Laminin β2 Gene Missense Mutation Produces Endoplasmic Reticulum Stress in Podocytes

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ABSTRACT

Mutations in the laminin β2 gene (LAMB2) cause Pierson syndrome, a severe congenital nephrotic syndrome with ocular and neurologic defects. LAMB2 is a component of the laminin-521 (α5β2γ1) trimer, an important constituent of the glomerular basement membrane (GBM). The C321R-LAMB2 missense mutation leads to congenital nephrotic syndrome but only mild extrarenal symptoms;the mechanisms underlying the development of proteinuria with this mutation are unclear. We generated three transgenic mouse lines, in which rat C321R-LAMB2 replaced mouse LAMB2 in the GBM. During the first postnatal month, expression of C321R-LAMB2 attenuated the severe proteinuria exhibited by Lamb2−/− mice in a dose-dependent fashion; proteinuria eventually increased, however, leading to renal failure. The C321R mutation caused defective secretion of laminin-521 from podocytes to the GBM accompanied by podocyte endoplasmic reticulum (ER) stress, likely resulting from protein misfolding. Moreover, ER stress preceded the onset of significant proteinuria and was manifested by induction of the ER-initiated apoptotic signal C/EBP homologous protein (CHOP), ER distention, and podocyte injury. Treatment of cells expressing C321R-LAMB2 with the chemical chaperone taurodeoxycholic acid (TUDCA), which can facilitate protein folding and trafficking, greatly increased the secretion of the mutant LAMB2. Taken together, these results suggest that the mild variant of Pierson syndrome caused by the C321R-LAMB2 mutation may be a prototypical ER storage disease, which may benefit from treatment approaches that target the handling of misfolded proteins.


Pierson syndrome (Online Mendelian Inheritance in Man #609049), a rare autosomal recessive disease, is characterized by renal failure from congenital nephrotic syndrome/diffuse mesangial sclerosis, severe ocular abnormalities, and neurodevelopmental impairments.1–3 Pierson syndrome is caused by mutations in the laminin β2 gene (LAMB2).4

Laminin, type IV collagen, nidogen, and sulfated proteoglycans comprise the glomerular basement membrane (GBM),5 an unusually thick BM formed by fusion of distinct BMs assembled by podocytes and glomerular endothelial cells.6 Laminins are obligate heterotrimeric glycoproteins containing one α-, one β-, and one γ-chain.7 The major laminin heterotrimer in the mature GBM is laminin α5β2γ1 (LM-521).8 Laminin trimerization occurs in the endoplasmic reticulum (ER) and involves association of the three chains along their laminin coiled-coil domains to form the long arm.9 After trimers are secreted into the extracellular space, they polymerize with each other to form the supramolecular laminin network through interactions among the amino-termini of the short arms (laminin amino-terminal [LN] domains).10,11 Lamb2−/− mice

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exhibiting congenital nephrotic syndrome, abnormal neuromuscular junctions, and abnormalities in the retina re-capitulate Pierson syndrome.

LAMB2 null mutations cause the full syndromic phenotype of Pierson syndrome, whereas certain LAMB2 missense mutations, including R246Q, C321R, L1393F, and N1380K, cause congenital nephrotic syndrome with mild extrarenal features. These mild variants of Pierson syndrome have expanded the clinical spectrum of LAMB2-associated disorders. We previously showed that defective laminin secretion from podocytes to the GBM underlies the R246Q mutation-caused congenital nephrotic syndrome.

Missense mutations can lead to protein misfolding and disruption of protein trafficking. Alterations in protein trafficking occur mainly in the ER, the central site for folding, post-translational modifications, and transport of secretory, luminal, and membrane proteins. Protein folding in the ER is facilitated by ER-resident molecular chaperones and enzymes, such as immunoglobulin binding protein (BiP)/glucose-regulated protein (GRP)78, GRP94, oxygen-regulated protein 150 calnexin, calreticulin, and protein disulfide isomerase. An imbalance between the load of misfolded proteins and the folding capacity of the ER leads to ER stress. The ER responds to stress by activating the unfolded protein response (UPR), which is initiated by three ER transmembrane proteins—protein kinase regulated by RNA-like ER kinase, inositol-requiring protein-1, and activating transcription factor-6. BiP/GRP78 is also a key sensor linked to the UPR in stressed cells. Protein kinase regulated by RNA-like ER kinase phosphorylates eukaryotic initiation factor-2, leading to attenuation of protein translation and induction of activating transcription factor-4 and its target CCAAT/enhancer-binding protein homologous protein (CHOP), which is an ER-specific proapoptotic transcription factor. Misfolded protein and the resultant ER stress represent one important cause of ER storage diseases, such as cystic fibrosis, α1-antitrypsin deficiency, retinitis pigmentosa, and Alzheimer’s disease. Accumulating evidence also suggests that ER stress contributes to the development and progression of kidney disease.

