Investigation of the Endoplasmic Reticulum Localization of UDP-Glucuronosyltransferase 2B7 with Systematic Deletion Mutants

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ABSTRACT

UDP-Glucuronosyltransferase (UGT) plays an important role in the metabolism of endogenous and exogenous compounds. UGT is a type I membrane protein, and has a dilysine motif (KXXX/KXKXX) in its C-terminal cytoplasmic domain. Although a dilysine motif is defined as an endoplasmic reticulum (ER) retrieval signal, it remains a matter of debate whether this motif functions in the ER localization of UGT. To address this issue, we generated systematic deletion mutants of UGT2B7, a major drug-metabolizing enzyme catalyzing glucuronidation, and compared their subcellular localizations with that of an ER marker protein calnexin (CNX), using subcellular fractionation and immunofluorescent microscopy. We found that although the dilysine motif functioned as the ER retention signal in a chimera that replaced the cytoplasmic domain of CD4 with that of UGT2B7, UGT2B7 truncated mutants lacking this motif extensively colocalized with CNX, indicating dilysine motif–independent ER retention of UGT2B7. Moreover, deletion of the C-terminal transmembrane and cytoplasmic domains did not affect ER localization of UGT2B7, suggesting that the signal necessary for ER retention of UGT2B7 is present in its luminal domain. Serial deletions of the luminal domain, however, did not affect the ER retention of the mutants. Further, a cytoplasmic and transmembrane domain–deleted mutant of UGT2B7 was localized to the ER without being secreted. These results suggest that UGT2B7 could localize to the ER without any retention signal, and lead to the conclusion that the static localization of UGT results from lack of a signal for export from the ER.

Introduction

In eukaryotic cells, the secretory pathway is a conserved biosynthetic transport route for secreted proteins and proteins localized in each organelle constituting the pathway. All of the proteins transported in this pathway are synthesized in the endoplasmic reticulum (ER) and delivered to their destinations by vesicular transport (Gomez-Navarro and Miller, 2016). In general, the following two strategies are believed to be necessary for ER proteins to reside in the ER: 1) retention (elimination from the vesicles involved in anterograde transport); and 2) retrieval (capture and return of the escaped ER residents by coat protein I complex (COPI) vesicles (Teasdale and Jackson, 1996; Barlowe and Helenius, 2016). The mechanism of retention remains controversial, but the second mechanism is more established and is known as motif-based retrieval to the ER. The carboxyl-terminal tetrapeptides KDEL and HDEL are necessary for ER-soluble proteins to be retrieved by COPI in mammals and yeast, respectively (Munro and Pelham, 1987; Pelham, 1988). The C-terminal dilysine motif (DM) (KKXX/KXXXX) is also defined as a retrieval signal for type I membrane residents in the ER (Nilsson et al., 1989; Jackson et al., 1993; Gaynor et al., 1994).

UDP-Glucuronosyltransferase (UGT) is one of the major drug-metabolizing enzymes catalyzing glucuronidation, and 22 isoforms have been identified so far. Among them, 19 isoforms have been reported to be involved in glucuronidation in humans (Rowland et al., 2013). Most research on UGT is...
focused on its enzymatic activity, because the enzyme is involved in the metabolism of the second largest number of drugs, following cytochrome P450 (Evans and Relling, 1999). However, UGT is also a model for studies on ER localization. UGT is a typical type I membrane protein, with most of the protein, including the substrate-binding site, on the luminal side of the ER membrane. Only a single transmembrane (TM) helix exists in its C-terminal region, followed by about 20 residues facing the cytosol (cytosolic tail, CT). UGT has a DM at its C-terminus (Fig. 1) (Iyanagi et al., 1986; Harding et al., 1987; Shepherd et al., 1989; Radominska-Pandya et al., 1999), and its functions were well examined. The DM of UGT was transplanted onto the end of the CT of plasma membrane proteins (e.g., CD4, CD8, and ErbB2), which altered their localization from the plasma membrane to the ER (Jackson et al., 1990; Kinosaki et al., 1993). These studies were enough to believe that the C-terminal motif was a major driving force for UGT localization in the ER. However, there is also evidence that UGT is able to localize in the ER in a DM-independent manner. Deletion mutants of rat UGT2B1, human UGT1A6, and other isoforms lacking the CT and TM region are still retained in the ER (Jackson et al., 1993; Meech et al., 1996; Ouzine et al., 1999). In our recent study, we also obtained similar results with UGT2B7. UGT2B7 is an important human isoform catalyzing the metabolism of a huge number of clinical drugs (e.g., morphine and zidovudine) (Rowland et al., 2013). In addition, UGT2B7 is also involved in the glucuronidation of endogenous compounds such as fatty acids, and steroids (Bowlagaha et al., 2007; Bock, 2012). Further, there are reports that UGT can localize to other organelles beside the ER (e.g., mitochondria, Golgi apparatus, and plasma membrane) (Chowdhury et al., 1985; Radominska-Pandya et al., 2005; Ménard et al., 2013; Ziegler et al., 2015). A growing number of studies demonstrates that the ER can make contact with other organelles, exchange membrane composition, and function as a central hub for organelle interaction (Schrader et al., 2015; Shim, 2017). Thus, elucidating the mechanisms by which UGT localizes to the ER is expected to lead to new insights into drug metabolism.

