATG ATA TAG ACG TTG-3'; YC forward primer 5'-TAT AAC CCG TGG ACA AGC AGA ACG GC-3' and reverse primer 5'-TAT ATG TAC AAG CCT TAC ATG ACG TCG TCC ATC-3'. An Age I site at the 5'-end and a Bsr GI site at the 3'-end were introduced with the primers. This allowed exchange of full-length EGFP in pEYFP-PTP1B and pEYFP-CNXC with the YC and YN fragments to generate YC-PTP1B and YN-CNXC plasmids. cDNA encoding the PTP1B deletion mutant missing the last 35 amino acid ER targeting domain (YC-PTP1BΔ35) was synthesized by PCR-driven amplification of YC-PTP1B plasmid DNA using the following primers: forward primer 5'-ATG CGT AC-3' and reverse primer 5'-TAT AGA ATT CCC CCT CTG CCT GGC TTT CTT GCG-3' and subcloned in-frame with GFP into pEGFP-C1 and pEGFP-N1 vectors, respectively.

**Expression and Purification of Recombinant Proteins and in Vitro SUMOylation**—pBAD-CNXC (encoding ER luminal domain of calnexin) and pBAD-CNXC-C (encoding the C-tail of calnexin) vectors were used for expression of calnexin in TOP 10 F' Escherichia coli. Top 10 F' recombinant protein expression was induced with L-arabinose followed by purification of the His-tagged proteins with a Ni2+-nitriloacetic acid-agarose
affinity column, concentrated by a centrifugal filter, and dissolved in a buffer containing 10 mM Tris, pH 7.0 and 1 mM EDTA. For NMR spectroscopy, expression of uniformly $^{15}$N-labeled C-tail of calnexin was performed in minimal medium supplemented with $^{15}$NH$_4$Cl. Recombinant human UBC9 encoded by the pET28b vector was expressed in BL21 E. coli cells and purified as previously described (20). Purified C-tail protein and the C-tail SUMOylation mutant were used with an in vitro SUMOylation kit (Vaxxon) as per the manufacturer’s protocol.

NMR Spectroscopy—All of the NMR experiments were performed on a Varian INOVA 600 MHz NMR spectrometer using Biopack pulse sequences. The NMR samples were prepared in 90% H$_2$O/10% D$_2$O, 100 mM KCl, 10 mM imidazole, 0.25 mM of deuterated 2,2-dimethyl-2-silapentane-5-sulfonic acid (DSS-d$^6$) as chemical shift reference and 0.05% H$_2$O/10% D$_2$O, 100 mM KCl, 10 mM imidazole, 0.25 mM of deuterated 2,2-dimethyl-2-silapentane-5-sulfonic acid (DSS-d$^6$) as chemical shift reference and a pH of 6.8 according to imidazole H2 proton chemical shifts. The spectra were all processed using standard nmrPipe processing (21); prior to the Fourier transform, a sinebell function of 60 to 90° offsets was applied, and then zero filling was used to extend a maximum of twice the number of real points without using any linear prediction. The two-dimensional spectra were visualized and analyzed using NMRViewJ (One Moon Scientific Inc.).

Transfection and siRNA—Mouse fibroblasts were derived from 7-day-old wild-type and calnexin-deficient mice, respectively (22). Cells were immortalized as described previously (23) and maintained in Dulbecco’s Modified Eagle’s Medium supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin at 37 °C with 5% CO$_2$. The Neon Transfection System (Invitrogen) was used for electroporation of cells as per the manufacturer’s protocol with the following settings: wild-type and cnx$^{-/-}$ mouse fibroblasts were electroporated with a pulse voltage of 1500 V, a pulse width of 20 ms, and a pulse number of 1; HeLa cells were electroporated with a pulse voltage of 1005 V, a pulse width of 35 ms, and a pulse number of 2; and NIH 3T3 cells were electroporated with a pulse voltage of 1200 V, a pulse width of 20 ms, and a pulse number of 3.

For knockdown of UBC9 expression, NIH3T3 cells were transfected with siGENOME SMARTpool Mouse UBE2I or a negative scrambled siRNA (Thermo Scientific) using DharmaFECT 1 transfection reagent (Thermo Scientific). Co-transfection of YN-CNX with YC fusion protein (or YC-PTP1B with YN fusion protein) was used as a negative control. EGFP fluorescence and BiFC signals were detected by confocal laser scanning microscopy (Leica TCS SP5).

