Membrane Topology and Intracellular Processing of Cyclin M2 (CNNM2)*

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Background: Mutations in CNNM2 cause severe dominant hypomagnesemia. Structure of CNNM2 consists of an extracellular N terminus and intracellular C terminus containing CBS domains, which are affected by the identified mutations.

Results: Structure of CNNM2 was exposed in detail. In the endoplasmic reticulum, CNNM2 was glycosylated at residue Asn-112, stabilizing CNNM2 on the plasma membrane. Interestingly, co-immunoprecipitation studies evidenced that CNNM2a forms heterodimers with the smaller isoform CNNM2b. These new findings on CNNM2 structure and processing may aid to elucidate CNNM2 function in renal magnesium transport.

As the second most abundant intracellular cation in humans, magnesium (Mg²⁺) constitutes an essential co-factor in vital processes such as energy metabolism, DNA transcription, and protein synthesis. Balance between intestinal absorption and renal excretion is tightly regulated to keep the plasma Mg²⁺ concentration in its physiological range (0.7–1.1 mM) (1). Specifically, bulk Mg²⁺ reabsorption in kidney from the pro-urine is dependent on passive, paracellular transport in the proximal tubule and the thick ascending limb of Henle’s loop, whereas fine-tuning is achieved by active, transcellular transport in the distal convoluted tubule (DCT)² (1). In the latter segment, tight regulation of Mg²⁺ transport will determine the final urinary Mg²⁺ excretion because no Mg²⁺ reabsorption takes place beyond the DCT. The TRPM6 ion channel, expressed at the apical side of the DCT, constitutes the gatekeeper in active Mg²⁺ uptake (2), whereas a basolateral Mg²⁺ extrusion mechanism remains to be identified.

Recently, mutations in the cyclin M2 (CNNM2) gene were described leading to a dominant form of hypomagnesemia caused by renal Mg²⁺ wasting (3). Patients suffer from muscle weakness, tremor, and headaches accompanied by low Mg²⁺ serum concentrations (0.3–0.5 mM) (3, 4). Interestingly, no other electrolyte disturbance was detected (3). CNNM2, formerly known as ancient conserved domain protein 2 (ACDP2), belongs to the CNNM family consisting of four proteins (CNNM1–4) that share homology to cyclins, although no cyclin-related function has been described (5). Two predicted CNNM2 protein isoforms are conserved between humans and mouse. Although the 875-amino acid CNNM2a is encoded by splice variant 1 (Cnnm2v1, NM_017649), the 22-residue-shorter isoform b is translated from splice vari-
ant 2 (Cnnm2v2, NM_199076), which lacks exon 6. CNNM2 expressed in *Xenopus laevis* oocytes and in the Mg$^{2+}$-deficient *Salmonella enterica* strain MM281 has been suggested to be involved in Mg$^{2+}$ transport (6, 7). However, electrophysiological analysis using mammalian cells expressing CNNM2 showed Mg$^{2+}$-sensitive Na$^+$ currents instead of Mg$^{2+}$ currents (3). This last finding could indicate that instead of transporting Mg$^{2+}$ itself, CNNM2 might be involved in a Mg$^{2+}$-sensing mechanism that, in turn, could regulate renal Mg$^{2+}$ uptake.

At present, no studies have been reported showing the structure of CNNM2 or any of the other CNNM family members. The aim of this study was to elucidate the membrane topology of CNNM2 and to provide a molecular characterization of the intracellular processing mechanisms.

**EXPERIMENTAL PROCEDURES**

**Expression Profiling**—Three C57BL/6 mice were sacrificed; kidney, duodenum, ileum, jejunum, colon, cecum, brain, lung, liver, spleen, testis, muscle, and heart tissues were collected. The transgenic parvalbumin-EGFP mice were a gift kind from Dr. Monyer (University of Heidelberg, Germany (8)), and TRPV5-GFP mice were kindly provided by Dr. Praetorius from the University of Aarhus, Denmark (9). They were used to isolate the DCT and connecting tubule (CNT), respectively. Mice were anesthetized by a mixture injection of ketamine (0.1 mg/g of body weight) and xylazine (0.01 mg/g of body weight) and were perfused with 20 ml of PBS (in mM: 137 NaCl, 2.7 KCl, 10 Na$_2$HPO$_4$, 1.76 KH$_2$PO$_4$) through the heart. The kidneys were removed, minced, and digested in collagenase (1 mg/ml collagenase A (Worthington), 1 mg/ml hyaluronidase) in Krebs buffer (in mM: 145 NaCl, 5 KCl, 10 HEPEs, 1 Na$_2$HPO$_4$, 2.5 CaCl$_2$, 1.8 MgSO$_4$, 5 glucose) at 37 °C for 18 min. The collagenase-digested tubules sized between 40 and 100 μm were sorted based on GFP fluorescence by COPAS (Complex Object Parametric Analysis and Sorting, Union Biometrica).