Here, we investigate the mechanisms by which the C321R-LAMB2 missense mutation causes nephrotic syndrome using a knockout/transgenic approach to replace endogenous mouse laminin β2 with different levels of C321R mutant rat β2 together with an in vitro system. Our studies suggest that the C321R mutation impairs secretion of mutant laminin-521 from podocytes, resulting in low GBM laminin levels and altered GBM composition. Our results also indicate that podocyte ER stress and injury are linked to the development of proteinuria and that a chemical chaperone can partially ameliorate the secretion defect in vitro. These data advance our understanding of the molecular pathogenesis of LAMB2 mutation-induced congenital nephrotic syndrome and suggest a strategy for therapeutic intervention in this disease.

RESULTS

Generation and Characterization of Podocyte-Specific Mouse Nephrin Promoter-C321R-LAMB2 Transgenic Mice

To investigate the mechanisms whereby the C321R-LAMB2 mutation causes congenital nephrotic syndrome, we generated transgenic mice, in which a full-length rat β2 cDNA with an engineered C321R mutation was placed under the control of the podocyte-specific mouse nephrin promoter (NEPH). Three individual NEPH-C321R-LAMB2 transgenomes were bred onto the Lamb2−/− background. The fact that the expression of the wild-type (WT) rat β2 cDNA in podocytes through NEPH is sufficient to restore the integrity of the glomerular filtration barrier (GBF) in Lamb2−/− mice is proof of principle that Lamb2−/− mice expressing mutant rat β2 in podocytes serve as useful models for mild variants of Pierson syndrome.

In Lamb2−/−; NEPH-C321R-LAMB2 mice, the endogenous WT mouse β2 in the GBM was replaced by the mutant C321R-LAMB2. Transgene expression was assayed by both quantitative confocal immunofluorescence microscopy and in situ hybridization on the Lamb2−/− background. As shown in Figure 1A, dual immunostaining of kidney sections for rat laminin β2 and the GBM marker agrin identified three independent lines of transgenic mice with differing levels of rat β2 in the GBM (TgLo, TgMed, and TgHi represent the low, medium, and high transgene expressors, respectively). It was also noted that the highest C321R expressor (C321R-TgHi) exhibited a lower level of β2 in the GBM than the lowest R246Q expressor (R246Q-TgLo) studied previously (Figure 1A). Moreover, as we previously showed, rat β2 protein levels in the GBM were much higher in Lamb2−/− mice expressing WT rat β2 (Lamb2−/−; Tg-WT) than in R246Q-TgLo mice (Figure 1B). Surprisingly, transgene-derived mRNA was easily detected by in situ hybridization in C321R-TgMed and TgHi and the WT kidneys in a podocyte-specific pattern, although not in C321R-TgLo or R246Q-TgLo kidneys (Figure 1C). Taken together, analysis of transgene-derived mRNA and protein levels suggests that, even with transcription levels comparable with those levels of the WT β2 transgene in podocytes (and higher than the endogenous mouse Lamb2 mRNA), C321R-LAMB2 protein levels in the GBM were comparatively very low.

Proteinuria in Lamb2−/−; NEPH-C321R-LAMB2 Mice

Because proteinuria is the most sensitive clinical manifestation of GBF defects, we examined the in vivo function of C321R-LAMB2 mutant protein in the GBM by comparing proteinuria in the three lines of Lamb2−/−; NEPH-C321R-LAMB2 mice with their Lamb2−/− and WT littermates. Spot urinary protein to creatinine ratios were determined every 2 weeks for up to 12 weeks. Lamb2−/− mice developed severe proteinuria and died by 4–5 weeks. Between 3 and 4 weeks of age, the average protein to creatinine ratios (gram/gram) for Lamb2−/− (n=12), TgLo mutants (n=10), TgMed mutants (n=9), TgHi
that even the lowest expression of C321R-LAMB2 provides significant benefit.