To address this issue, in the present study, we used two separate procedures (cellular fractionation and immunocytochemistry) to ascertain the role of the DM in UGT2B7 localization. We then generated systematic deletion mutants lacking not only the C-terminal TM and/or cytoplasmic domains, but also one or several sections of the ER luminal domain of UGT2B7 to look for another ER retention sequence.

Materials and Methods

Materials. Synthetic oligonucleotides were purchased from Fasmac (Kanagawa, Japan). Restriction enzymes and other DNA-modifying enzymes were from Takara Bio (Shiga, Japan). Glass products for chemistry were from Matsunami Glass (Osaka, Kanagawa, Japan). Restriction enzymes and other DNA-modifying enzymes were from Corning Gentest (Woburn, MA). All other reagents were of the highest quality commercially available.

Subcloning of UGT2B7 and Its Mutants into a Baculoviral Vector. Recombinant baculoviruses coding UGT2B7 wild-type (WT), ΔCT (∆511-529), ΔTM (∆493-529), and human calnexin (CNX) were prepared using Bac-to-Bac Baculovirus Expression System (Life Technologies, Carlsbad, CA) as described previously (Miyauchi et al., 2015). ΔCT (∆525-529) was generated by polymerase chain reaction (PCR) using pFastBac1-WT as a template, KOPI-Plus-Neo DNA polymerase (Toyobo Life Science, Osaka, Japan), and the following a pair of primers: sense, 5'-GGGGTACCCTTATTGTGTTTCTAGAGCGTAATCTGGAACATCGTATGGGTAATGCTAGAAAAGCAAAGAAGGGAAAAATGAT; anti-sense, 5'-GGTACCTTTTTGGCTTTTCTAGC-98°C, 2 minutes; cycling step (×40 rounds): 98°C, 10 seconds; 68°C, 1 minute; hold, 4°C. The PCR product was purified with FastGene Gel/PCR Extraction Kit (NIPPON Genetics, Tokyo, Japan) and subcloned into pFastBac1 vector using KpnI sites. Hemagglutinin (HA) epitope tag (HA-tag) was introduced into the C-terminus of WT to yield WT-HA by QuickChange site-directed mutagenesis. Primers were designed by Agilent QuickChange Primer Design Program (https://www.genomics.agilent.com/primerDesignProgram.jsp), and non-tagged pFastBac1 construct was used as a template. Sequences of the primers were as follows: sense, 5'-GGGGTACCCTTATTGTGTTTCTAGAGCGTAATCTGGAACATCGTATGGGTAATGCTAGAAAAGCAAAGAAGGGAAAAATGAT; anti-sense, 5'-GGTACCTTTTTGGCTTTTCTAGC-3' (underlining indicates coding HA-tag). Thermal cycle parameters were “a two-step cycle” as follows: initial denaturation, 94°C, 2 minutes; cycling step (×20 rounds): 98°C, 10 seconds; 68°C, 4 minutes; hold, 4°C. Template vectors were digested by DpnI treatment, and the reaction mixtures were used in transformation with Competent Quick DH10a (Toyobo, Life Science Department, Osaka, Japan). Further deletion mutants with HA-tag (Δ1-Δ525) were also prepared by site-directed mutagenesis using the pFastBac-WT-HA construct as a template. Primers are listed in Supplemental Table 1. Mutagenesis was carried out under the two-step cycling mentioned above. The nucleotide sequences of the constructs were confirmed by an Applied Biosystems 3130xl Genetic Analyzer (Thermo Fisher Scientific, Waltham, MA), using a BigDye Terminator version 3.1 Cycle Sequencing Kit (Life Technologies). Recombinant pFastBac1 were transfected into competent Escherichia coli DH10Bac strain (Life Technologies). After blue/white selection, a positive single clone was picked up and cultured to obtain recombinant bacmid, a part of baculoviral DNA, according to the user manual.