Total RNA was isolated using TRIzol total RNA isolation agent (Invitrogen). Obtained RNA was treated with DNase (Promega) to remove genomic DNA. Subsequently, reverse transcription of the RNA by M-MLV reverse transcriptase (Promega) to remove genomic DNA. Subsequently, reverse transcription was performed for 1 h at 40 °C. Gene expression levels were determined by quantitative real-time PCR on a Bio-Rad analyzer and normalized for Gapdh expression levels. Primer sequences are provided in supplemental Table 1.

**Immunohistochemistry**—Immunohistochemistry was performed as described previously (10). In short, co-staining for CNNM2 with Na$^{+}$-K$^{+}$-Cl$^{-}$ cotransporter (NKCC2), thiazide-sensitive Na$^{+}$-Cl$^{-}$ cotransporter (NCC), calbindin-D$_{28k}$, and aquaporin-2 (AQP2) was performed on 7-μm sections of fixed frozen mouse kidney samples. The sections were incubated for 16 h at 4 °C with the following primary antibodies: guinea pig anti-CNNM2 (1:100 3)), rabbit anti-NKCC2 (1:100 (39)), rabbit anti-NCC (1:100 40), rabbit anti-calbindin-D$_{28k}$ (1:500 (41)), and rabbit anti-AQP2 (1:300, kindly provided by Dr. Deen, Nijmegen, The Netherlands). For detection, kidney sections were incubated with Alexa Fluor-conjugated secondary antibodies. Images were taken with an AxioCam camera (Zeiss) and AxioVision software (Zeiss).

**DNA Constructs**—Mouse Cnnm2 was cloned into the pCINeo HA IRES GFP vector as described previously (3). To obtain a C-terminal vesicular stomatititis virus epitope, an XbaI restriction site was introduced before the stop codon using the QuikChange site-directed mutagenesis kit (Agilent, Amstelveen, The Netherlands) according to the manufacturer’s protocol. Subsequently, oligonucleotides (forward, 5′-GATGCGGCTCTTGGGAAGTGAGCTCGAGAATTCCGCCCC-3′, reverse, 5′-GGGGCGGAATTCGTCACTTCCCCAAGCGATTCC-3′) encoding the vesicular stomatititis virus epitope were hybridized and ligated into the pCINEO HA-CNNM2 IRES GFP vector using the XbaI and XhoI restriction sites. CNNM2 mutations were inserted in the construct using the QuikChange site-directed mutagenesis kit according to the manufacturer’s protocol. Human CNNM4 full-length cDNA clone (GenBank™ accession number BC063295) was amplified using Phusion polymerase (Finnzymes, Vantaa, Finland) and primers (forward, 5′-CGGCCTAGCGCCACCATGGGCGCCTGTTGG-3′, reverse, 5′-CGACCGGTCTATGCGTAGTTCTGGCACTGTGATAGGTAGTGAGATGAGATGTACTCTGTTGGAG-3′) to introduce an in-frame HA tag before the stop codon and Argel and Nhel restriction sites. The ampiclons were cloned into the pCINEO IRES GFP expression vector using the Argel and Nhel restriction sites. All constructs were verified by sequence analysis.