Effects of C321R-LAMB2 on GBM Composition

During glomerulogenesis, there is a transition in laminin deposition, such that the $\alpha1\beta1\gamma1$ and $\alpha5\beta1\gamma1$ trimers are present in the nascent GBM but are replaced by LM-521 at maturation proceeds. In the absence of laminin B2, laminin B1 persists in the GBM, and laminins $\alpha1$, $\alpha2$, $\alpha3$, $\beta3$, and $\gamma2$ are ectopically deposited, perhaps as a compensatory response to abnormal GBM that might also be pathogenic. We, therefore, investigated how the replacement of WT LAMB2 with mutant C321R-LAMB2 impacted the laminin composition of the GBM. We examined laminin chain repertoire in TgHi mutants at 3 weeks, when proteinuria was minimal. By immunostaining, the normally mesangial laminins $\alpha1$ and $\alpha2$ (Figure 3, A and E and Supplemental Figure 1, A and B) were ectopically deposited in the GBMs of TgHi (Figure 3, B and F and Supplemental Figure 1, A and B) and nontransgenic Lamb2 mice (Figure 3, C and G and Supplemental Figure 1, A and B). In contrast, laminins $\alpha1$ and $\alpha2$ were found only in the mesangium of the Lamb2 mice (Figure 3, D and H and Supplemental Figure 1C). This finding suggests that high-level podocyte expression of WT B2 suppresses ectopic deposition of laminins $\alpha1$, $\alpha2$, and $\beta1$ in the GBM, whereas the reduced level of C321R-LAMB2 and absence of B2 in the GBM are associated with accumulation of these potentially pathogenic laminins, which may contribute to the progression of proteinuria.

Histologic and Ultrastructural Features at Early and Later Stages

Light and transmission electron microscopy (TEM) were used to examine renal histopathology in the three lines of Lamb2 mice and their WT and Lamb2 littermates. Lamb2 mice had developed severe proteinuria by 3 weeks, which was associated with mesangial matrix (MM) expansion and glomerulosclerosis (Figure 4A, d) compared with

Figure 1. There are three different levels of NEPH-C321R-LAMB2 transgene expression in the three independent lines. These are revealed by (A and B) quantitative confocal immunofluorescence and (C) in situ hybridization. Transgenes were on the Lamb2 background in all cases. (A) Quantitative confocal immunofluorescence showed different levels of transgene-derived rat C321R-LAMB2 and R246Q-LAMB2 proteins in the GBM. LAMB2 protein accumulation in the GBM was detected at ~3 weeks of age by colocalization of mutant LAMB2 (green) and agrin (red) in overlay images (merge). As shown in the histogram, significant differences in total fluorescence intensity were observed in glomeruli of the different groups (n=20–25 glomeruli for each line). **P<0.001, *P<0.05 by t test. Scale bar, 10 μm. (B) Panels from our previous paper show that Tg-WT mice exhibited much higher rat B2 in the GBM compared with R246Q-TgLo mutants. To avoid image saturation and the resulting quantification errors, different laser intensities and gains were used for the upper and lower panels; R246Q-TgHi staining was used as a common reference for comparison between R246Q-TgLo mutants and the Tg-WT mice. Original magnification, ×800. (C) Transgene-derived mRNA was easily detected by in situ hybridization in C321R-TgMed, C321R-TgHi, and Tg-WT mice but not C321R-TgLo or R246Q-TgLo mutants (at approximately 3 weeks). Scale bars, 100 μm.
controls (Figure 4A, a). Light microscopic examination of hematoxylin and eosin (H&E)-stained kidney sections also revealed diffuse MM expansion in TgLo mutants (Figure 4A, b). In contrast, there were no obvious renal histopathological changes in TgHi mutants at 3 weeks (Figure 4A, c). TgMed mutants showed limited glomerulosclerosis (data not shown). Periodic acid–Schiff (PAS) staining at 3 weeks showed that Lamb2<sup>−/−</sup> mice exhibited abundant renal tubular protein casts and complete loss of brush borders in their proximal tubules (Figure 4A, h). TgLo mutants showed fewer protein casts and only partial loss of brush borders (Figure 4A, f). In contrast to the above findings, in both TgMed and TgHi mutants, tubular protein casts were rarely seen, and tubular brush borders were preserved (Figure 4A, g) (data not shown).

