Laminin β₂ variants associated with isolated nephropathy that impact matrix regulation

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Graphical abstract

GBM: Glomerular basement membrane

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Laminin β₂ variants associated with isolated nephropathy that impact matrix regulation

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Introduction

The glomerular filtration barrier allows the efficient flow of water and small molecules while preventing plasma proteins from leaking into the urine. The barrier consists of podocytes, endothelial cells, and the glomerular basement membrane (GBM). Nephrotic syndrome, characterized by proteinuria, edema, hypoalbuminemia, and hyperlipidemia, is associated with impaired glomerular filtration. Nephrotic syndrome can be caused by dysfunctions of podocytes, endothelial cells, or GBM, indicating that all 3 components are essential for maintaining the glomerular filtration barrier. Extensive genetic studies of nephrotic syndrome have identified mutations in genes important for establishing and maintaining the barrier (1).

The GBM is a specialized extracellular matrix separating 2 cellular layers, endothelial cells and podocytes. The GBM plays critical roles in filtration and in maintenance of glomerular morphology (2). Basement membrane (BM) is usually composed of 4 major extracellular matrix proteins: laminin, collagen IV, nidogen, and sulfated proteoglycan. The GBM contains an atypical assortment of BM protein isoforms, including collagen α₃α₄α₅(IV) and laminin-521 (LM521; α₂β₂γ₁; ref. 3). Mutations in collagen α₃α₄α₅(IV) cause Alport syndrome, a kidney disease with variable ear and eye defects (4). LM521 is composed of α₂, β₂, and γ₁ chains. Mutations in the laminin β₂ gene (LAMB2) cause Pierson syndrome, which is characterized by congenital nephrotic

Mutations in LAMB2, encoding laminin β₂, cause Pierson syndrome and occasionally milder nephropathy without extrarenal abnormalities. The most deleterious missense mutations that have been identified affect primarily the N-terminus of laminin β₂. On the other hand, those associated with isolated nephropathy are distributed across the entire molecule, and variants in the β₂ LEa-LF-LEb domains are exclusively found in cases with isolated nephropathy. Here we report the clinical features of mild isolated nephropathy associated with 3 LAMB2 variants in the LEa-LF-LEb domains (p.R469Q, p.G699R, and p.R1078C) and their biochemical characterization. Although Pierson syndrome missense mutations often inhibit laminin β₂ secretion, the 3 recombinant variants were secreted as efficiently as WT. However, the β₂ variants lost pH dependency for heparin binding, resulting in aberrant binding under physiologic conditions. This suggests that the binding of laminin β₂ to negatively charged molecules is involved in glomerular basement membrane (GBM) permselectivity. Moreover, the excessive binding of the β₂ variants to other laminins appears to lead to their increased deposition in the GBM. Laminin β₂ also serves as a potentially novel cell-adhesive ligand for integrin α₄β₁. Our findings define biochemical functions of laminin β₂ variants influencing glomerular filtration that may underlie the pathogenesis of isolated nephropathy caused by LAMB2 abnormalities.
syndrome with severe ocular and neuromuscular defects (5). Zenker et al. originally identified homozygous or compound heterozygous mutations in the LAMB2 gene, which contains 32 exons, in patients with Pierson syndrome (6). More than 49 mutations have since been reported to be associated with Pierson syndrome (5). Most disease-associated alleles are truncating mutations localized across the entire gene, eliminating laminin β2 function completely (7). On the other hand, the patients with LAMB2 missense mutations occasionally exhibit higher mean age at onset of renal disease or oligosymptomatic disease variants of Pierson syndrome (5). For example, p.R246Q and p.R246W lead to congenital or infantile nephrotic syndrome with milder or no extrarenal symptoms (5, 6). A patient with Pierson syndrome carrying p.S80R presented with late-onset nephrotic syndrome that was diagnosed when the patient was 6.5 years old (5). Another missense mutation, p.C321R, is associated with congenital nephrotic syndrome, despite less severe extrarenal defects (5). p.C321R affects the formation of disulfide bonds in an EGF-like module and disrupts the structure of the LEa domain, leading to defective secretion of laminin β2 and podocyte ER stress possibly due to misfolding of the aberrant laminin β2 (8). Previously, Matejas et al. demonstrated that missense mutations found in Pierson syndrome are apparently clustered in the laminin N-terminal (LN) domain, which is required for the polymerization of laminins (5). Therefore, mutations in the LN domain have been implicated in the perturbation of BM formation.