Mouse Cnnm2 was cloned into pcDNA3 (Invitrogen) as described previously (3). To obtain splice variant 2 with a 66-nucleotide deletion corresponding to exon 6, site-directed mutagenesis (QuikChange, Agilent) was used, according to the two-stage PCR mutagenesis protocol described by Wang and Malcolm (11). The same protocol was used to delete the C-terminal tag and to insert internal HA tags with an additional serine residue (NH$_2$-YPYDVPDYAS-COOH) (12) after residues Thr-24, Gly-210, or Pro-744 or to replace the C-terminal HA tag with a FLAG sequence. Mutagenesis primer sequences are available upon request. To obtain an N-terminal HA-tagged expression clone, the ORF in pFLCI-mCnnm2WT (3) was amplified using Expand polymerase (Roche Applied Science) using forward primer 5′-ATGGGGTAGGCCATGGTATGGCGACTCAGCTGTGATAGTGAGATGAGATGTACTCTGTTGGAG-3′ containing an in-frame HA tag after the Kozak sequence and reverse primer 5′-ATCTTCTAGAGTTTCACTGAGGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGA
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**Immunocytochemistry**—COS-7 cells were seeded on glass coverslips and transfected using Lipofectamine 2000 according to the manufacturer’s protocol with constructs expressing epitope-tagged CNNM2 or CNNM4. After 2 days of growth in medium supplemented with 20 mM MgCl₂, the cells were epitope-tagged CNNM2 or CNNM4. After 2 days of growth in medium supplemented with 20 mM MgCl₂, the cells were fixed for 10 min with 4% (w/v) methanol-free formaldehyde solution (Thermo Scientific) in PBS. Cells in a control experiment (supplemental Fig. S1, lower panels), living cells were incubated with primary antibody rat anti-HA (Roche Applied Science, high affinity 3F10, 1:250) for 60 min at 4 °C before formaldehyde fixation. After rinses in PBS, some of the cells were incubated for 10 min with 0.2% (v/v) Triton X-100 in PBS for permeabilization. After PBS rinses, the cells were incubated with blocking buffer (5% (v/v) donkey serum (Sigma) in PBS) for 1 h and immunolabeled with the primary antibodies rat anti-HA (1:250), mouse anti-FLAG M2 (Sigma, 1:250), rabbit anti-calnexin (C5C9, Cell Signaling Technology, 1:50), or rabbit anti-CNNM2 (Abcam, Ab111351, 1:50) in blocking buffer for 90 min. After rinses in PBS, cells were incubated with Alexa Fluor 488-conjugated donkey anti-rabbit or anti-mouse IgG or Alexa Fluor 594-conjugated donkey anti-rat IgG secondary antibodies (Invitrogen, 1:500) and DAPI (2 µg/mL) for 45 min. After PBS and ethanol rinses, the cells were mounted on slides using ProLong Gold antifade reagent (Invitrogen). Fluorescence microscopy was performed with a Leica DMIRE2 inverted microscope, and images were taken with the Openlab software.

**Homology Modeling**—A homology model was built using the modeling script in the WHATIF and YASARA Twissnet with standard parameters (14, 15). We used the structure of a cystathionine B-synthase (CBS) domain pair structure of the Mg²⁺ and Co²⁺ efflux protein CorC from *Bordetella parapertussis* (Protein Data Bank (PDB) file 3jtf) as a template for modeling.

**Cell Surface Biotinylation, PNGase Treatment, and Immunoblotting**—Biotinylation experiments were performed as described previously (16). In short, HEK293 cells were transfected for 48 h, and cell surfaces were biotinylated at 4 °C for 1 h by adding sulfo-NHS-LC-LC-biotin. Subsequently, protein lysates were incubated overnight with NeutrAvidin-agarose beads (Pierce).

Protein lysates were denatured in Laemmli containing 100 mM DTT for 30 min at 37 °C and subsequently subjected to SDS-PAGE. Then, immunoblots were incubated with mouse anti-HA (Cell Signaling Technology) primary antibodies and peroxidase conjugated sheep anti-mouse secondary antibodies (Sigma). PNGase F (New England Biolabs) was added to the protein lysates in Laemmli–DTT buffer and incubated at 37 °C for 1 h before loading the protein samples on gel.