Ultrastructural analysis revealed diffuse foot process (FP) effacement and mild GBM thickening in 3-week-old Lamb2<sup>−/−</sup> mice (Figure 4B, e). However, although some FP effacement was observed in most glomeruli of the transgenic mutants (Figure 4B, b–d), there were still focal areas with much less severe FP effacement in all three lines at 3 weeks (Figure 4B, f–h), which is consistent with the lower proteinuria compared with their nontransgenic Lamb2<sup>−/−</sup> littermates. There was no significant GBM thickening in the C321R mutants at 3 weeks (Figure 4B, b–d); however, a moth-eaten appearance of the GBM was sometimes observed in TgLo (Figure 4B, b) and TgMed (Figure 4B, c) mutants but not TgHi mutants (Figure 4B, d). These findings suggest that the increased transgene expression and the subsequent increased deposition of C321R-LAMB2 partly ameliorate the severe GFB defect in Lamb2<sup>−/−</sup> mice in a dose-dependent manner.

By 6 weeks of age, the three lines of transgenic mutants had developed more severe proteinuria and MM expansion (data not shown). TEM analysis showed diffuse FP effacement and mild GBM thickening in 3-week-old Lamb2<sup>−/−</sup> mice (Figure 4B, e). However, although some FP effacement was observed in most glomeruli of the transgenic mutants (Figure 4B, b–d), there were still focal areas with much less severe FP effacement in all three lines at 3 weeks (Figure 4B, f–h), which is consistent with the lower proteinuria compared with their nontransgenic Lamb2<sup>−/−</sup> littermates. There was no significant GBM thickening in the C321R mutants at 3 weeks (Figure 4B, b–d); however, a moth-eaten appearance of the GBM was sometimes observed in TgLo (Figure 4B, b) and TgMed (Figure 4B, c) mutants but not TgHi mutants (Figure 4B, d). These findings suggest that the increased transgene expression and the subsequent increased deposition of C321R-LAMB2 in the GBM partly ameliorate the severe GFB defect in Lamb2<sup>−/−</sup> mice in a dose-dependent manner.
The fact that a high level of mutant Defective Secretion of C321R-LAMB2 GFB ultrastructure and kidney histopathological features. C321R-LAMB2 in the GBM correlates with improvements in of nephrotic syndrome indicate that increased accumulation of

of nephrotic syndrome indicate that increased accumulation of C321R-LAMB2 in the GBM correlates with improvements in GFB ultrastructure and kidney histopathological features.

**Defective Secretion of C321R-LAMB2–Containing Trimmers from Podocytes to the GBM and Induction of Podocyte ER Stress**

The fact that a high level of mutant β2 transcription in podocytes was associated with a low level of β2 protein in the GBM (Figure 1) suggests that secretion of the mutant from podocytes may be defective. To test this hypothesis, we established stable human embryonic kidney 293T (HEK293T) cell lines expressing fusion proteins, in which the LN domain and laminin epidermal growth factor-like domain a (LEa) of rat WT or C321R β2 were fused to Gaussia luciferase (Gluc), the brightest known luciferase. Secretion of the LAMB2-LN/LEa-Gluc fusion proteins was directed by the β2 signal peptide, and trafficking of the fusion proteins was analyzed. Cells were cultured for 48 hours, and media were collected for both Western blot analysis with an anti-Gluc antibody and luciferase assay. Although the WT fusion protein was detected in the medium, the mutant fusion protein was not (Figure 6A). Similarly, the luciferase activity in the medium was significantly higher in the WT-transfected cells compared with the mutant-transfected cells (Figure 6B). In contrast, when cell lysates were subjected to Western blot analysis using the anti-Gluc antibody, both WT and mutant laminin β2 fusion proteins were easily detected, and the mutant seemed to be more abundant (Figure 6C). Quantitative RT-PCR analysis showed similar LAMB2 transcript levels in the WT and C321R clones (Figure 6D). Together, these in vitro studies directly show that the C321R mutation inhibits secretion of the C321R mutant protein, leading to its intracellular accumulation.