Culture of Sf9 Cells and Preparation of Microsomes. Sf9 insect cells were cultured under the conditions in our previous study with slight modifications (Ishii et al., 2014). In short, the cells were grown in a 500 ml plastic Erlenmeyer flask (Corning, Lowell, MA) containing SF-9001 medium (Life Technologies) supplemented with 5% FBS and 10 μg/ml gentamycin. To obtain recombinant baculovirus,
Sf9 cells (2 × 10⁶ cell) were seeded in a 175 cm² t-flask (Thermo Fisher Scientific) and then transfected with recombinant bacmids with CellfectinII reagent (Life Technologies) diluted with SF-900II medium in the absence of FBS or antibiotics. After 5 hours of incubation, the medium was replaced with fresh medium, and the cells were cultured for 7 days. The cells were pelleted by low-speed centrifugation, and the supernatant was collected as primary (P1) virus. The pelleted cells were resuspended in 10 ml of PBS, and an aliquot of the suspension was mixed with the same volume of 2 × SDS-PAGE sampling buffer [125 mM Tris-HCl (pH 6.8); 4% SDS; 20% glycerol; 0.006% bromophenol blue; 10% 2-mercaptoethanol] to prepare whole-cell lysate for confirmation of target protein expression. Titer of the recombinant baculovirus was determined using a BacPAK qPCR Titration Kit (Clontech, Mountain View, CA). Baculoviral titer was amplified by several rounds of transfection until a titer of over 1.0 × 10⁹ plaque-forming units/ml was obtained. For the expression of recombinant enzymes, Sf9 cells (2 × 10⁶ cells/ml, 200 ml) were infected with recombinant baculovirus, and cultured for 48 hours. Microsomes were prepared from the protocols according to previously (Ishii et al., 2014).

**Construction of Recombinant Mammalian Expression Vectors.**

cDNA of CD4 was purchased from Ori-Gene Technologies (Rockville, MD) and amplified by PCR with a pair of primers: sense, 5'-CCGCTCGAGAATATGGACGCGGGGATCTCCTTTTAGG-3' (sense primer A; underlining indicates XhoI site); anti-sense, 5'-CCGGGTACCTCAAAATGGCTCATGTCATGTC-3' (underlining indicates KpnI site). The PCR product was digested with XhoI and KpnI and subcloned into the pcDNA3.1/hygro(+) vector using XhoI-KpnI sites. The HA-tag was also added to the C-terminal end of DM, ACT, and ΔTM by site-directed mutagenesis using pFastBac constructs as templates. Primers used in the mutagenesis are listed in Supplemental Table 2, and amplification was carried out in a two-step cycle, as mentioned above. After HA-tag addition, cDNAs of ΔDM-HA, ΔCT-HA, and ΔTM-HA were subcloned into pcDNA3.1/hygro(–) as was done with WT-HA. In addition, mammalian constructs coding UGT2B7 Δ1-Δ11 with HA-tag were generated by two-step cycle mutagenesis. Primers and templates used for preparing Δ1-Δ5 and Δ6-Δ11 are listed in Supplemental Tables 3 and 4, respectively. After DpnI treatment, reaction mixtures were used for the transformation of E. coli. Nucleotide sequences were checked according to the protocol described above.

**Culture of COS-1 Cells and Transfection of Mammalian Expression Vectors.**

COS-1 cells were grown in Dulbecco's modified Eagle medium containing high glucose (Wako Pure Chemical, Osaka, Japan) and 10% FBS. For immunoblotting, cells were seeded in 35-mm dishes the day before transfection, and pcDNA3.1 vectors (2 µg) were transfected with FuGENE6 (Promega, Madison, WI) according to the manufacturer protocol. Twenty-four hours after transfection, the cells were lysed in lysis buffer, 20 mM Tris-HCl (pH 7.4) containing 150 mM NaCl, 1% Triton X-100, 10% glycerol, and protease inhibitor cocktail (Nacalai Tesque, Kyoto, Japan); and sonicated to prepare whole-cell lysates. For immunofluorescence staining, cells were plated onto 13-mm coverslips the day before transfection. The amount of transected pcDNA3.1 was 1 µg with FuGENE6 as a transfection reagent, and immunofluorescence staining was performed 24 hours after transfection.