Co-immunoprecipitation—COS-7 cells were cultured and transfected as described above with expression vectors for HA- and FLAG-tagged wild type CNNM2 or CNNM4 isoforms. 48 h after transfection, cells were scraped from dishes in ice-cold 50 mM Tris (pH 8.0) followed by centrifugation at 5000 × g. The pellets were lysed in lysis buffer (150 mM NaCl; 0.1% (v/v) Nonidet P-40; 50 mM Tris, pH 7.5; and Complete protease inhibitor mixture (Roche Applied Science)) by repeated passage through a 26-gauge needle. Protein concentration measurement was performed using a BCA assay (Pierce). The next incubation steps were all done under rotary agitation at 4 °C. Protein aliquots of 100 µg in lysis buffer were precleared by incubation with protein G-Sepharose (Sigma-Aldrich) for 2 h after which the precleared lysate was incubated for 18 h with 2.5 µg of anti-HA antibody (Roche Applied Science, high affinity 3F10). The antibody-lysate mixture was added to 50 µL of fresh protein G-Sepharose beads and incubated for 1 h. The beads were collected by centrifugation at 1000 × g for 1 min at 4 °C and washed four times with lysis buffer. The proteins were separated from the beads by boiling for 5 min in 1× Laemmli sample buffer and detected by immunoblotting using a mouse monoclonal antibody against FLAG M2 (Sigma-Aldrich, 1:1000).

**In Vitro Transcription/Translation**—The cDNA template for *in vitro* transcription contained the Cnnm2-HA sequence downstream of a T7 promoter sequence in a pT7Ts vector. The coupled *in vitro* transcription/translation reaction was incubated at 30 °C for 90 min. The reaction mixture contained 2 µg of cDNA template, 2 µL of reaction buffer, 1 µL of T7 RNA polymerase, 1 µL of amino acid mix without methionine, 3 µL of [³⁵S]Met (1 µCi/µL), RNasin, and 25 µL of rabbit reticulocyte lysate (Promega). The reaction was performed with or without signal peptidase inhibitor MeoSuc-Ala-Ala-Pro-Val-chloromethylketone (APAV-CMK, Sigma). The protein mixture was separated on an 8% (w/v) SDS-PAGE gel. The gel was fixed in a mixture of acetic acid, methanol, and H₂O (20:10:70) and incubated in 20% (w/v) 2,5-diphenyloxazole in dimethyl sulfoxide (DMSO) to intensify the radioactive signal. The proteins were visualized using CL-XPosure films (Pierce, Etten-Leur, The Netherlands).

**Statistical Analysis**—In all experiments, data are expressed as means ± S.E. Statistical significance was determined using an unpaired Student’s *t* test, and differences with *p* < 0.05 were regarded as statistically significant.

**RESULTS**

**Expression Profile of CNNM Family**—To examine the transcript expression levels for CNNM1–4 in various tissues, a mouse tissue cDNA panel was constructed, and *Cnnm* mRNA expressions were quantified by real-time PCR analysis. *Gapdh* expression was used to normalize values. *Cnnm3v1* and *Cnnm3v2* (containing an alternative last exon) show a ubiquitous expression pattern with highest expression in kidney, brain, lung, spleen, and heart (Fig. 1, D and E). *Cnnm1* expression is predominantly found in brain and testis (Fig. 1A), whereas *Cnnm4* shows highest expression along the gastrointestinal tract (Fig. 1F). Interestingly, *Cnnm2* shows a ubiquitous expression pattern with highest expression in kidney, lung, spleen, and testis (Fig. 1, B and C). CNNM2A and CNNM2B (encoded by *Cnnm2v1* and *Cnnm2v2*, lacking exon 6) share a similar pattern; only minor differences between the tissue expression levels were measured.

Because CNNM2 has been described to be causative for hypomagnesemia due to excessive renal Mg²⁺ wasting (3), its kidney localization was investigated in detail. Using immunohistochemistry, co-expression of CNNM2 with markers of different distal nephron segments was examined: the Na-K-Cl cotransporter NKCC2 (thick ascending limb of Henle’s loop), the thiazide-sensitive Na-Cl cotransporter NCC (DCT), calcin-
din-D28k (CNT and cortical collecting duct), and aquaporin-2 (collecting duct). CNNM2 was expressed on the basolateral side of the cells (Fig. 2, A–D, green), opposed to the luminal expression of the marker proteins (Fig. 2, A–D, red). CNNM2 immunopositive tubules were negative for aquaporin-2 staining, but did co-stain with NCC-positive tubules. Only a part of the tubules positive for calbindin-D28k showed CNNM2 staining. Although most of NKCC2-positive tubules are negative for CNNM2, only a few NKCC2-positive tubules show faint CNNM2 staining. These results indicate that CNNM2 is predominantly expressed in DCT. mRNA expression levels measured by real-time PCR on GFP-sorted DCT and CNT tubules showed an equivalent pattern; expression of Cnnm2v1 and Cnnm2v2 was high in DCT and to a lesser extent also in the CNT (Fig. 2E). As a control for DCT and CNT specificity of the mRNA, we examined the expression of the transcripts for the marker proteins TRPM6 and TRPV5, respectively (Fig. 2E). These and previous results (3) indicate that CNNM2 is predominantly expressed basolaterally in DCT and also, but in lower amounts, present in thick ascending limb of Henle’s loop and CNT (Fig. 2F).