Isolation of Microsomal ER Membranes—Microsomes were isolated from HeLa cells grown to 80–90% confluency. A protease inhibitor mixture (0.5 mM PMSE, 0.5 mM benzamidine, 0.05 μg/ml aprotonin, 0.025 μg/ml phosphorhodamine, 0.05 μg/ml TLCK, 0.05 μg/ml APMSF, 0.05 μg/ml E-64, 0.025 μg/ml leupeptin, and 0.01 μg/ml pepstatin) (24) and deSUMOylation inhibitor mixture (800 μM N-ethyl maleimide (NEM) and 200 μM iodoacetamide (IAA)) were included in all buffers. Cells were harvested, spun at 1,000 × g for 3 min at 4 °C, resuspended in 20 ml of PBS followed by transfer to LIS Buffer containing 10 mM Tris-HCl, pH 7.5, and 0.5 mM MgCl$_2$ for 10 min. Cells were homogenized with a Dounce homogenizer in a buffer containing 0.5 M sucrose, 10 mM Tris-HCl pH 7.5, 40 μM CaCl$_2$, 300 mM KCl, and 6.3 mM β-mercaptoethanol. The solution was spun at 8,000 × g followed by centrifugation at 120,000 × g for 1 h at 4 °C. The pellet containing the microsomal fraction was washed and resuspended in a buffer containing 5 mM Tris-HCl, pH 7.5, 20 μM CaCl$_2$, and 150 mM KCl. Immunoprecipitation and Western Blot Analysis—Cells at 80% confluency were lysed in a buffer containing 10 mM tris pH 8.0, 150 mM NaCl, 1% Nonidet P-40, 1 mM EDTA pH 8.0, and 10% glycerol, with the protease/deSUMOylation inhibitor mixture or in RIPA buffer containing 50 mM Tris, pH 7.0, 150 mM NaCl, 1 mM EGTA, 1 mM EDTA, 1% Triton X-100, 0.1% SDS, and 0.5% sodium deoxycholate with the protease/deSUMOylation inhibitor mixture. After 30 min of incubation on ice, the lysed cells were spun at 15,000 × g for 15 min at 4 °C. The supernatant was pre-cleared with protein A or a 1:1 protein A/protein G bead slurry for 1 h at 4 °C with rotation after which the primary antibodies were added to the supernatant for overnight incubation at 4 °C with rotation. Protein A or a 1:1 protein A/protein G bead slurry (10%) was added for 4–5 h at 4 °C with rotation. The beads were spun down, washed 5 times with lysis buffer, and resuspended in SDS-PAGE sample buffer containing 10% glycerol, 1% β-mercaptoethanol, 2% SDS, 65 mM Tris, pH 6.8, and 0.01% w/v bromphenol blue for analysis. Proteins were separated on 6% or 10% acrylamide SDS-PAGE gels and transferred onto nitrocellulose for Western blot analysis. Two polyclonal rabbit anti-calnexin antibodies were used: SPA-860 (Enzo Life Sciences) raised against a synthetic peptide corresponding to the C terminus of calnexin (amino acid residues 575–593) and SPA-865 (Enzo Life Sciences) raised against a synthetic peptide near the N terminus. Antibodies were used at a 1:1,000 and 1:500 dilution, respectively. Other antibodies used included anti-UBC9 (Abgent), 1:200 dilution; anti-GFP (Abcam), 1:10,000 dilution; β-tubulin, (Abcam), 1:500 dilution; anti-PTP1B (Abcam), 1:100,000 dilution; and anti-GAPDH (Abcam), 1:1,000 dilution.

Subcellular Fractionation—Fractionation was carried out using the Opti-prep (Sigma) system. Wild-type mouse fibroblasts were washed twice with PBS and resuspended in homogenization buffer (10 mM Hepes-NaOH pH 7.4, 250 mM sucrose, 1 mM EDTA, 1 mM EGTA). Cells were lysed via 25 passes through an 18 micron clearance ball bearing homogenizer (Iso-biotec), then centrifuged at 800 × g for 10 min at 4 °C to remove nuclei and cellular debris. The homogenate supernatant was layered onto an eight-step gradient of 25, 22, 19, 16, 13, 10, 7, and 4% (% iodixanol) dilutions of Opti-prep in homogenization buffer. The gradient was centrifuged in a Beckman Sw50.1 rotor for 6 h at 4 °C at 150,000 × g. Twelve fractions were harvested from the top of the gradient, followed by acetone precipitation of proteins. Protein pellets were washed in 100% ethanol, then resuspended in sample buffer, separated on SDS-PAGE followed by Western blot analysis. Antibodies used included GAPDH (Pierce), 1:5,000; GFP (Abcam), 1:10,000; Lamp2b (Abcam), 1:1,000; mannosidase II (Santa Cruz Biotechnology), 1:1,000; and ribophorin I (Santa Cruz Biotechnology), 1:400.

FLT-3 Phosphorylation Assay—Expression constructs for calnexin and calnexin K506A mutant, UBC9 and pcDNA3.1 empty vector were used for calnexin rescue expression cell assay. Calnexin-deficient cells were plated at 1 × 10$^6$ cells per
6-well plate 20 h prior transfection. Cells were transfected with expression vector (3.75 μg) encoding calnexin or calnexin K506A mutant along or with vector encoding UBC9 or empty vector (3.75 μg) using Lipofectamine 2000 according to the manufacturer’s recommendations. Following transfection, cells were grown in a medium containing 10% FBS. Cells were harvested in RIPA containing complete mixture of protease inhibitors. Polyclonal antibodies against phospho-FLT3 (Tyr591) and FLT3 were purchased from Cell Signaling Technology. PTP1B polyclonal antibodies were purchased from Millipore and actin polyclonal antibodies from Sigma Aldrich. Calnexin antibodies were a kind gift from Dr. John J.M. Bergeron (McGill University).