**Immunoblotting.** Proteins separated by SDS-PAGE were electroblotted onto a polyvinylidene difluoride membrane (Merek Millipore, Burlington, MA). UGT2B7 was detected either by goat anti-mouse polyclonal UGT antibody (Mackenzie et al., 1984) or rabbit anti–HA-tag antibody (Sigma-Aldrich, St. Louis, MO). Rabbit anti-CNX (GeneTex, Irvine, CA) and mouse anti-β-actin (Sigma-Aldrich) were purchased from the sources indicated. The primary antibodies were diluted 2000-fold when used. Immunchemical detection was conducted either with horseradish peroxidase (HRP)–conjugated secondary antibodies, HRP-rabbit anti-goat IgG (MP Biomedicals, Santa Ana, CA) or HRP-donkey anti-rabbit IgG (GE Healthcare, Little Chalfont, UK). The secondary antibodies were diluted 10,000-fold in use. EzWestLumi plus (ATTO, Tokyo, Japan) was used as a substrate of HRP, and the signals were visualized with a ChemiDoc MP System (Bio-Rad, Hercules, CA) and quantified with ImageJ software.

**Immunofluorescence Microscopy.** Immunofluorescence staining was performed according to the protocol described previously with slight modification (Hirata and Tanaka, 2009; Fujimoto et al., 2015). Briefly, COS-1 cells were immobilized on coverslips with 4% paraformaldehyde in PBS (pH 7.4) for 30 minutes at room temperature, and subsequently quenched with 50 mM NH₄Cl in PBS for 15 minutes. Permeabilization and blocking was simultaneously conducted by treating the samples with PBS containing 0.05% saponin and 1% bovine serum albumin (BSA; fraction V) for 30 minutes. The cells were incubated with the following primary antibodies for 1 hour at room temperature: rabbit anti-CD4 antibody (H-370; Santa Cruz Biotechnology, Dallas, TX); rabbit anti–HA-tag antibody (Sigma-Aldrich); rabbit anti-UGT2B7 antibody (ProteinTech, Rosemont, IL); and mouse anti-CNX antibody (BD Biosciences, San Jose, CA). The cells were diluted 300-fold with PBS containing 0.05% saponin and 1% BSA. After a PBS wash (5 minutes, ×3), the cells were incubated for 30 minutes at room temperature with Alexa Fluor 488- and Cy3-labeled secondary antibodies, together with 4′,6-diamidino-2-phenylindole (DAPI) as a nuclear counterstain. The secondary antibody was diluted 300-fold with PBS containing 0.05% saponin and 1% BSA, and the concentration of DAPI was 1 µg/ml in use. After a PBS wash (5 minutes, ×3), coverslips were mounted in MOWIOL 4-88 (Merck Millipore) onto glass slides, and the samples were analyzed by the LSM 700 Laser-Scanning Confocal Microscope (Carl Zeiss, Oberkochen, Germany) equipped with four lasers (405, 455, 555, and 639 nm).
The objective lens used in the analysis was Plan-Apochromat 63×/1.4 oil differential interference contrast. To obtain multiple stained images, laser and filter were optimized automatically for each fluorescence label. Image files were created from raw data with ZEISS Efficient Navigation (ZEN) software, and photographic images were processed using Photoshop (Adobe Systems, San Jose, CA). Colocalization of UGT2B7 and CNX were quantified with ZEN software.

Detection of Aggregated Forms of UGT2B7-HA and Its Mutants by Triton X-100 Treatment. Triton X-100 soluble and insoluble fractions were prepared as described previously with slight modifications (Hirota and Tanaka, 2009). In brief, COS-1 cells transiently expressing UGT2B7-HA and its mutants were harvested in cold sucrose solution (0.25 M sucrose, 1 mM EDTA, 10 mM Hepes, pH 7.4), homogenized by 25 passages through a 23-gauge needle connected to a 1-ml syringe, and subsequently centrifuged at 1000 rpm for 10 minutes. The resulting postnuclear supernatants (homogenate) were centrifuged at 105,000 g for 1 hour, and fractionated into membrane (pellet) and cytosol (supernatant). Membrane fractions were solubilized in 1% Triton X-100 in PBS, and divided into soluble and insoluble fractions by centrifugation (105,000g, 4°C, 1 hour). Insoluble fractions were further solubilized in radioimmunoprecipitation assay buffer, as follows: 50 mM Tris-HCl (pH 7.5) containing 150 mM NaCl, 1 mM EDTA, 1% NP40, 0.1% SDS, and 0.05% sodium deoxycholate. Volumes of each fraction were kept equal, and then homogenate (15 μg protein) and equivalent volumes of the later fractions were analyzed by immunoblotting.