To delineate the subcellular localization of the WT and mutant proteins, stable 293T-Gluc cells were subjected to confocal immunofluorescence staining using antibodies against Gluc and protein disulfide isomerase A3 (PDI A3), an ER resident foldase catalyzing disulfide bond formation and an ER stress marker. Although C321R/ Gluc overlapped exclusively with PDI A3 (Figure 6E, b, d, and f), WT/Gluc showed only a partial overlap with PDI A3 (Figure 6E, a, c, and e). This result suggests that the WT/Gluc fusion protein was able to exit the ER and proceed through the secretory pathway, whereas C321R/Gluc was retained inside the ER. In addition, the upregulation of PDI A3 in cells expressing the mutant (Figure 6E, c and d) suggests that C321R/Gluc is misfolded and has triggered ER stress.

We next examined if BiP, an important ER molecular chaperone and a central sensor for the increased load of misfolded proteins, was induced. Western blot analysis showed increased expression of BiP in C321R cells versus WT cells; treatments with tunicamycin, which blocks N-linked glycosylation in the ER and causes ER stress, were included as positive controls (Figure 6F). Similarly, double immunofluorescence staining of BiP and the podocyte nucleus marker WT-1 showed a very low level of constitutive expression of BiP in the podocytes of WT and Lamb2−/−; Tg-WT mice (Figure 6G, a, e, and i and Figure 6 G, b, f, and j, respectively). In contrast, even at 3 weeks, when TgHHi mutants had developed only trace proteinuria, TgHi podocytes...
exhibited BiP upregulation in all glomeruli (Figure 6G, c, g, and k). Interestingly, significant BiP induction was also observed in the podocytes of 5%–10% of Lamb2−/− glomeruli at 3 weeks, when proteinuria was nephrotic range (Figure 6G, d, h, and l). In summary, our in vitro and in vivo data clearly suggest that the C321R mutation causes protein misfolding, which leads to podocyte ER stress. The fact that induction of BiP in podocytes of TgLo mutants with minimal proteinuria was observed in all glomeruli, which is in contrast to the limited increase of BiP in Lamb2−/− mice with heavy proteinuria (only seen in 5%–10% of glomeruli), supports the concept that the primary cause of podocyte ER stress is the misfolded C321R-LAMB2 protein rather than proteinuria.

Podocyte ER Distention, CHOP Activation, and Associated Podocyte Injury

ER stress triggers both survival and apoptotic signals. If the UPR is unable to resolve protein folding defects, ER dysfunction can lead to apoptosis.37 We investigated whether induction of ER stress in podocytes led to ER morphologic changes by TEM examination. At 3 weeks, we observed significant podocyte rough ER (rER) distention in TgHi mutants (Figure 7A, b) and a prominent vesiculated rER in the podocytes of Lamb2−/− littermates (Figure 7A, c). In contrast, Lamb2−/−; Tg-WT mice did not show significant podocyte rER dilation (Figure 7A, a).

Chronic and persistent ER stress may activate cell apoptosis by induction of CHOP, a proapoptotic transcription factor specifically related to ER stress. Indeed, at 3 weeks, a marked induction of CHOP was detected in the podocytes of some glomeruli of TgHi mutants (Figure 7B, b, e, and h) and Lamb2−/− mice (Figure 7B, c, f, and i) versus the Tg-WT mice (Figure 7B, a, d, and g). Moreover, compared with the Tg-WT podocytes (Figure 7C, a, d, and g), there was desmin expression in some podocytes (an indicator of podocyte injury) of the mildly proteinuric TgHi mutants (Figure 7C, b, e, and h) and much stronger desmin expression in the podocytes of heavily proteinuric Lamb2−/− mice (Figure 7C, c, f, and i). Collectively, these data suggest that podocyte ER dysfunction, manifested by ER distension and upregulation of CHOP, and the associated podocyte injury caused by the misfolded protein precede the occurrence of significant proteinuria in the TgHi mutants. These results also indicate that misfolding of the mutant protein, rather than simply overexpression of any LAMB2 protein (WT or mutant) in podocytes, induces all the above-mentioned events. Associated podocyte ER stress in Lamb2−/− mice with heavy proteinuria is perhaps from albumin overload inside Lamb2−/− podocytes; indeed, it has been shown that increased albumin endocytosis in cultured podocytes can induce ER stress and apoptosis.38