Laminin is a family of heterotrimeric glycoproteins that mediate cell adhesion via various receptors. Five α, 3 β, and 3 γ chains have been characterized, and 19 isoforms have been identified in various tissues and cell culture media (9). Of the isoforms, 9 contain laminin β2, and LM521 is the most studied of these. LM521 trimers are recognized by several receptors, including integrin αβ1, αβ6, and αβ7; α-dystroglycan; and Lutheran/basal cell adhesion molecule. The C-terminal 20–amino acid sequence of the laminin β2 chain modulates the binding affinities of laminins to X2-type integrins, such as αβ1 and αX2β1, (10). However, because the laminin globular domains of laminin α chains have been implicated in various cellular interactions, the X2-type integrins are not specific for laminin β2. The β2 chain of laminin-421 and -521 are also identified as ligands that bind to N- and P/Q-type voltage-gated calcium channels (VGCCs, ref. 11). Although VGCCs specifically bind to laminin β2, at neuromuscular junction presynaptic terminals, it is unlikely that these mediate cell adhesion to the β2 chain. So far, the cell-adhesive activity of laminin β2 has not been fully examined.

Next-generation sequencing approaches have also identified LAMB2 missense mutations in congenital or childhood-onset nephrotic syndrome without apparent extrarenal abnormalities (12–19). Here we demonstrate that most of these missense variants found in cases with isolated nephropathy affect a region in the LEx-LF-LExb domains in the laminin β2 short arm. Notably, variants in this region have not been found in patients with classic Pierson syndrome. We examined the effects of these variants on secretion of laminin β2, in vitro. To explore the biochemical properties that might influence glomerular filtration, recombinant laminin β2 chains carrying the variants were produced in mammalian cells. We found that laminin β2 serves as a potentially novel cell-adhesive ligand for integrin αβ1. These results unraveled roles of the laminin β2 short arm in matrix regulation that may underlie the pathogenesis of isolated nephropathy as a mild form of Pierson syndrome.

Results
We previously reported a Japanese patient with isolated nephropathy carrying LAMB2 p.R469Q and p.G699R variants (12). He was born to a healthy parent and appeared normal at birth. He exhibited systemic edema by 21 months and was diagnosed with steroid-resistant nephrotic syndrome. A kidney biopsy at 23 months of age revealed focal segmental glomerulosclerosis (FSGS) and mesangial matrix expansion (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.145908DS1). As seen by electron microscopy, the GBM exhibited a stratified structure (Supplemental Figure 1B). The patient’s renal function declined and progressed to end-stage kidney disease (ESKD) at 26 months of age. After renal transplantation at 7 years of age, he is presently healthy without extrarenal symptoms. Our whole-exome analyses of the patient and his parents showed that c.1406G>A (p.R469Q) and c.2095G>C (p.G699R) in LAMB2 are derived from his father and mother, respectively (Supplemental Figure 2, A and B, and ref. 12). The former, c.1406G>A (p.R469Q), has not been reported in Genome Aggregation Database (gnomAD) (20) or TogoVar (21). Although c.1406G is adjacent to a splice acceptor site, the predictions of bioinformatics tools such as NNSplice and ASSP (22, 23) were comparable to that of WT (Supplemental Figure 2C), suggesting that c.1406G>A does not influence splicing. The allele frequency of c.2095G>C (p.G699R) is 0.01173 in East Asians (gnomAD) and 0.007 in 2 Japanese genome databases (ToMMo 4.7KJPN and GEM-J WGA; available from TogoVar, ref. 21, and Supplemental Table 1).
A Hispanic patient was found to have massive proteinuria by 20 months and diagnosed with steroid-resistant nephrotic syndrome. Sanger sequencing identified a heterozygous c.3232C>T (p.R1078C) variant in LAMB2; its allele frequency is 0.000276 (gnomAD). Additional pathogenic variants were not identified in LAMB2 or in genes associated with early-onset nephrotic syndrome in this patient, but we suspect there is an additional pathogenic variant in LAMB2 that could not be identified. In silico algorithms (SIFT, PolyPhen-2, Align-GVGD) suggested that p.R1078C is likely to be disruptive (24–26). In the ClinVar database (27) there are multiple submissions of this variant in nephrotic syndrome or Pierson syndrome but with scarce clinical or genetic information.