Treatments with Proteasome and Lysosome Inhibitors. UGT2B7-HA, ΔCT-HA, and ΔTM-HA were transiently expressed in COS-1 cells. Twelve hours after transfection, the cells were treated with 1 μM proteasome inhibitor (MG132) and lysosome inhibitor cocktail, 10 μg/ml E-64d, 20 μg/ml leupeptin, and 10 μg/ml pepstatin A for 12 hours. Whole-cell lysates were prepared and analyzed by immunoblotting.

Secretion of the Cytoplasmic and Transmembrane Domain-Deleted Mutants of UGT2B7 and CD4. COS-1 cells were transfected with each construct and cultured for 12 hours in 1 ml of FBS-free Dulbecco’s modified Eagle medium. Cells were harvested in cold sucrose solution, and whole lysates were prepared as described above. Medium was also collected and 400 μl of medium was mixed with 100 μl of 50% trichloroacetic acid, and incubated on ice for 10 minutes. After centrifugation, precipitants were washed with ice-cold acetone twice and analyzed by immunoblotting.

Other Methods. Protein concentrations of baculosomes were determined by the method of Lowry et al. (1951), and those of COS-1 cell lysates were by Protein Assay CBB Solution (Nacalai Tesque) with BSA as a standard.

Results

To assess the role of the C-terminal region of UGT2B7, including the DM, in its ER localization, we designed a series of deletion mutants lacking the ΔDM, ΔCT, and ΔTM, respectively, as illustrated in Fig. 2A. Each deletion mutant, as well as UGT2B7 WT, were expressed in Sf9 insect cells, and their expression was analyzed by immunoblotting with goat anti-mouse low pi form UGT antibody (Mackenzie et al., 1984). Using this antibody, we observed a single band in cell lysates expressing each UGT2B7 construct, which had the predicted molecular weight based on the size of the deletion (Fig. 2B). To determine subcellular localization of the UGT2B7 mutants, we fractionated the homogenates prepared from insect cells simultaneously infected with recombinant baculovirus coding human CNX as an ER marker. We obtained three fractions,

![Fig. 2](attachment:image.png)