RESULTS

UBC9 Binds to the Calnexin Cytoplasmic Tail—We employed yeast-2-hybrid techniques to identify proteins that interact with the cytoplasmic tail (C-tail) domain (Fig. 1A) of calnexin (26). Using the 88 amino acid mus musculus calnexin C-tail (Cys484-Glu571) as bait, we screened a mouse brain cDNA library and identified nine positive clones (26), one of which encoded UBC9. A second, independent yeast-2-hybrid screen of a human T cell leukemia (Jurkat) library with canine calnexin C-tail (Cys485-Glu573) as bait identified ten putative clones, and also included UBC9.

To further examine the UBC9/calnexin C-tail interaction, we used immunoprecipitation techniques (Fig. 1B) and yeast 2-hybrid analysis (Fig. 2A). Calnexin could be efficiently immunoprecipitated with anti-calnexin antibodies from wild-type cells but not from calnexin-deficient cells (Fig. 1B). Western blot analysis with antibodies against both calnexin and UBC9 revealed that calnexin co-immunoprecipitated with UBC9 in wild-type fibroblasts (Fig. 1B), indicating that UBC9 was bound to calnexin.

Next, we performed subcellular fractionation using Optiprep gradients to assess the amount of UBC9 that co-distributed with calnexin (Fig. 1C). As expected, UBC9 was found in both the cytoplasmic fractions (using glyceraldehyde 3-phosphate dehydrogenase (GAPDH) as a cytoplasmic marker) but a significant amount of the protein (>20%; Fig. 1D) was associated with ER fractions (Fig. 1C, fractions 7–9), along with calnexin and the ER marker ribophorin I (Fig. 1C), suggesting that UBC9 co-fractionated with ER fractions. We also probed for other organelle markers, including mannosidase II (Golgi) and Lamp2b (lysosomes). Finally, we used confocal microscopy techniques to investigate the co-distribution of UBC9 with calnexin. UBC9 and calnexin immunostaining were found in the nuclear envelope and at the ER (Fig. 1E). Quantitative analysis of confocal images revealed an average threshold Pearson’s correlation coefficient of 0.558 ± 0.060, indicating some but not complete co-localization of UBC9 and calnexin at the ER, consistent with UBC9 not being restricted to the ER membrane but also being localized to the nuclear membrane and throughout the cytoplasm (Fig. 1E).

UBC9-dependent Association of Calnexin and PTP1B at ER

The Calnexin C-tail Lacks Structure and Binds UBC9 in a Ca2+-independent Manner—To date, only the structure of the luminal domain of calnexin has been solved (27) and there is no structural information available about the calnexin C-tail. To understand the nature of the C-tail/UBC9 interaction, we employed NMR spectroscopy to examine the structural and biophysical properties of the C-tail alone and with UBC9, in the presence and absence of Ca2+ (28). The small chemical shift dispersion observed for resonances in both the amide and aliphatic regions of the one-dimensional 1H NMR spectrum, combined with the few number of NOE cross-peaks observable in a two-dimensional 1H-1H NOESY NMR spectrum, indicated that the C-tail was mostly unfolded (data not shown). The two-dimensional 1H-15N HSQC NMR spectrum of [U-15N]C-tail also showed little chemical shift dispersion for the amide NH resonances (Fig. 2B, black), confirming the absence of structure, and showed small chemical shift changes after the addition of excess Ca2+ (Fig. 2B, red), indicating weak Ca2+ binding. These results are in agreement with earlier studies on Ca2+-independent GST fusion calnexin fragments (29).

To further examine the nature of the C-tail/UBC9 interaction, we looked at the two-dimensional 1H-15N HSQC spectra of [15N]C-tail and monitored the effect of adding unlabeled UBC9. In the absence of Ca2+, the HSQC of [U-15N]C-tail (Fig. 2C, black) showed approximately ten to twelve resonances which underwent strong chemical shift broadening or disappearance after the addition of UBC9 (Fig. 2C, red). This suggested that these residues were in conformational exchange, created by the binding of UBC9 to calnexin. Several cross-peaks remained unaffected with the addition of UBC9, which was not surprising considering the size of the C-tail of calnexin. Most of the unperturbed resonances came from the resonances with the highest intensities and smaller chemical shift dispersion, suggesting that the N and C termini of calnexin most likely remained labile in the bound form. Two-dimensional 1H-15N HSQC NMR spectra were acquired after adding 6 mM Ca2+ to investigate the Ca2+ requirement for UBC9 binding. Negligible changes were observed, indicating that Ca2+ saturation was not required for UBC9 interaction with the C-tail.