Including these cases, 14 cases carrying LAMB2 variants that presented with nephropathy without ocular or neurologic abnormalities have been reported in the literature (Table 1). These patients presented with abnormal urinalysis to varying degrees, ranging from mild hematuria to nephrotic range proteinuria. Among the patients, 4 of their cases progressed to ESKD in the infantile period (cases 2, 3) or in early childhood (cases 4, 5). Other cases maintained normal kidney function in childhood and even at school age. This feature is quite different from the clinical picture of typical Pierson syndrome, in which patients often progress to ESRD in the early infantile period. With respect to genetic involvement, 1 patient (case 14) carries splice site variants in LAMB2, and the other patients carry at least 1 missense variant. These missense variants are distributed in exons 2–29 (Figure 1A). This is in clear contrast to missense mutations found in typical patients with Pierson syndrome, which cluster in exons 2–9 and 25–28. Missense variants in a region spanning from exon 10 to 24, which encode the LEa-LF-LEb domains in the laminin $\beta_2$ short arm, were found in 9 patients (64.3%) with isolated nephropathy (cases 4–13), despite the mutations in that region not being reported in previously reported classical Pierson syndrome patients with ocular or neurological manifestations (Supplemental Table 2). Furthermore, none of these 9 patients progressed to ESRD in the infantile period. Therefore, we hypothesized that variants in this region may be associated with the mild phenotype caused by LAMB2 abnormalities.

**Table 1. The missense variants in isolated nephropathy**

<table>
<thead>
<tr>
<th>Case</th>
<th>Variants</th>
<th>homo/compound (c)</th>
<th>Onset</th>
<th>ESRD</th>
<th>Maintained kidney function (y.o.)$^a$</th>
<th>Biopsy findings</th>
<th>Ethnicity</th>
<th>Publications</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>p.D53N</td>
<td>p.R464H</td>
<td>c.hetero</td>
<td>8 mo</td>
<td>1 y</td>
<td>FSGS</td>
<td>Chinese</td>
<td>17</td>
</tr>
<tr>
<td>2</td>
<td>p.R246Q</td>
<td>p.R246Q</td>
<td>homo</td>
<td>1 mo</td>
<td>&lt;1 y (expired at 5 y)</td>
<td>FSGS</td>
<td>Turkish</td>
<td>16</td>
</tr>
<tr>
<td>3</td>
<td>p.R246Q</td>
<td>p.R246Q</td>
<td>homo</td>
<td>&lt;1 mo</td>
<td>&lt;1 y</td>
<td>NA</td>
<td>Turkish</td>
<td>16</td>
</tr>
<tr>
<td>6</td>
<td>p.F560I</td>
<td>p.P1024L</td>
<td>c.hetero</td>
<td>10 y</td>
<td>Expired by ESRD</td>
<td>IgM nephropathy</td>
<td>Pakistani</td>
<td>18</td>
</tr>
<tr>
<td>7</td>
<td>p.R644H</td>
<td>p.G1623D</td>
<td>c.hetero</td>
<td>0.8 y</td>
<td>4 y</td>
<td>FSGS</td>
<td>European</td>
<td>19</td>
</tr>
<tr>
<td>10</td>
<td>p.I992V</td>
<td>p.R1148H</td>
<td>c.hetero</td>
<td>12 y</td>
<td></td>
<td>FSGS</td>
<td>Pakistani</td>
<td>18</td>
</tr>
<tr>
<td>11</td>
<td>p.Q1048H</td>
<td>p.Q1048H</td>
<td>homo</td>
<td>13.5 y</td>
<td></td>
<td>FSGS</td>
<td>Pakistani</td>
<td>18</td>
</tr>
<tr>
<td>12</td>
<td>p.R1078C</td>
<td>Not identified</td>
<td>c.hetero</td>
<td>20 mo</td>
<td></td>
<td>DMS</td>
<td>Hispanic</td>
<td>This study</td>
</tr>
<tr>
<td>13</td>
<td>p.A1556V</td>
<td>p.A1556V</td>
<td>homo</td>
<td>1 y</td>
<td></td>
<td>FSGS</td>
<td>Pakistani</td>
<td>18</td>
</tr>
<tr>
<td>14</td>
<td>p.Q1692Vfs</td>
<td>c.3797+5G</td>
<td>c.hetero</td>
<td>NA</td>
<td>5 y</td>
<td>TBM, BWC</td>
<td>Japanese</td>
<td>13</td>
</tr>
</tbody>
</table>