**Fig. 2.** Subcellular localization of Sf9 cells infected with CT- and/or TM domain deletion mutants of UGT2B7. (A) Schematic sequences of UGT2B7 deletion mutants. The numbers represent the residue position of WT UGT2B7 counted from the N-terminus. SP, signal peptide. (B) Immunoblotting to confirm the expression of WT UGT2B7 and its mutants in insect cells. Sf9 cells, transfected with recombinant bacmid to obtain P1 viruses, were lysed and analyzed. After the collection of P1 viruses, the pelleted cells were resuspended in PBS, and an aliquot of the suspension was mixed with 2× SDS-PAGE sample buffer to prepare whole-cell lysates. HLM and Mock represent HLM (5 μg protein) and baculosomes prepared from Sf9 cells infected with control viruses (10 μg), respectively. Anti-mouse UGT antibody was used as a primary antibody (Mackenzie et al., 1984). (C) Fractionations of Sf9 cells to determine localization of the UGT2B7 mutants. UGT2B7 WT/mutants were simultaneously overexpressed with CNX in Sf9 cells. Sf9 homogenates were fractionated by sequential centrifugation. The fractions were analyzed by immunoblotting to determine the UGT2B7 WT/mutant subcellular localization. Protein amounts of loaded fractions are indicated below: homogenate (Homo), 10 μg; nuclei & mitochondria (Mt), 2.5 μg; microsomes (MS), 5 μg; cytosol (Cyt), 7 μg. Anti-mouse UGT antibody and anti-CNX antibody were used as primary antibodies. Each band was quantified with ImageJ software, and was shown as the percentage of the total.
Fig. 3. Localization of the CT and/or TM domain–deleted mutants of UGT2B7 and the CD4-UGT chimeric protein in COS-1 cells. COS-1 cells transfected with UGT2B7 WT or the mutants were fixed and stained 24 hours after transfection. (A) Localization of UGT2B7 and UGT2B7-HA (UGT2B7 with a C-terminal HA-tag) were compared with anti-UGT2B7 antibody (green). Cells were also stained red with an antibody to endogenous CNX, a marker protein of the ER. In the merged panels, nuclei stained by DAPI are shown in cyan. (B) Localization of the C-terminal–deleted mutants of UGT2B7. The sequences of the mutants are shown in Fig. 2A. An HA-tag was conjugated to their C-terminus for immunochemical detection. Cells were stained with anti-HA antibody (green) and anti-CNX (red). (C) Colocalization of UGT2B7 WT/mutants and CNX were quantified, and the mean ± S.D. values are shown. (D) Schematic sequences of CD4 and its chimeric proteins whose cytoplasmic domains were replaced with that of UGT2B7. The numbers represent the residue position of WT CD4 counted from the N-terminus. The DM of UGT2B7 is highlighted in the C-terminus of each chimera. SP, signal peptide. (E) Localization of CD4 chimeric proteins. CD4-UGT represents a CD4 chimera whose cytoplasmic domain was replaced with that of UGT2B7. Cells were stained with anti-CD4 antibody, which recognizes the extracellular domain of CD4 (green) and anti-CNX (red). In merged panels, the nuclei are shown in cyan. Scale bar, 10 µm.
nuclei and mitochondria, microsomes, and cytosol, for each sample. All of the deletion mutants were exclusively collected in the microsomal fraction, similar to CNX (Fig. 2C), in agreement with our last report (Miyauchi et al., 2015). To further investigate their subcellular localization, we conducted immunofluorescence staining to analyze colocalization of UGT2B7 and endogenous CNX in COS-1 cells. Overexpressed WT UGT2B7 showed a reticular staining pattern and colocalized with CNX (Fig. 2C). These results support the results of the fractionation approach (Fig. 2C). A chimeric protein that replaced the cytoplasmic domain of CD4 with that of UGT2B7 (CD4-UGT) also colocalized with CNX. However, it was redistributed to the cell surface by the substitution of two lysines with alanines within the cytoplasmic domain (CD4-UGT-2KA) (Fig. 3, D and E), suggesting that the DM acts as an ER retention signal in this chimera. Intriguingly, in contrast to the case of UGT2B7 (Fig. 3A), fusion of the HA-tag to the C-terminus of CD4-UGT (CD4-UGT-HA) altered the ER distribution of CD4-UGT to the cell surface (Fig. 3, D and E), consistent with a previous report.

![Fig. 4. Subcellular localization of a series of C-terminal–truncated mutants of UGT2B7 in Sf9 infected cells. (A) Schematic sequences of C-terminal–truncated UGT2B7 mutants. The numbers represent the residue position of WT UGT2B7 counted from the N-terminus. An HA-tag was conjugated to the C-terminus for detection by immunochemical methods. SP, signal peptide. (B) Immunoblotting to detect UGT2B7 mutants in insect cells. Sf9 cells transfected with recombinant bacmids to obtain P1 viruses were lysed and analyzed. After the collection of P1 viruses, the pelleted cells were resuspended in PBS, and an aliquot of the suspension was mixed with 2× SDS-PAGE sample buffer to prepare whole-cell lysates. WT-HA and Mock represent UGT2B7-HA microsomes (5 μg) and control microsomes (10 μg), respectively. Anti-HA-tag antibody was used as a primary antibody. (C) Fractionations of Sf9 cells to determine the localization of the UGT2B7 mutants. Human CNX was also transfected in cells expressing UGT2B7 deletion mutants. Sf9 homogenates were fractionated by sequential centrifugation. Protein amounts of loaded fractions were as follows: homogenate (Homo), 15 μg; nuclei & mitochondria (Mt), 6 μg; microsomes (MS), 5 μg; cytosol (Cyt), 10 μg. Anti-HA antibody and anti-CNAX antibody were used as primary antibodies. Each band was quantified with ImageJ software and shown as a percentage of the total.](image-url)
demonstrating that the proper positioning of two lysine residues from the C-terminus is critical for the retention of ER resident membrane proteins (Jackson et al., 1990). Collectively, our data suggest that the DM within the cytoplasmic domain of UGT2B7 is sufficient to localize CD4 to the ER but is not necessary for the ER retention of UGT2B7.