The C-tail of Calnexin is SUMOylated—UBC9 is a SUMO-conjugating enzyme that plays an important role in the SUMOylation of specific target proteins (30). SUMO conjugation modifies a lysine residue, typically within the core consensus motif ψKxExE where ψ is a bulky hydrophobic residue (31). In addition to the core consensus motif, additional flanking amino acid sequences have been proposed to extend the SUMO consensus motif (32, 33). These include the phosphorylation-dependent SUMOylation motif (PSDM) of ψKxExSP (32), and the negatively charged amino acid-dependent SUMOylation motif (NDSM) (33). A global search for NDSM-containing proteins in the SWISSPROT database identified that the highly acidic calnexin C-tail contains a NDSM (amino acid residues 505–518), VK506EEEEEEEEKDK (Fig. 1A), a motif previously used successfully in determining in vivo SUMO substrates (34). To examine if the C-tail of calnexin can be a substrate for SUMOylation at this site, we used an in vitro SUMOylation system with purified canine calnexin C-tail and a K506A C-tail.
UBC9-dependent Association of Calnexin and PTP1B at ER

FIGURE 1. The C-tail interacts with UBC9 in yeast 2-hybrid system, lacks structure and binds UBC9 in a Ca^{2+}-independent manner and, A, yeast 2-hybrid analysis of interaction between calnexin and UBC9. First row: yeast AH109 strain transformed with calnexin C-tail cloned into pGBK7 vector (pGBK7-CNXC) and blank pGADT7. Calnexin C-tail did not interact under these conditions and consequently yeast cells did not grow on the SC/-Leu/-Trp/-His/-Ade plate and no activity of β-galactosidase was observed. Second row: yeast AH109 strain transformed with pGBK7-CNXC and UBC9 cloned in to pGADT7 vector (pGADT7-UBC9). Yeast cells did grow in the SC/-Leu/-Trp/-His/-Ade media with activated β-galactosidase indicating the two molecules interacted. Third row: yeast AH109 strain transformed only with pGBK7-CNXC did not on SC/-Leu/-Trp plate indicative of no interaction between calnexin C-tail alone under these experimental conditions. B, two-dimensional $^1$H-$^1$N HSQC NMR spectra of the C-tail of calnexin in the apo (black) and Ca$^{2+}$ (red) states. The small chemical shift dispersion indicates that the C-tail lacks nascent structure. Small chemical shift changes were observed for some amide NH cross-peaks (indicated by black arrows) after addition of 6 mM CaCl$_2$ suggesting only a weak Ca$^{2+}$ interaction. C, two-dimensional $^1$H-$^1$N HSQC NMR spectra of the $^{15}$N-labeled C-tail of calnexin before (black) and after addition of unlabeled UBC9 (red). Approximately ten to twelve resonances in calnexin (indicated by black arrows) showed significant chemical shift broadening or disappeared with the addition of UBC9. This suggests that these residues were in conformational exchange, created by the binding of UBC9 to calnexin.

mutant. In the presence of SUMO-1, there was a shift in the C-tail mobility of $\sim 12$ kDa as detected with anti-calnexin antibodies (Fig. 3A), a size shift consistent with the addition of SUMO-1. Mutation of Lys$^{506}$ to an alanine abrogated in vitro SUMOylation of the C-tail (Fig. 3A), indicating that the predicted NDSM motif in the calnexin C-tail was the site of SUMO addition. Mutation of Lys$^{534}$ or Lys$^{554}$, two other potential SUMOylation sites in calnexin C-tail, did not eliminate in vitro SUMOylation of the C-tail (Fig. 3A). We concluded that the calnexin C-tail can be SUMOylated in vitro at Lys$^{506}$ within the predicted SUMOylation NDSM consensus site.

Next we examined if calnexin can be SUMOylated in vivo. The SUMO enigma is a well-documented phenomenon, where only a small portion of the total substrate is detectably SUMOylated at any given time, even when functional evidence indicates complete SUMOylation (35). To increase the probability of detecting SUMOylated calnexin (and the corresponding molecular shift) in vivo, we isolated ER microsomes from cultured cells and tested them for the presence of SUMOylated endogenous calnexin. HeLa cells were transfected with an expression vector encoding GFP-SUMO-1 followed by the isolation of ER enriched microsomes. Anti-GFP antibodies were used to pull down GFP-SUMO-1 followed by Western blot analysis with anti-GFP antibodies (to report the presence of GFP-SUMO-1) and anti-calnexin antibodies (to identify calnexin protein). Immunoprecipitation of solubilized microsomes with goat anti-GFP antibodies and immunoblotting with rabbit anti-GFP antibodies indicated that a GFP-SUMO-1 modified protein at $\sim 100$ kDa was immunoprecipitated with anti-GFP antibodies (Fig. 3B). Western blot analysis of the same
UBC9-dependent Association of Calnexin and PTP1B at ER

A purified recombinant calnexin C-tail (wt-CNXC) or Lys506 to Ala calnexin C-tail mutant (CNXC-K506A) (left panel) or Lys534 to Lys554 to Ala calnexin C-tail mutants (CNXC-K534A and CNXC-K554A, respectively) (right panel) were SUMOylated followed by Western blot analysis with anti-calnexin antibodies. The location of the recombinant calnexin C-tail (CNXC) and SUMO-conjugated calnexin C-tail (SUMO-CNXC) is indicated by the arrows. The location of the recombinant calnexin C-tail (CNXC) and SUMO-conjugated calnexin C-tail (SUMO-CNXC) is indicated by the arrows.

The data represent more than three biological replicates.