$^a$Renal function is maintained until at least the stated age. FSGS, focal segmental glomerulosclerosis; ESRD, end-stage renal disease; DMS, diffuse mesangial sclerosis; TBM, thin BM; BWC, basket weave change.

Secretion of laminin $\beta_2$ short arm carrying p.R469Q, p.G699R, and p.R1078C in vitro. To examine the impact of missense variants in the LEa-LF-LEb domains of the laminin $\beta_2$ short arm, we searched for associated functional alterations. We chose 3 variants found in our patients, p.R469Q, p.G699R, and p.R1078C, affecting the LEa, LF, and LEb domains, respectively (Figure 1A). Although Arg469 and Gly699 are highly conserved among species, Arg1078 is replaced with His in the rhesus monkey and mouse (Figure 1B). Our previous study showed that some laminin $\beta_2$ mutations result in impaired secretion (28). Therefore, we investigated whether...
the 3 β2 mutations influenced the secretion of laminin β2 by expressing N-terminal fragments of rat β2 mutants in HEK293 cells using a system in which the β2 short arm was fused to a human immunoglobulin Fc domain (Figure 2A). As shown in our previous study, although the p.C321R mutant was synthesized in the transfectants, its secretion was impaired (Figure 2, B and C). In contrast, the p.R469Q, p.G699R, and p.R1078C mutant fusion proteins were secreted into the medium similarly to the WT fusion protein. These results indicate that these 3 variants did not inhibit LM521 secretion.

Heparin binding properties of mutant laminin β2 short arms. Lamb2-knockout mice exhibit mislocalized anionic sites in the GBM, indicating that laminin β2 is associated with negative charge either directly or indirectly (29). The negative charge of the GBM is mainly imparted by heparan sulfate chains of the proteoglycan agrin (30, 31). Based on the known locations of both the laminin binding site and the glycosaminoglycan (GAG) attachment sites on agrin, together with super-resolution imaging data (32), the laminin β2 short arm could be capable of interacting with agrin’s negatively charged GAG side chains. Therefore, we investigated whether the laminin β2 variants bound to heparan sulfate using recombinant proteins. We purified recombinant rat laminin β2 short arms carrying the variants fused with human IgG 1 Fc from conditioned media (Figure 3A). Human IgG 1 Fc and mouse laminin β1 short arm fused with human IgG 1 Fc were prepared as control proteins. SDS-PAGE analysis of the purified proteins revealed a single protein band corresponding to the expected size. Heparin, a highly sulfated form of heparan sulfate, was used for solid phase binding assays. Heparin acts as a cation exchanger because of its polyanionic features and can bind to cationic proteins in an ion strength–dependent manner. Whereas both WT and mutant laminin β2 short arms bound to heparan sulfate under physiologic conditions (Figure 3B). These results indicated that the variants increased the binding of laminin β2 to heparin under physiologic conditions. Our results also show that laminin β1 did not bind heparin, indicating that heparin binding is a characteristic feature of the β2 chain.