The results described above led us to speculate that the luminal domain of UGT2B7 is required for ER retention. In addition, it has been reported that the 141–240 region of human UGT1A6 is another membrane-associated domain and functions as an ER retention signal (Ouzzine et al., 1999). When this region was transplanted to the cytoplasmic enhanced GFP, the UGT 141–240/enhanced GFP chimera was localized to the ER. Based on these findings, we first focused on the N-terminal half-domain of UGT2B7 (24–252; except signal peptide) to explore the regions necessary for ER retention. We generated five deletion mutants (Δ1-HA to Δ5-HA) systematically lacking 30 residue sections containing two or three predicted helices from the C-terminal end of UGT2B7 (24–252), and an HA-tag was added to the C-terminal end of each deletion mutant (Fig. 4A). We transiently expressed these UGT2B7 mutants in Sf9 insect cells and confirmed their expression by immunoblotting (Fig. 4B). We further performed subcellular fractionation of insect cells expressing these deletion mutants using the same procedure, as in Fig. 2C. The results showed that all of the mutants were exclusively detected in the microsomal fraction with coexpressed CNX (Fig. 4C). Moreover, consistent with the results of subcellular fractionation, the results of immunofluorescence microscopy in COS-1 cells also showed that all of the deletion mutants colocalized with CNX in reticular-like structures (Fig. 5A), and that 90% of these mutants colocalized with CNX (Fig. 5B). Taken together, these results suggest that at least the N-terminal 24-133 region may be necessary for the retention of UGT2B7 in the ER.

To verify the necessity of the 24-133 residue region for ER retention of UGT2B7, we created a deletion mutant of this domain (Δ6-HA) and analyzed its expression and localization in COS-1 cells (Fig. 6, A–C). To our surprise, UGT2B7 Δ6-HA localized to CNX-positive reticular-like structures. We further generated a series of deletion mutants of the UGT2B7 luminal domain (Fig. 6A). These deletion mutants were transiently expressed in COS-1 cells, and their expression and localization analyzed using immunoblotting and immunofluorescence microscopy, respectively (Fig. 6, B and C). As shown in Fig. 6C, despite the systematic truncation, all of the mutants were observed in the reticular-like structures and colocalized with CNX. Moreover, 80%–95% of the serial mutants colocalized with CNX (Fig. 6D) as same as UGT2B7 WT (Fig. 3C). Collectively, these data imply that UGT2B7 could localize in the ER without any signal sequence rather than with the presence of a possible signal for the retention to the ER. Moreover, it is well known that proteins that aggregate in cells are largely detergent insoluble (Imai et al., 2001; Hirota and Tanaka, 2009). When we solubilized microsomes expressing these deletion mutants with 1% Triton X-100, they were predominantly recovered in solubilized fractions (Fig. 7A), indicating that they were properly folded, and thereby ruling out the possibility that the ER localization of these mutants resulted from accumulation in the ER by unfolding or misfolding. Moreover, we examined the stability of UGT2B7-HA deletion mutants (ΔCT and ΔTM) using inhibitors of proteasome and lysosomal degradation. Treatment with MG132, a proteasome inhibitor, but not with lysosomal protease inhibitors, increased protein levels of WT, ΔCT, and ΔTM of UGT2B7 (Fig. 7, B and C), thereby suggesting that the deletion of both the cytoplasmic and TM domains did not affect the stability of mutant proteins.

Deletions of the TM domain of membrane proteins transported by the secretory pathway should result in their secretion into the medium, unless they contain a retention signal in the remaining domain (Pääbo et al., 1986, 1987). We examined whether both cytoplasmic– and TM domain–deleted mutants of UGT2B7 and CD4 were secreted. Although all of the deletion mutants of UGT2B7-HA (ΔTM, Δ1, and Δ7) were
detected only in cell lysates, CD4-ΔTM was significantly secreted into the cultured medium (Fig. 7D).