A Chemical Chaperone Promotes Secretion of Mutant C321R-LAMB2 Protein

Chemical chaperones, such as tauroudoxicholyl acid (TUDCA), can assist protein folding in the ER and facilitate the trafficking of mutant proteins. To determine whether chemical chaperones can improve secretion of the mutant C321R protein, 293T cells expressing either the WT/Gluc or the C321R/Gluc fusion protein were incubated with 1 mM TUDCA or vehicle, and the luciferase activity in the medium was determined. TUDCA selectively enhanced secretion of the mutant (Figure 8B) but not the WT fusion protein (Figure 8A) into the medium. These data suggest that TUDCA could be a good therapeutic candidate for the treatment of proteinuria in patients carrying the C321R and perhaps, other missense LAMB2 mutations.

DISCUSSION

In contrast to the typical Pierson syndrome phenotype, which has been associated primarily with null mutations in LAMB2,1–4 missense mutations, such as R246Q and C321R, cause mild variants of Pierson syndrome, in which congenital nephrotic syndrome is the predominant manifestation and extrarenal features are less pronounced.18 LAMB2 missense mutation is one of the most common disease-causing mutations in early onset nephrotic syndrome.19 Previously, we investigated the mechanisms whereby the R246Q mutation causes congenital nephrotic syndrome; here, we determined how the C321R mutation causes proteinuria.

Our results suggest that the mechanisms responsible for nephrotic syndrome in patients harboring the C321R-LAMB2 mutation involve both severely impaired laminin secretion and concomitant podocyte ER stress-associated injury. Our hypothesis is supported by both in vivo and in vitro data. In vivo, we generated three lines of transgenic mice, in which
C321R-LAMB2 mRNA was expressed in podocytes at differing levels, two of which were comparable with the expression level of the WT β2 transgene. Our analysis revealed that ample levels of C321R-LAMB2 mRNA in podocytes were associated with unexpectedly low levels of C321R-LAMB2 protein in the GBM, suggesting a secretion defect. In vitro, biochemical studies confirmed that secretion of C321R-LAMB2 was severely inhibited compared with WT-LAMB2. Moreover, the fact that podocyte desmin activation and FP effacement preceded overt proteinuria in the C321R mutant indicates that a primary podocyte injury is also crucial in the development of proteinuria. The mutant C321R-LAMB2 protein induced podocyte ER stress, which was indicated by the upregulation of the ER chaperones PDIA3 and BiP. Failure to relieve sustained and excessive ER stress led to podocyte injury, which was associated with activation of the ER stress-mediated apoptotic signal CHOP and ER dilation and dysfunction. In addition, the fact that the accumulation of ectopic laminins α1, α2, and β1 in the GBM occurred before marked proteinuria suggests another possible mechanism to accelerate the progression of proteinuria, which is most likely secondary to the reduced level of LM-521 in the GBM.

An important caveat here is that we did not find accumulation of C321R-LAMB2 protein inside podocytes because of deficient secretion. It is known that ER quality control mechanisms include ER-associated degradation (ERAD), and induction of the UPR may increase ERAD capacity. Thus, the bulk of the misfolded C321R-LAMB2 that is not secreted into the GBM may be degraded by ERAD rapidly and become depleted. The AF508 mutation in the cystic fibrosis transmembrane conductance regulator and misense mutations of Wolfram syndrome gene 1 in Wolfram syndrome are other prominent examples of functional proteins that are retained in the ER and degraded by ERAD. Our findings in mice suggest that the mild variant of Pierson syndrome caused by the C321R mutation may be a prototypical ER storage disease, in which the missense mutation leads to a recessive loss-of-function state. The autosomal recessive pattern of inheritance in the mild variants of Pierson syndrome is in agreement with this hypothesis. The aberrantly folded β2 protein may be retrotranslocated into the cytoplasm, tagged by ubiquitin ligase, and degraded by the 26S proteasome. However, it is also possible that the misfolded protein is degraded by the lysosome-autophagy pathway coupled with ER stress. Additional studies will be aimed at investigating the mechanisms of mutant protein degradation.