To narrow the heparin binding region on the laminin β2 short arm, we produced truncated recombinant fragments (Figure 4, A and B). Solid phase binding assays were performed under low ionic conditions. Heparin bound to laminin β2 LN-LEa domains, indicating the heparin binding region is localized in the N-terminus.
These results suggest that the variants structurally influence the heparin binding affinity of the N-terminus under physiologic conditions rather than providing additional heparin binding sites. We next examined the heparin binding of trimeric laminins and found that under low ionic conditions, LM521 containing the $\beta_2$ chain bound heparin significantly stronger than did laminin-511 containing the $\beta_1$ chain (Figure 4D). There was no difference in heparin binding under physiologic conditions. These results indicate that laminin $\beta_2$ influences heparin-binding properties of the whole laminin trimer.

Although heparin bound to laminin $\beta_2$ at low ionic strength, it is unlikely that there are low-salt conditions in vivo. Another factor that can modulate the charge of laminin $\beta_2$ for heparin binding is the concentration of hydrogen ion (pH). To explore this, we first calculated the theoretical isoelectric point (pI) of each domain based on amino acid sequence (Figure 5A and Supplemental Figure 3). Of the laminin $\beta_2$ domains, the pI of the LN domain was more than 7.4, which is the physiologic pH in blood. On the other hand, each pI of laminin $\beta_1$ domains exhibited acidity. These results suggest that the laminin $\beta_2$ LN domain has more positive charge under acidic conditions. Therefore, solid phase binding assays were performed under different pH conditions (Figure 5B). Heparin binding to the laminin $\beta_2$ short arm was enhanced at pH = 6.1 versus pH = 7.1. These results indicate that laminin $\beta_2$ modulates GBM charge in a pH-dependent manner.

The binding of laminin $\beta_2$ variants to laminin-111. Laminins polymerize to form a network in BMs. The polymerization of laminins is mediated by the 3 short arms of the cross-shaped heterotrimer (33). The short arms are capped with LN domains that mediate the interactions among laminin trimers. To investigate the interaction of mutant laminin $\beta_2$ short arms with other laminins, solid phase binding assays were performed using laminin-111 (Figure 6A). Laminin $\beta_2$ mutants readily bound to laminin-111, but the WT $\beta_2$ short arm did not bind at this concentration (64 nM). The binding of $\beta_1$ and $\beta_2$ LN domains to laminin-111 was also
observed at higher concentration (128 nM), indicating that they bound less robustly than did the mutants (Supplemental Figure 4). Divalent cations such as Ca\textsuperscript{2+} are required for self-polymerization of laminins (33).

To investigate the manner of laminin-111 binding, we also performed solid phase binding assays in the presence of EDTA (Figure 6B). Chelation of divalent cations significantly enhanced laminin-111 binding rather than inhibiting it. That the laminin \( \beta_2 \) short arm bound to laminin-111 in a divalent cation-independent manner suggests that the variants unmask a novel laminin binding site.