**CNNM2 Has an Intracellular C Terminus Containing CBS Domains**—The consensus topology prediction for the plasma membrane protein CNNM2 indicates five potential transmembrane (TM) domains and an extracellular C terminus (3), which would locate the CBS domains (see below) in an unprecedented, extracellular position. We decided to investigate the topology in detail using an intramolecular epitope strategy by expression of various epitope-tagged CNNM2 proteins (Fig. 3A) in COS-7 cells with subsequent immunocytochemistry. A similar approach was used successfully to aid in the topology determination of other plasma membrane transporters such as multidrug resistance proteins (17) and an iron transporter (18). Immunocytochemistry of cells overexpressing CNNM2a proteins with HA epitopes at positions 211 (HA211), 745 (HA745), or at the extreme C terminus revealed these proteins to be targeted to the plasma membrane of the cells (Fig. 3B, upper panels), similar to the results obtained in MDCK-C7 cells (3). Importantly, C-terminally HA-tagged CNNM2b showed a localization that is undistinguishable from that of CNNM2a, indicating that the absence of the amino acids translated from exon 6 does not influence sorting to the plasma membrane.

To determine whether the epitopes are located on the inside or the outside of the cells, formaldehyde-fixed cells, both intact and permeabilized, were immunostained for the HA epitope. The HA tag at position 211 could be detected on intact cells (Fig. 3B, lower panels), indicating extracellular localization. On the contrary, detection of HA745 or the epitope at the C termi-
nus required cell permeabilization, suggesting that these latter two are located intracellularly. Similar results were obtained when living cells were incubated with anti-HA antibodies prior to fixation (supplemental Fig. S1).

As an alternative approach to confirm that the C terminus is located intracellularly, cells that expressed HA211 were immunostained with Abcam antibody 111351, raised against amino acids 590–640, located just after the pair of CBS domains. Also here, the epitope could only be detected after permeabilization (Fig. 3C), indicative for an intracellular position of the C terminus that importantly includes the two CBS domains.

To investigate whether the intracellular localization of the C terminus is shared with other members of the CNNM family, the same permeabilization assay was performed on cells transfected with C-terminally HA-tagged CNNM4 (Fig. 3D). Again, the plasma membrane staining was only observed after permeabilization, indicating that also for CNNM4, the C terminus resides inside the cell. In conclusion, the CNNM2 protein has a membrane topology with its C terminus inside of the cell.

CNNM2 N Terminus Contains a Large Signal Peptide and a Glycosylation Site—We observed that the HA epitope at the extreme N terminus or at position 25 of CNNM2a was not...
detected at the plasma membrane, but rather in distinct perinuclear structures (supplemental Fig. S2), which suggested that the N terminus might contain a signal peptide and, therefore, ultimately results in cleavage of the HA epitope. To exclude incorrect processing of the recombinant protein, a construct was expressed with the HA epitope at position 25 of CNNM2a and a FLAG epitope at the C terminus. We observed that the FLAG epitope was correctly targeted at the plasma membrane, but the HA epitope was retained in perinuclear structures (Fig. 3E). Colocalization studies with an antibody against calnexin (supplemental Fig. S2) showed that these perinuclear structures are part of the endoplasmic reticulum (ER), which is in line with the presence of a signal peptide.