PTP1B-1 is the first protein to be identified as an ER-associated SUMO modified protein (36). Since we showed here that calnexin interacted with UBC9 and was SUMOylated, we hypothesized that there may be an association between PTP1B and calnexin at the ER membrane. To test this, we carried out immunoprecipitation experiments. PTP1B was immunoprecipitated from both wild-type and calnexin-deficient fibroblasts (Fig. 4A). Next, we tested for the presence of calnexin in PTP1B immunoprecipitated complexes using anti-calnexin antibodies (Fig. 4B). Western blot analysis of immunoprecipitates from wild-type cells and cnx−/− cells (control) revealed that calnexin co-immunoprecipitated with PTP1B from wild-type cells (Fig. 4B), indicating that the two proteins formed complexes. To examine if calnexin and PTP1B interact in situ, we used two independent methods: the DUAL membrane yeast-2-hybrid method (Fig. 4C) and BiFC (Fig. 4D). For DUAL membrane yeast-2-hybrid, full-length calnexin and PTP1B were constructed into the pBT3-SUC and pPR3 vector, respectively, and co-transformed into the yeast strain NYM51. Co-transformation of calnexin and PTP1B expression vectors demonstrated comparable growth to the CNX-pBT3/Nub-Ostl positive control, as well as positive β-galactosidase activity, indicating PTP1B and calnexin formed complexes under these conditions (Fig. 4C).

Next, we used BiFC to examine the in situ interactions between PTP1B and calnexin in mammalian cells (Fig. 4D). Both GFP-calnexin and GFP-PTP1B fusion proteins localized to an ER-like intracellular network (Fig. 4D). For BiFC, we constructed vectors encoding calnexin as a fusion protein with the N-terminal portion of YFP (YN-CNXC) and PTP1B as a fusion protein with the C-terminal portion of YFP (YC-PTP1B). Co-transfection of the YN-CNXC and YC-PTP1B constructs resulted in a robust BiFC signal (Fig. 4D), indicative of an in situ interaction between calnexin and PTP1B. Therefore, based on immunoprecipitation, yeast-2-hybrid, and BiFC techniques, we concluded that calnexin and PTP1B interacted.

PTP1B Association with Calnexin Is UBC9-dependent—Considering that calnexin is an integral ER membrane protein, while PTP1B is a cytoplasmic protein, we envisaged that the PTP1B-calnexin interaction was most likely to occur on the cytoplasmic face of the ER and thus involve the cytoplasmic C-tail of calnexin. Since calnexin and PTP1B are SUMOylated proteins and the calnexin C-tail interacted with UBC9, we asked if the interaction between calnexin and PTP1B was dependent on UBC9 and the SUMOylation machinery. NIH3T3 cells treated with siRNA for UBC9 were transfected with WN-CNXC and YC-PTP1B expression vectors followed by confocal microscope analysis. siRNA treatment reduced the abundance of UBC9 by over 90% and did not affect abundance of WN-CNXC or YC-PTP1B in NIH3T3 cells (Fig. 5A). Importantly, the BiFC signal, reporting PTP1B binding to calnexin, was significantly reduced in UBC9 knockdown cells when compared with cells treated with control, scrambled siRNA (Fig. 5B). Quantitative analysis of confocal images revealed an average intensity of fluorescence with scrambled siRNA: 9.77 ± 0.3 and with UBC9 siRNA: 4.57 ± 0.18, p < 0.05 (Fig. 5C), indicating a
reduction of the signal in cells with silenced UBC9. Taken together these observations indicate that silencing UBC9 expression suppressed the interaction between calnexin and PTP1B.

PTP1B Association with Calnexin Is Independent of the PTP1B ER Localization Domain—PTP1B consists of multiple functional domains including an N-terminal catalytic domain
followed by a proline-rich domain and an ER localization domain (35 amino acids at the C terminus) (Fig. 6A) that is believed to anchor PTP1B to the cytoplasmic face of the ER (14, 17). To test whether PTP1B association with calnexin involved the 35 amino acid C-terminal ER targeting domain, we created an expression vector encoding full-length or truncated PTP1B missing the last 35 amino acid residues as a fusion protein with a proline-rich domain and an ER localization sequence (14). Next, we silenced UBC9 with specific siRNA. Cells were also transfected with YN-CNX and YC-PTP1B expression vectors for BiFC analysis. There was a strong BiFC signal in cells expressing YN-CNX and YC-PTP1B followed by BiFC analysis of cells expressing YN-CNX and YC-PTP1B fusion proteins with silenced UBC9. sc siRNA, scrambled small interference RNA; siRNA, small interference RNA; The data represent more than three biological replicates.

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**FIGURE 5. Silencing of UBC9 affects interactions between calnexin and PTP1B.** A, NIH3T3 cells were treated with scrambled siRNA (sc siRNA) or UBC9 specific siRNA. Cells were also transfected with YN-CNX and YC-PTP1B expression vectors for BiFC analysis. Western blot analysis was carried out with anti-GFP, anti-UBC9, and anti-β-tubulin antibodies. β-tubulin was used as a loading control. * indicates the location of nonspecific protein bands. The location of the YN-calnexin and YC-PTP1B is indicated by the arrows. The data represent more than three biological replicates. B, BiFC in NIH3T3 cells expressing YN-calnexin and YC-PTP1B fusion proteins with silenced UBC9. sc siRNA, scrambled small interference RNA; siRNA, small interference RNA; The data represent more than three biological replicates. C, quantitative analysis of BiFC (YFP) signals. The data represent more than three biological replicates. *, p < 0.05.