Protein overexpression can also induce ER stress. To ensure that the identified podocyte ER stress response and ER distortion are not caused by transgene overexpression, Lamb2−/− mice expressing WT rat β2 (Lamb2−/−; Tg-WT) were included as important controls in the relevant experiments. Our data clearly show that protein misfolding, but not overexpression, causes ER stress. At a molecular level, the mutation of Cys 321 to
lysosome inhibitor with a chemical chaperone may be required. Degraded, an alternative strategy of combining a proteasome or TUDCA and other chemical chaperones in our animal model.

**CONCISE METHODS**

**Generation of Mutant Rat Laminin β2 Transgenic Mice**

Site-directed mutagenesis of the rat Lamb2 cDNA, construction of the NEPH-C321R-LAMB2 transgene, and production and identification of transgenic mice were performed as described previously. All animal experiments conformed to the National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the Washington University Animal Studies Committee.

**Antibodies, Immunofluorescence, and In Situ Hybridization**

Commercially available antibodies were obtained as follows: mouse anti-rat laminin β2 carboxy-terminal coiled-coil domain mAbs D5 and D7 were from the Developmental Studies Hybridoma Bank (Iowa City, IA); rabbit anti-mouse agrin laminin globular domains was from Takako Sasaki (University of Erlangen-Nürnberg, Erlangen, Germany); rat anti-mouse laminin α2 mAb 4H8 was from Enzo Life-sciences (Farmingdale, NY); mouse IgG1 anti-human desmin clone D33, which crossreacts with mouse desmin, was from DAKO (Carpinteria, CA); rat anti-mouse nidogen clone ELM1 was from Millipore (Billerica, MA); rabbit anti-mouse BiP and CHOP antibodies were from Santa Cruz (Santa Cruz, CA); and mouse IgG1 anti-mouse WT-1 antibody was from Thermo Scientific (Kalamazoo, MI). Other primary antibodies were gifts from generous colleagues: rat anti-mouse laminin α1 mAb 8B350 was from Dale Abrahamson (University of Kansas Medical Center, Kansas City, KS); rabbit anti-mouse laminin β1 was from Takako Sasaki (University of Erlangen-Nürnberg, Erlangen, Germany); and rabbit anti-mouse podocin was from Corinne Antignac (Necker Hospital, Paris, France). Alexa 488- and Alexa 594-conjugated secondary antibodies were purchased from Invitrogen (Carlsbad, CA).

**Chemical Chaperones**

Chemical chaperones can stabilize protein conformation and rescue trafficking-defective but otherwise functional misfolded proteins. They have been used in the treatment of cystic fibrosis, α1-antitrypsin deficiency, nephrogenic diabetes insipidus, Gaucher disease, Fabry’s disease and Fabry’s disease in vitro or in animal models. We have shown here that increased accumulation of the C321R mutant protein in the GBM alleviates proteinuria, and TUDCA partially rescues the secretion defect of the mutant protein in vitro. Attempts to test TUDCA and other chemical chaperones in our animal model are in progress. Considering that the mutant protein may be degraded, an alternative strategy of combining a proteasome or lysosome inhibitor with a chemical chaperone may be required.
Confocal Microscopy and Quantitative Image Analysis
Confocal microscopy was used to quantify the levels of laminin β2 accumulation in the GBM in the three C321R and the one R246Q mutant β2 transgenic lines. In the individual comparative studies, 8-μm kidney cryosections of the different genotypes were placed on the same slide and immunolabeled with the same mixture of primary antibodies (D5 and D7) and Alexa 488-conjugated secondary antibody. The slides were then examined under a Nikon TE-2000 scanning laser confocal microscope (Melville, NY), and Z-series images were captured at 0.3-μm intervals. For each genotype, 20–25 glomeruli were scanned, and the images were captured on the same day using the same laser intensity, confocal aperture, and gain. Raw confocal images taken from the mid-regions of the glomeruli were imported into Image J software. Total pixel density within a fixed circle, which was slightly smaller than the smallest glomerular field, was used to measure glomerular immunofluorescence intensity. The mean intensity for each genotype was compared statistically using a t test.