Signal peptide prediction software (SignalP (19)) predicts a long N-terminal signal peptide of 64 amino acids. Examination of the N-terminally HA-tagged CNNM2 construct by immunoblotting confirmed that the CNNM2 N terminus is cleaved (Fig. 4A, upper panel) because the HA tag was undetectable. Mutation of the predicted cleavage region (62GCTA65) to hydrophobic residues 62LLV65 marred this cleavage event, evidenced by
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In eukaryotic cells, trafficking of membrane proteins to the plasma membrane is often regulated by glycosylation of asparagine residues present in extracellular domains. To determine the CNNM2 glycosylation profile, four potential candidate asparagine residues were mutated to alanine, namely Asn-112, Asn-327, Asn-527, and Asn-591 (based on prediction software NetNGlyc 1.0, Center for Biological Sequence Analysis (CBS)). Subsequently, the presence of N-linked glycans in the mutants was discerned by immunoblotting the proteins after treatment with PNGase F, an enzyme causing N-glycan cleavage. The glycosylated wild type CNNM2 runs at ~105 kDa, but PNGase F treatment reduced its molecular mass to ~96 kDa (Fig. 5A). Of all the candidates, only residue Asn-112 was identified as a glycosylation target in CNNM2 because the N112A mutation was observed when compared with wild type CNNM2 protein (Fig. 5B, right panel), suggesting that Asn-112 glycosylation is necessary for CNNM2 membrane stability. Taken together, the CNNM2 extracellular N terminus provides important sites for post-translational modifications necessary for proper plasma membrane expression.

C-terminal CBS Domains Are Disrupted by T568I Mutation—One of the mutations recently described to be causative for hypomagnesemia is located in the CBS domains within the C terminus (3). Therefore, emphasis was placed on determining the structure of the intracellular CBS domains (Fig. 3A) that have been associated with ATP binding, among other functions (20). Here, we provide a homology model of the CNNM2 CBS domains based on the B. parapertussis Mg²⁺/Co²⁺ CorC protein (Fig. 6A). Analysis of this model showed conservation of the potential ATP-binding site in the CBS dimer of CNNM2 (Fig. 6A). Interestingly, the T568I mutation, identified in a Czech family to be causative for hypomagnesemia (3), is lining the ATP-binding site in the CBS domain (Fig. 6B). In our model, mutation of the threonine into the bigger isoleucine residue causes steric bumps with the ATP molecule. Besides that, Thr-568 also forms a hydrogen bond that stabilizes the position of the Glu-570 and Asp-571 residues (Fig. 6B). These two residues form important hydrogen bonds and ionic interactions to the ATP molecule and residues lining the ATP-binding pocket, respectively. According to the model, the T568I mutation of the patient will also affect the position of the Glu-570 and Asp-571 residues and could, therefore, severely affect ATP binding. In conclusion, this model might explain why the T568I mutation of the patient is causative for nonfunctional CNNM2.

CNNM2a and CNNM2b Form Multimers—In GenBank, two splice variants exist that are conserved between human and mouse, the first being full length, the second lacking exon 6. Interestingly, splice variant 2 was consistently found in conjunction with that of splice variant 1 in all tissues examined, including kidney (Fig. 1). Further examination of potential interactions between CNNM2a and CNNM2b was performed by studying the dimerization of CNNM2 isoforms. Interestingly, besides the CNNM2 monomer (with an expected molecular mass of 105 kDa) (5), additional complexes were visualized in the immunoblot at ~200 kDa (Fig. 7A). This molecular mass matches the mass of a hypothetical CNNM2 dimer. To confirm CNNM2 oligomerization events, co-immunoprecipitation experiments were performed using anti-HA antibody. To avoid signal peptide cleavage, as shown by a higher molecular weight on gel (Fig. 4A), a double-tagged CNNM2-HA was transfected in HEK293 cells against the N-terminal HA tag (upper blot). As a control for CNNM2 expression, the noncleavable CNNM2 C terminus was targeted by immunoblotting against the vesicular stomatitis virus tag (lower blot). The blot shown is representative of five independent experiments. WT, wild type; MU, mutant. 8, coupled in vitro transcription/translation protein synthesis of the CNNM2-HA protein in the presence (+) and absence (−) of SPC inhibitor AAPV-CMK. The gel shown is representative of three independent experiments.
assays of CNNM2 isoforms were performed. CNNM2a-HA and CNNM2b-HA were each able to co-immunoprecipitate both CNNM2a and C-terminally FLAG-tagged CNNM2b. CNNM4-HA only weakly co-immunoprecipitated, although it was highly expressed, indicating the specificity of these interactions (Fig. 7B). These data show that CNNM2a and CNNM2b are able to form specific homo- and heteromultimeric complexes.
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FIGURE 7. CNNM2a and CNNM2b form homo- and heterotypic interactions. A. Western blot analysis of CNNM2 protein showing CNNM2a and CNNM2b monomers at ~100 kDa and potential CNNM2 dimers at ~200 kDa. B. Co-immunoprecipitation (IP) studies of CNNM proteins in COS-7 cells. C-terminal FLAG-tagged CNNM2a or CNNM2b were co-expressed with C-terminal HA-tagged CNNM2a, CNNM2b, or CNNM4 as indicated in the two top rows. The lower two blots show the detection of the FLAG-tagged proteins in anti-HA precipitated cell lysates. The lower two blots show input controls (10%) of HA-tagged and FLAG-tagged proteins, respectively. The results shown are representative of three individual experiments. IB, immunoblot.