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Finally, we tested if SUMOylation of the calnexin C-tail played a role in the formation of calnexin-PTP1B complexes. The K506A mutant of YN-calnexin, previously shown to be SUMOylation-deficient, was expressed in NIH3T3 cells together with YC-PTP1B followed by BiFC analysis. There was a significant reduction of the BiFC signal between YN-CNX-K506A and YC-PTP1B (BiFC with for YN-CNX: 11.11 ± 65; for YN-CNX-K506A: 8.33 ± 0.24, p < 0.01) (Fig. 6C) indicating that SUMOylated Lys506 in calnexin C-tail was important to support calnexin-PTP1B interactions.

To further investigate the subcellular localization of wild-type PTP1B and PTP1B lacking its ER targeting C terminus (PTP1BΔ35), we transfected wild-type and calnexin-deficient fibroblasts with expression vectors encoding either full-length GFP-tagged PTP1B (PTP1B-GFP) or GFP-tagged truncated PTP1B (GFP-PTP1BΔ35). 48 h post-transfection, we performed subcellular fractionations and Western blot analysis to monitor the localization of PTP1B. As expected, we observed PTP1B-GFP primarily in the same subcellular fractions as ER markers (calnexin, ribophorin I) with a small co-fractionation with a cytoplasmic marker (GAPDH) (Fig. 6D). In comparison, GFP-PTP1BΔ35 mostly co-fractionated with GAPDH, indicative of a primarily cytoplasmic distribution (Fig. 6E). However, some GFP-PTP1BΔ35 signal was observed in the same fractions as ER markers, suggesting that PTP1B lacking its classical ER targeting sequence still has some propensity to localize to the ER membrane (Fig. 6E). Similar distribution of GFP-PTP1B was seen in fractions derived from calnexin-deficient cells (Fig. 6, F and G). In conjunction with our BiFC data, we therefore concluded that the deletion of the 35 C-terminal residues of PTP1B severely abrogated but did not completely delete the localization of PTP1B to the ER membrane. We concluded that PTP1B formed UBC9-dependent complexes with calnexin independently of the presence of a PTP1B C-terminal ER tar-
UBC9-dependent Association of Calnexin and PTP1B at ER

A

Catalytic domain
Proline-rich domain
ER targeting domain

NH₂

276

COO⁻

197 432 aa

PTP1B

PTP1BΔ35

B

YFP

DAPI

Merge

anti-UBC9

anti-GAPDH

BiFC: YN-CNX+YC-PTP1Δ35

20 µm

40 µm

* BiFC: YN-CNX+YC-PTP1Δ35

C

YFP

DAPI

Merge

YN-CNX-K506A + YC-PTP1B

BiFC: YN-CNX-K506A + YC-PTP1B

20 µm

40 µm

BiFC signal intensity (AU)

2

4

6

8

10

YN-CNX-K506A + YC-PTP1B

D

Fraction density

25%

Wild-type + GFP-PTP1B

25-kDa

anti-GFP

70-kDa

anti-ribophorin I

100-kDa

anti-calnexin

25-kDa

anti-GAPDH

Fraction number

1 2 3 4 5 6 7 8 9 10 11 12

E

Fraction density

25%

Wild-type + GFP-PTP1Δ35

25-kDa

anti-GFP

70-kDa

anti-ribophorin I

100-kDa

anti-calnexin

25-kDa

anti-GAPDH

Fraction number

1 2 3 4 5 6 7 8 9 10 11 12

F

Fraction density

25%

cnx⁻/⁻ + GFP-PTP1B

25-kDa

anti-GFP

70-kDa

anti-ribophorin I

25-kDa

anti-GAPDH

Fraction number

1 2 3 4 5 6 7 8 9 10 11 12

G

Fraction density

25%

cnx⁻/⁻ + GFP-PTP1Δ35

25-kDa

anti-GFP

70-kDa

anti-ribophorin I

25-kDa

anti-GAPDH

Fraction number

1 2 3 4 5 6 7 8 9 10 11 12
Calnexin Association with PTP1B Affects Its Activity—After concluding that the association of calnexin and PTP1B was not likely to serve as a mechanism for mediating the association of PTP1B with ER membranes, we next asked whether calnexin may instead alter the activity of PTP1B. We looked at the phosphorylation of FLT3, a receptor tyrosine kinase that has been shown to be dephosphorylated by PTP1B. Additionally, certain mutants of FLT3 have been shown to associate with calnexin, leading us to hypothesize the existence of a complex containing calnexin, FLT3, and PTP1B (37). We transfected calnexin-deficient cells with vectors encoding either wild-type calnexin or calnexin K506A, which cannot be SUMOylated. We found that in cnx<sup>−/−</sup> cells expression of wild-type calnexin noticeably increased the levels of phosphorylated FLT3 (P-FLT3) (Fig. 7, A and B), which suggested that PTP1B activity was reduced in the absence of calnexin. Importantly, expression of calnexin K506A mutant was not effective in a full recovery of P-FLT3 levels as compared with cells expression wild-type calnexin (Fig. 7, A and B). As calnexin K506A cannot be SUMOylated and interacts with PTP1B less efficiently, we concluded that calnexin’s interaction with PTP1B may act to inhibit its phosphatase activity (at least with respect to FLT3). Further research is needed to determine if calnexin acts to inhibit global PTP1B activity or instead to regulate the substrate specificity of PTP1B.