**Discussion**

The DM is one representative peptide for motif-based retrograde transportation, but it is controversial whether this motif is indeed necessary for ER localization of UGTs. In this study, we present evidence that C-terminal deletion mutants of UGT2B7, including ΔDM, ΔCT, and ΔTM, can localize to the ER in their folded states (Fig. 2C; Fig. 3, B and C; Fig. 7). It has been demonstrated previously that the proper positioning of two lysines at −3 and −4 or −5 from the C-terminus is critical for the retention of ER resident membrane proteins, because the addition or deletion of residues from the C-terminus decreased retention efficiency (Jackson et al., 1990). Indeed, we also confirmed this finding, using the chimeric protein CD4-UGT (Fig. 3, D and E). However, our findings that not only UGT2B7ΔDM, but also UGT2B7-HA, in which the positions of two lysine residues were shifted from −3 and −5 to −12 and
and −14 by adding an HA-tag to the C-terminus, localized to the ER (Fig. 3B), suggest that the C-terminal DM of UGT2B7 is unnecessary for its ER localization, supporting previous reports (Meech et al., 1996; Meech and Mackenzie, 1997; Ouzzine et al., 1999). Of note, there are ER-located splice variants of human UGT, which lack the TM and cytoplasmic domains (Girard et al., 2007; Lévesque et al., 2007; Bellemare et al., 2010; Guillemette et al., 2014; Rouleau et al., 2014).
Similar variants of UGT2B7 showed ER localization despite lacking the DM (Ménard et al., 2013). These findings also reinforce our conclusion that the cytoplasmic region including the DM is not required for ER retention.

In contrast, previous studies have brought attention to the role of the DM as a retrieval signal for UGT. Transplantation of the motif altered subcellular localizations of some plasma membrane proteins to the ER (Jackson et al., 1990; Kinosaki et al., 1993). Moreover, it was reported that the interaction of rat UGT2B1 with β-COP, a subunit in the COPI complex, was abolished by deletion of the C-terminal cytoplasmic tail of UGT2B1 (Meech and Mackenzie, 1998). Given that UGT2B7 is statically retained in the ER, what is the physiologic role of the C-terminal region containing the DM? One possibility is to retrieve UGT2B7 that has escaped to post-ER compartments via nonspecific transport known as “bulk flow.” Details for the transport of proteins from the ER by bulk flow remain unclear (Barlowe and Helenius, 2016). Retrieval mediated by the DM should return UGT2B7 to the ER against such a leak mechanism. The importance of this region on catalytic activity was also reported with another UGT isoform. The activity of a water-soluble UGT1A9 mutant lacking the TM and cytoplasmic domains (UGT1A9-sol), which were purified from an insect cell expression system, was one order of magnitude lower than microsomal WT UGT1A9 (Kurkela et al., 2004). In addition, we have reported previously that the C-terminal region of UGT2B7 is important for its functional interaction with CYP3A4 (Miyauchi et al., 2015). UGT2B7 WT significantly suppressed CYP3A4 activity, but this suppression disappeared with ΔCT and ΔTM. These lines of evidence clearly indicate a novel role for the UGT C-terminal region as a domain regulating UGT activity and the activity of other drug-metabolizing enzymes, which may improve our understanding of the large interindividual differences in the in vivo catalytic activity of UGT and cytochrome P450 (Shimada et al., 1994; Lamba et al., 2002; Court, 2010).

ER controls the quality of the synthesized proteins, and unfolded and misfolded proteins are degraded by ER-associated degradation (Nakatsukasa and Brodsky, 2008). In this study, we showed that all UGT mutants were predominantly collected in the soluble fraction, thereby suggesting that they were folded correctly (Fig. 7A). Moreover, there was no difference in protein stability among WT and the mutants. These results demonstrated that the ER localization of the mutants is not due to protein misfolding.

Serial mutation analyses suggest that retention or retrieval signals are not needed for ER localization of UGT2B7 in this study. Although many ER membrane proteins do not bear a DM in their C-terminal ends, they are positioned in the ER,
indicating the presence of another mechanism underlying ER localization of membrane protein (Teasdale and Jackson, 1996). One postulated mechanism is that the proteins should have a novel retrieval/retention signal. Previous studies have suggested the presence of ER retention signals of UGT isoforms in either the luminal domain or the cytoplasmic and TM domains (Table 1). However, all of the UGT2B7 deletion mutants we have constructed were observed in the ER without being secreted (Fig. 7D). Thus, it seems more reasonable to assume that UGT2B7 lacks an active signal required for exit from the ER, which results in passive and static retention in the ER (Fig. 8), although little is known about such an export signal.

Further studies are necessary to clarify the molecular mechanism underlying UGT localization in the ER, which may improve our understanding of how ER membrane proteins reside in their respective membrane compartments without retention and/or retrieval motifs.

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