DISCUSSION

Mutations in CNNM2 have, recently, been described to be causative for a severe form of hypomagnesemia (3). Initially described as nuclear proteins involved in cell cycle regulation (5), the family of CNNM proteins has now been shown to reside at the plasma membrane (3, 6). Although their physiological role is still poorly understood, all members are likely involved in electrolyte homeostasis. CNNM1 has been identified as a cytosolic chaperone, CNNM2 was shown to mediate Mg2+ -sensitive Na+ currents in HEK293 cells, and Cnnm2, Cnnm3, and Cnnm4 have been linked with serum Mg2+ levels (3, 21, 22). Furthermore, Cnnm2 has been associated with coronary artery disease and hypertension (23–25). Finally, mutations in Cnnm4 were shown to be causative for recessive cone-rod dystrophy with amelogenesis imperfecta (26, 27). In the present study, we aimed to elucidate the CNNM2 protein topology and characterize its post-translational modifications.

Although Gómez García et al. (28) have been working on structural elucidation of the CBS domains of CNNM4, complete structural information for any of the CNNM proteins is presently unknown. Using immunocytochemical methods, we examined the membrane topology of CNNM2. Based on our results showing an extracellular N terminus and an intracellular C terminus (Fig. 3), we propose a structure comparable with that of the glutamate receptor (GluR) ion channels (29), containing three full membrane-spanning regions and an additional re-entrant loop. In silico analysis of the CNNM2 protein sequence showed that of the four hydrophobic regions, the second one is the shortest and the least hydrophobic, indicating that the second hydrophilic region might not be completely membrane-spanning, but instead forms a re-entrant loop (Fig. 8).

Similar to the glutamate receptor family, CNNM2 contains an N-terminal signal peptide that is cleaved in the ER. Here, the SPC seems to be the protease complex responsible for CNNM2 cleavage (Fig. 4B). The human SPC is a complex of five proteins expressed on the ER membrane and recruited post-translationally to the signal recognition particle to perform signal peptide cleavage (30). Our results do not identify the exact position of the cleavage site. Signal peptide recognition is known to be dependent on the residues at positions −1 and −3 of the actual cleavage site (31). Nevertheless, conversion of the −1 and −3 position of the predicted site (Gly-62/Thr-64) to hydrophobic residues did not prevent cleavage. Likewise, the cleavage was also not impaired when the alternative cleavage site (Cys-63/Ala-65) was mutated. Only mutation of both sites prevented CNNM2 cleavage, suggesting that the SPC can cleave CNNM2 at the preferential site between Thr-64 and Ala-65, but also at an alternate site between Ala-65 and Ala-66. Thus, CNNM2 contains a remarkable lengthy signal peptide of ~64 amino acids. Long signal peptides have been associated with multiple functions, but might serve as a mechanism to achieve proper membrane insertion of the protein (32). This also suggests that the first predicted transmembrane domain (Fig. 3A) actually represents the hydrophobic core of the signal peptide that, after cleavage, will remain at the ER membrane (Fig. 8).

After cleavage in the ER, CNNM2 is further processed by glycosylation in the Golgi apparatus. N-Glycosylation is known to facilitate plasma membrane trafficking for several membrane proteins (33). Here, Asn-112 was identified as the residue that is glycosylated and implicated in CNNM2 membrane stability (Fig. 5, A and B). This can be interpreted as an additional argument showing that the N terminus is located on the extracellular side of the membrane. Interestingly, our immunoblotting data showed a shift of the molecular weight of the CNNM2 protein when the alternative residue Asn-327 was mutated. However, PNGase F treatment still lowers the molecular weight of the N327A mutant, indicating that the protein is not deglycosylated (Fig. 5A). Because Asn-327 is predicted to be located in the second TM domain, we suspect that mutation of this residue instigates an altered protein folding that runs different on gel, explaining the observed shift.