Adhesion of HEK293 cells to the laminin \( \beta_2 \) short arm in vitro. The heparin binding activity of laminin \( \beta_2 \) led us to hypothesize that it binds to cell surface receptors containing heparan sulfate such as syndecans and glypicans. We therefore explored the cell adhesion activity of the laminin \( \beta_2 \) short arm. HEK293 cells suspended...
in serum-free DMEM were plated on wells coated with the recombinant proteins. The cells weakly adhered to the laminin \( \beta_2 \) LN-L Ea domains but not to the full \( \beta_2 \) short arm (Figure 7A). Based on the heparin binding at low ionic strength, the cell adhesion assay was next performed in diluted DMEM. The low ionic strength markedly promoted cell adhesion not only to the laminin \( \beta_2 \) LN-L Ea domains but also to the \( \beta_2 \) short arm. The cells readily adhered to the laminin \( \beta_2 \) LN-L Ea domains, indicating that the cell attachment site is localized in the N-terminus. The laminin \( \beta_2 \) mutants were more cell adhesive than the WT \( \beta_2 \) short arm (Figure 7B). Thus, similar to heparin binding, the variants appeared to structurally influence cell-adhesive activity of the laminin \( \beta_2 \) N-terminus. Furthermore, to clarify the mechanism of cell adhesion, we identified the receptors’ binding to the laminin \( \beta_2 \) N-terminus. Although cell surface receptors carrying heparan sulphate were initially expected, the cell adhesion assays revealed divalent cation dependency, indicating the involvement of integrins. Cell adhesion assays were performed in the presence of function-blocking antibodies to integrins (Figure 7C). Anti–integrin \( \beta_1 \) and \( \alpha_4 \) monoclonal antibodies significantly inhibited the adhesion of HEK293 cells to laminin \( \beta_2 \) LN-L Ea domains, indicating that the cell adhesion is mediated by integrin \( \alpha_4 \beta_1 \).

**Figure 4. Characterization of heparin binding to the laminin \( \beta_2 \) short arm.** (A) Schematic diagrams of truncated laminin \( \beta_1 \) and \( \beta_2 \) short arms fused with an Fc-tag. (B) SDS-PAGE analysis of B1SA_Fc (\( \beta_1 \) short arm, lane 1), B1LN-L Ea_Fc (lane 2), B2SA_Fc (\( \beta_2 \) short arm, lane 3), and B2LN-L Ea_Fc (lane 4). (C) Heparin binding of \( \beta \) chain LN-L Ea domains in 10 mM Tris-HCl (pH 7.5). Ninety-six-well microtiter plates were coated with 10 \( \mu \)g/mL of recombinant proteins. After blocking, 10 \( \mu \)g/mL of biotinylated heparin was applied to the wells and incubated for 1 hour at room temperature. The bound heparin was detected as described in Methods. Box-and-whisker plots show median, 25th and 75th percentiles, and minimum and maximum values (n = 6 from 3 independent experiments). Data were analyzed by 1-way ANOVA with Tukey’s multiple-comparison test, *P < 0.001. (D) The binding of heparin to whole laminin containing \( \beta \) chain in 10 mM Tris-HCl (pH 7.5), without (left panel) or with 150 mM (right panel) NaCl. Ninety-six-well microtiter plates were coated with laminin-111 (LM111), laminin-511 (LM511), and laminin-521 (LM521) at 10 \( \mu \)g/mL. Data were analyzed by 1-way ANOVA with Tukey’s multiple-comparison test, *P < 0.001 and are shown as box-and-whisker plots (n = 6 from 3 independent experiments).
Discussion

Recently, variants in LAMB2 have been found in patients with nephropathy but without apparent extrarenal manifestations. These cases also present with less severe renal phenotypes. Proteinuria is often found in childhood, and some cases maintain renal function in adolescence, which is a clear contrast to the clinical course of typical Pierson syndrome. Electron microscopic analysis of kidneys of these patients demonstrated a thin or stratified GBM forming a basket weave pattern (Supplemental Figure 1 and ref. 13). Notably, laminin β2 in the GBM from a proteinuric patient carrying compound heterozygous truncating (p.Y76TfsTer) and missense (p.G699R) variants was detected (13), indicating that p.G699R did not cause a secretion defect. We noted that the distribution of missense variants causing Pierson syndrome and those of isolated nephropathy were different: variants in a region spanning from exon 10 to exon 24, which correspond to the LEa-LF-LEb domains in the laminin β2 short arm, were exclusively found in patients with isolated nephropathy. Although it is known that LN domains mediate the self-polymerization of laminins, the functions of the LEa-LF-LEb domains have been unclear. Here we found that the p.R469Q, p.G699R, and p.R1078C variants affecting the LEa-LF-LEb domains found in isolated nephropathy cases did not abrogate laminin secretion but altered heparin and laminin-111 binding. These data suggest that the LEa-LF-LEb domains regulate matrix interactions, and conformational changes in this segment lead to altered GBM structure. Because these variants do not cause extrarenal defects, the altered LEa-LF-LEb domains must maintain those laminin β2 functions required in ocular and neurological tissues. The present results suggest a novel mechanism underlying isolated nephropathy caused by LAMB2 defects, which is clinically and genetically distinct from Pierson syndrome.