**DISCUSSION**

Calnexin is a type I integral ER membrane molecular chaperone that, in conjunction with the oxidoreductase Erp57, facilitates protein folding (38–40). Calnexin can be partitioned into distinct functional domains: the ER luminal domain, and the transmembrane and cytoplasmic domain. The ER luminal domain of the protein binds monoglucosylated glycans and consequently plays a role in the folding of virtually all glycosylated proteins including many cell surface proteins, such as receptor tyrosine kinases (4). The C-terminal domain of calnexin (C-tail) extends from the transmembrane α-helix into the cytoplasm. This unique arrangement of calnexin domains across the ER membrane (chaperone domain in the lumen of the ER and the C-tail exposed to the cytoplasm) confers the protein the ability to modulate cellular functions on both sides of the ER membrane (Fig. 7C). Indeed, the C-tail of calnexin has been proposed to play an important role in the regulation of protein-protein interactions via post-translational modification: the C-tail has been shown to be phosphorylated and palmitoylated, both of which promote calnexin interaction with the ribosome-translocon complex (6, 9, 11, 12, 42–45).

Phosphorylation of the C-tail may additionally act as a molecular switch regulating calnexin-SERCA2b interactions, thereby coupling Ca<sup>2+</sup> signaling with Ca<sup>2+</sup>-sensitive chaperone functions in the ER (7). Here, we showed a novel post-translational modification of calnexin: namely, its SUMOylation at Lys<sup>506</sup>. We also discovered that calnexin interacted with PTP1B in a UBC9 and SUMOylation-dependent manner linking ER luminal events with cytoplasmic signaling (Fig. 7C). These findings strongly support an important role for the C-tail of calnexin in maintaining ER and cellular functions.

Our discovery that the calnexin C-tail underwent SUMOylation in vitro and in vivo was consistent with previous large-scale mass spectrometry screens of SUMOylated proteins identifying calnexin as a potential SUMOylated substrate (46, 47). While UBC9 activity and SUMOylation have been extensively studied in the nuclear compartment as a constituent of transcriptional control (48, 49), examples of integral membrane proteins modified by SUMO have recently emerged, revealing a wider range of cellular SUMO substrates. SUMOylation of these proteins regulates their biological functions and/or intracellular trafficking (50, 51). With respect to ER transmembrane proteins, in addition to calnexin (this study), Ssc2, an ER membrane protein that regulates phosphatidylinositol synthesis and lipid trafficking (52) and SERCA2a are SUMOylated (53). Here we identified calnexin as a new, ER associated, SUMO substrate. Importantly, SUMOylation of calnexin supports PTP1B binding to calnexin and, consequently, it contributes to the association of the phosphatase with ER membrane. This indicates a new role for the SUMOylation machinery in bringing together two functionally diverse molecules to potentially provide a link between ER chaperone function and energy metabolism.

PTP1B is a prototype non-transmembrane tyrosine phosphatase associated with the cytosolic face of the ER membrane (17). Over 20 years ago, Fragion et al. identified a 35 amino acid C-terminal hydrophobic “ER targeting” sequence responsible for its association with the ER membrane (14). It has been widely accepted that the protein partitions into the outer leaflet of the ER phospholipid bilayer by direct insertion of the ER targeting C-terminal domain (14). Interestingly, protein-protein interactions have not been shown to play a role in PTP1B association with the ER membrane (14). With respect to the localization of PTP1B, our results were largely in agreement with the current consensus: namely, that PTP1B was associated with the cytoplasmic face of the ER membrane. Further, we confirmed that PTP1B lacking its C-terminal 35 residues had a severely reduced ability to associate with the ER membrane. However, our subcellular fractionation and BiFC experiments