Once at the plasma membrane, the functional CNNM2 unit likely represents a multimer. Interestingly, we showed that both CNNM2a and CNNM2b can form homomultimers as well as heteromultimers (Fig. 7B). This interaction is specific, as evi-
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FIGURE 8. Schematic model of CNNM2 structure. A model representing the structure of the CNNM2 protein at the plasma membrane and showing the signal peptide cleavage in the endoplasmic reticulum is illustrated. Crosses represent the locations of the mutations. The glycosylation site at position Asn-112 is shown in the extracellular N terminus. The CBS domains are represented in purple, and binding ATP is in red. SP, signal peptide; TM, transmembrane helix; N112, Asn-112.

CNNM2 are highly homologous to those found in the ing and ionic strength sensing (35, 36). The CBS domains in its role in a variety of functions such as adenosine phosphate bind-
domains are found in 100–200 mammalian proteins and play a identification of CNNM2 protein partners to further elucidate its role in Mg2+ homeostasis.

REFERENCES


dened by the fact that CNNM2a and CNNM2b hardly dimerize with CNNM4, which shares the highest sequence homology with CNNM2 (96.7% similarity and 81.4% identity at the protein level) (5).

In magnesium research, CNNM2 function is heavily debated. CNNM2 was first reported as a Mg2+ transporter (6, 7), but recent findings questioned the ability of CNNM2 to transport Mg2+ and proposed a Mg2+-sensing function (3). We, therefore, examined whether our topological findings could further elucidate the function of CNNM2. We reasoned that although CNNM2 might contain a re-entrant region, it is unlikely that this region forms a pore for Mg2+ transport. The re-entrant loop does not include negatively charged residues that could provide Mg2+ specificity. Furthermore, the protein topology of only three transmembrane segments would be small in comparison with known eukaryotic Mg2+ transporters such as TRPM6/7 (tetramers of six TM proteins) (2) and SLC41A1 (10/11 TM domains) (34). Thus, our topological findings strengthen the previously reported patch clamp data (3), suggesting that CNNM2 might be a Mg2+ sensor indirectly regul-
ulating Mg2+ transport.

To further substantiate this hypothesis, we focused on the CBS domains during analysis of the CNNM2 structure. CBS domains are found in 100–200 mammalian proteins and play a role in a variety of functions such as adenosine phosphate binding and ionic strength sensing (35, 36). The CBS domains in CNNM2 are highly homologous to those found in the B. parapertussis Co2+/Mg2+ protein CorC, of which the structure has been crystallized recently (PDB 3jtf). As shown by homology modeling based on the CorC structure, the CBS domains in the CNNM2 C terminus contain a potential ATP-binding domain (Fig. 4B). ATP binding in CBS domains has been reported to activate or further inhibit protein function in eukaryotic CBS domain-containing proteins (20).

Our model does not allow discrimination between ATP and Mg-ATP binding, but we expect that the CBS domains of CNNM2 are also able to bind Mg-ATP. This theory is strengthened by the recently elucidated crystal structure of the bacterial Mg2+ transporter MgtE in which its CBS domains are involved in Mg2+ binding (PDB 2yvy (37, 38). As the majority of intra-
cellular Mg2+ is ATP bound, the intracellular Mg-ATP concentrations are representative for the cellular Mg2+ content. The suggestion that CNNM2 binds Mg-ATP implies that rather than ATP being an energy source for active transport, the Mg-ATP binding would provide a Mg2+-sensing mechanism. Given that the CNNM2 T568I loss-of-function mutation found in patients (3) is located at the region predicted to form the ATP-binding pocket (Fig. 6B), the (Mg-)ATP binding is likely of major importance for CNNM2 function. Because the T568I mutation is specifically disturbing the ATP-binding pocket, Mg2+ alone will possibly not be able to activate CNNM2.

In conclusion, we propose a protein topology of CNNM2 consisting of three membrane-spanning domains and a re-entrant loop (Fig. 8). CNNM2 is extensively post-translationally modified by signal peptide cleavage and glycosylation before being functional in multimers at the plasma membrane. Rather than transporting Mg2+ itself, we hypothesize that CNNM2 would sense intracellular Mg2+ concentrations and regulate other Mg2+ transporters. Future research should focus on the identification of CNNM2 protein partners to further elucidate its role in Mg2+ homeostasis.
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