Light and Electron Microscopy
For light microscopy, kidneys were fixed in 10% buffered formalin, dehydrated through graded ethanol, embedded in paraffin, sectioned at 4 μm, and stained with H&E, PAS, and Gomori’s Trichrome by standard methods. For TEM, tissues were fixed, embedded in plastic, sectioned, and stained as described previously.

Establishment of Stable 293T Cells Expressing Gluc-Tagged Laminin β2 Fragments
We used a mammalian expression vector, pCMV-Gluc (New England Biolabs), containing a recombinant Gluc-Tag to which WT and site-directed mutant rat laminin β2 fragments containing the LN and LEa domains were fused. The expression constructs were transfected into HEK293T cells in 24-well plates using Lipofectamine 2000 (Invitrogen). Individual clones were isolated, and stable transfectants were selected in medium containing 25 μg/ml Zeocin (Invitrogen). Established clones were maintained in DMEM (Gibco, Grand Island, NY) supplemented with 10% heat-inactivated FBS (Gibco) and 25 μg/ml Zeocin.

Western Blot Analysis
Stable 293T-Gluc cells were cultured for 48 hours. Cells were lysed by RIPA buffer (Sigma) containing protease inhibitor cocktail (Roche, Indianapolis, IN), and the protein concentration of each cell lysate was determined by Bio-Rad protein assay (Hercules, CA) using BSA as a standard. Denatured proteins were separated on polyacrylamide gels and then transferred to polyvinylidene difluoride membranes. Blots were blocked with 5% nonfat milk for 1 hour and then incubated overnight with primary antibodies. The membranes were washed with Tris-buffered saline/Tween buffer and incubated with the appropriate horseradish peroxidase-conjugated secondary antibody. The proteins were then visualized in an x-ray developer using ECL-plus detection reagents (GE, Pittsburgh, PA). To ensure equal protein loading, the same blot was stripped with stripping buffer (25 mM glycine + 1% SDS, pH=2.0) and then incubated with a horseradish peroxidase-conjugated mouse anti-human β-actin antibody (Sigma). To examine secretion of LAMB2/Gluc fusion proteins, 40 μl conditioned media were subjected to Western blot analysis with a fluorescence-conjugated (IRDye 680LT) goat anti-rabbit secondary antibody (Li-Cor Biosciences, Lincoln, NE) and visualized in an infrared imaging system (Li-Cor Biosciences). Rabbit anti-Gluc antibody was from New England Biolabs.

Luciferase Assay
The Gluc activity in the media of 293T-Gluc cells stably expressing either WT or mutant LAMB2/Gluc was assayed by a BioLux Gaussia Luciferase Assay Kit (New England Biolabs) according to the manufacturer’s instructions.
instructions and quantified with a Femtometer FB12 Luminometer (Zylux, Oak Ridge, TN). The actual values of raw light units were normalized with respect to total cell protein for each group.

**mRNA Quantification by Real-Time PCR**

Total RNA from the individual WT and mutant 293T-Gluc clones was extracted using the RNeasy kit (QIAGEN, Valencia, CA) with subsequent DNase treatment; 1 μg RNA was then reverse-transcribed using an RT-PCR Kit (Superscript III; Invitrogen). mRNAs of the WT or mutant LAMB2/Gluc were evaluated by quantitative real-time PCR; 1 μl cDNA was added to SYBR Green PCR Master Mix (Qiagen) and subjected to PCR amplification (one cycle at 95°C for 20 seconds and 40 cycles at 95°C for 1 second and 60°C for 20 seconds) in an Applied Biosystems 7900HT Fast Real-Time PCR System (Life Technologies, Grand Island, NY) using human β-actin as an internal control. Quantitative PCR was conducted in triplicate for each sample. The sequences of primers were LAMB2/Gluc forward: CATGGAG-CAGTTTCATCGCAC, reverse: GTCAGAACACTGACGTTGG; human β-actin forward: GGCACCCAGCACAATGAAG, reverse: GGACCAGGCAATTAAG.

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**DISCLOSURES**

None.

**REFERENCES**


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