**FIGURE 6. Interaction between calnexin and PTP1B is independent of the PTP1B 35 amino acid ER-targeting sequence.** A, schematic representation of PTP1B and truncated PTP1B (PTP1BΔ35). Mouse PTP1B (NP_035331.3) consists of 432 amino acids and is composed of a catalytic phosphatase domain (yellow), a proline-rich domain (green), and a 35 amino acid C-terminal ER targeting domain (red). B, BiFC analysis was carried out in NIH3T3 cells expressing an YC-calnexin fusion protein (YN-CNX) and a PTP1B deletion mutant missing its 35 amino acid residue ER targeting domain (YC-PTP1BΔ35). DAPI was used as a nuclear stain. Cells were treated with scrambled siRNA (sc siRNA) or with UBC9 specific siRNA. The graph shows a quantitative analysis of the BiFC signal. The data represent more than three biological replicates. *p < 0.05, C, BIFC analysis in NIH3T3 cells expressing YC-calnexin fusion protein (YN-CNX) or YN-calnexin K506A mutant (YN-CNX-K506A) and YC-PTP1B. DAPI was used as a nuclear stain. The graph shows a quantitative analysis of the BiFC signal. The data represent more than three biological replicates. *p < 0.01, D-G, Opti-prep gradient fractionation of wild-type (D, E) or cnx<sup>−/−</sup> (F, G) mouse fibroblast expressing GFP-PTP1B (D, F) or GFP-PTP1BΔ35 (E, G). Lower density fractions (1–4) correspond to cytoplasmic components, while higher density fractions (5–12) correspond to heavier cellular compartments, including the ER. Western blot analysis was carried out with anti-GFP (to visualize the location of GFP-PTP1B or GFP-PTP1BΔ35 protein), anti-calnexin, anti-GAPDH, anti-ribophorin I antibodies.
FIGURE 7. FLT3 phosphorylation in calnexin rescue cell lines and a proposed model for PTP1B association with ER membrane. A, FLT3 phosphorylation in calnexin rescue cell line. cnx−/− cells were transfected with CNX or calnexin K506A mutant (CNX-K506A) expression vectors. Total lysates from non-transfected and transfected cells were analyzed by immunoblotting for phosphorylated FLT3 (P-FLT3) and membrane were then stripped and re-probed for total FLT3. High and low exposure of blots probed with anti-P-FLT3 antibodies are shown. B, quantitative analysis of the Western blots shown in A. A ration between P-FLT3 to non-phosphorylated FLT3 is presented. A.U., arbitrary units. C, PTP1B can associate with ER membranes via its C-terminal 35 amino acid ER targeting domain (red), and by binding to the calnexin C-tail. The interaction between PTP1B and the calnexin C-tail is independent of the PTP1B ER targeting domain but it is sensitive to the presence of Lys506, an amino acid residue in the calnexin C-tail that can be SUMOylated. 3D structures of the calnexin ER luminal domain, PTP1B and SUMO protein are based on pdb1jhn/pdb, PTN1_Human/pdb, and 1A5R, respectively.
both indicated that even without this ER targeting sequence, a small proportion of PTP1B was still able to localize to the ER membrane. Despite forming complexes with calnexin at the ER membrane, we saw no gross differences in the subcellular distribution of PTP1B in the absence of calnexin. Our data therefore did not support a model where the binding of PTP1B to calnexin serves to anchor or recruit PTP1B to the ER membrane. Instead, we favor a model where the interaction between PTP1B and calnexin affects the pool of targets upon which PTP1B may act; calnexin-bound PTP1B preferentially targets a certain subset of proteins, while free PTP1B may have a bias toward a different subset of targets. Indeed, we found that calnexin inhibits the dephosphorylation of the receptor tyrosine kinase FLT3. This suggested that the interaction of calnexin and PTP1B inhibited its phosphatase activity, whether specific to FLT3 or as a more general mechanism to lower of PTP1B activity.

To date, PTP1B has been shown to modify 30 known substrates and has several protein binding partners (54), and undergoes several post-translational modifications including phosphorylation, oxidation, proteolytic cleavage, and SUMOylation (17). There are multiple functions ascribed to PTP1B including its role in insulin and leptin signaling (55, 56) as a part of energy metabolism. Localization of PTP1B at the ER, and its binding to calnexin, might be important for the regulation of insulin receptors during their biosynthesis (57). Calnexin is the obligate molecular chaperone for the insulin receptor, and is involved in the folding efficacy and homodimerization of the receptor (58, 59). The association of PTP1B with calnexin could provide measures during biosynthesis of the insulin receptor to prevent autoactivation. Many PTP1B substrates are membrane-associated molecules that are translated on the rough ER and are also folding substrates for calnexin (4, 54). Furthermore, PTP1B’s interaction with calnexin may recruit PTP1B to the ribosome-translocon complex, ensuring proximal access to specific phosphatase targets and preventing autoactivation and aberrant cellular signaling events. PTP1B targets and dephosphorylates receptor tyrosine kinases after endocytosis, as they traffic in close proximity to the ER (15, 41, 60). Recently, we showed that the calnexin C-tail binds Src homology 3-domain growth factor receptor-bound 2-like (Endophilin) interacting protein 1 (SGIP1) and influences clathrin-dependent endocytosis due to modulation of the internalization of the receptor-ligand complexes (26). It is plausible that calnexin’s interaction with PTP1B supports a similar function in clathrin-dependent endocytosis, as most of the receptor tyrosine kinases activated at the cell surface maintain an interaction with PTP1B after internalization (15).

In summary, this work demonstrated that the SUMOylation machinery and UBC9 link two ER proteins from divergent pathways, calnexin and PTP1B, in a common pathway that may affect cellular quality control and energy metabolism. SUMOylation is a novel modification of the unique calnexin C-tail that has important implications for regulation of a ubiquitous ER molecular chaperone. In addition, this study demonstrated an interaction between a major metabolic enzyme and a classical ER protein quality control chaperone. As our study linked PTP1B to calnexin as a major quality control chaperone of the secretory pathway, it is possible that the mediation of the interaction between PTP1B and calnexin by the SUMOylation machinery serves as an additional cellular measure of quality control.

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