The glomerular filtration barrier is permeable to water and small molecules and selectively restricts macromolecules such as albumin (34). Because proteinuria precedes podocyte abnormalities in Lamb2-knockout mice, the GBM plays a critical role in selective filtration (29). The GBM had been believed to exhibit not only size selectivity but also charge selectivity to produce urine (31). However, because the reduction of negative charge using knockout mice does not affect the filtration barrier (35, 36), the hypothesis of charge selectivity has been challenged (37). Therefore, the mechanism whereby laminin β2 influences the size selectivity of the GBM is an important concept. In this study, WT laminin β2 bound heparin under not only low ionic but also acidic conditions, indicating that heparin binding is pH dependent. The pH dependency of laminin β2 binding to heparin is likely due to the basic pl of the β2 LN domain. The basic or neutral pl of the β2 LN domain is conserved in the mouse and human (Supplemental Figure 4) whereas that of the β1 LN domain is acidic. This may be why laminin β1 imperfectly compensates for the functions of β2, despite their structural similarity (38). Our solid phase binding assays also showed that the laminin β2 variants, but not WT β2, bound to heparin under neutral conditions. The recombinant proteins carrying p.R469Q, p.G699R, and p.R1078C bound similarly to heparin. The binding of mutant amino acid residues to heparin was minimal under physiologic conditions.
The solid phase binding assay using truncated recombinant proteins showed that heparin binding sites are localized in the N-terminus of β2. The variants may lead to a conformational change in the β2 short arm, causing heparin binding sites to become exposed. The heparin binding property of laminin β2 allows us to hypothesize a model in which the short arm of β2 variants binds to heparan sulfate on agrin in the GBM (Figure 8A). This binding seems to disturb laminin polymerization under physiologic conditions, resulting in a large-pore GBM that can be permeable to macromolecules such as albumin. Similar to the β2 variants, the WT β2 short arm binds to heparan sulfate under acidic conditions. To test this model, future investigations on the effect of pH variation in mice would be required.

Once laminins are secreted into the extracellular space, they polymerize to form a meshwork via interactions among LN domains of the α, β, and γ short arms (33). The pairings of α-α, α-β, α-γ, and β-γ are possible in the presence of Ca²⁺ (39). Although the LN domain of β2 binds to the N-termini of α1, α5, β3, and γ1, 3, our solid phase binding assays showed that WT β2 short arm weakly bound to laminin-111. On the other hand, the mutant β2 short arm readily bound to laminin-111 in a calcium-independent manner. These in vitro results suggest a model in which the variants expose a cryptic laminin binding site in the β2 short arm under physiologic conditions. Structural or biochemical analyses using samples from patients or knock-in mice carrying these variants are required to confirm the model.

Our results indicate that laminin β2 serves as a potentially novel cell-adhesive ligand for integrin αβ1. Integrin αβ1 plays a major role in the regulation of immune cell recruitment to inflamed endothelia and other
sites of inflammation. Previous studies have reported that integrin $\alpha_4\beta_1$ is a receptor for vascular cell adhesion molecule-1 (VCAM-1) and fibronectin (40). In fibronectin, integrin $\alpha_4\beta_1$ recognizes a motif containing the sequence EILDVPST within the alternatively spliced connecting segment-1 (41), with the LDV sequence being most critical (42). Homology search of the rat laminin $\beta_2$ amino acid sequence revealed that 2 LDVs

Figure 7. Analysis of cell adhesion to laminin $\beta_2$. (A) Adhesion of HEK293 cells to the WT $\beta_2$ short arm in diluted DMEM. Ninety-six-well microtiter plates were coated with 10 $\mu$g/mL of Fc (control tag protein), B1SA_Fc ($\beta_1$ short arm), BILN-LNEa_Fc ($\beta_1$ LN-LNEa domains), B2SA_Fc ($\beta_2$ short arm), and B2LN-LNEa_Fc ($\beta_2$ LN-LNEa domains). After blocking, HEK293 cells were suspended in DMEM diluted with H_2O and incubated in the wells at 37°C for 30 minutes. Attached cells were stained with Diff-Quik and counted. Box-and-whisker plots show median, 25th and 75th percentiles, and minimum and maximum values ($n = 6$ from 3 independent experiments). Data were analyzed by 1-way ANOVA with Tukey’s multiple-comparison test, $^*P < 0.001$. (B) Adhesion of HEK293 cells to the mutant $\beta_2$ short arms in the 1:3 DMEM dilution. Ninety-six-well microtiter plates were coated with 20 $\mu$g/mL of the mutant recombinant proteins. The number of attached cells is shown as box-and-whisker plots ($n = 6$ from 3 independent experiments) with analysis by 1-way ANOVA with Tukey’s multiple-comparison test, $^*P < 0.001$. (C) Inhibitory effects of integrin-blocking antibodies on adhesion of HEK293 cells to $\beta_2$ LN-LNEa domains. HEK293 cells preincubated with function-blocking antibodies against the indicated integrin subunits were added to B2LN-LNEa_Fc–coated wells. After incubation for 30 minutes, the attached cells were stained and counted. The attached cells were quantified and analyzed as described above. Data are shown as box-and-whisker plots ($n = 6$ from 3 independent experiments) with analysis by 1-way ANOVA with Tukey’s multiple-comparison test, $^*P < 0.001$. 

![Figure 7](https://example.com/figure7.png)
are localized in the LN and LCC domains, respectively. The N-terminal LDV sequence is likely exposed to integrin $\alpha_4\beta_1$ because the position is close to the signal peptide cleavage site. However, this LDV is conserved in rodents but not humans. We also searched for the LDV sequence in human laminin $\beta_2$ and found that it is localized in the LF domain. This LDV is conserved in mammals except for rodents. Pulido et al. reported the

Figure 8. Schematic of laminin $\beta_2$ activity in the GBM. (A) The disorder of permselectivity on GBM with $\beta_2$ variants. The polymerization of laminin-521 containing WT $\beta_2$ chain forms the GBM that restricts permeability of macromolecules such as albumin (ALB) (left). The N-terminus of $\beta_2$ variants binds to heparan sulfates on agrin, when the LEa-LF-LEb variants are present (right). This binding disturbs laminin polymerization, resulting in larger GBM pores with increased permeability to albumin. (B) Dysfunction of GBM with laminin $\beta_2$ variants. The heparin-binding activity of laminin $\beta_2$ variants leads to a large-pore GBM under neutral conditions, resulting in proteinuria. Increased laminin binding also results in the accumulation of $\beta_2$ variants in the GBM. The increased cell adhesive activity would affect glomerular cells.
crystal structure of the laminin β1 LF domain (43). Because the position of the LDV is located at the molecular surface, integrin αβ1 would be able to bind to the sequence. Our recent study showed that the β1 short arms are polymerized toward the center of the GBM, suggesting that it is difficult for the binding site to be accessible to integrins (32). The excessive deposition of β1 variants in the GBM may expose adhesive sites for integrin αβ1-positive cells (Figure 8B). In this regard, expression of integrin αβ1 was detected in the endothelial cells and mesangial cells of glomeruli (Supplemental Figure 5). Therefore, the exposure of the integrin binding site might strengthen cell adhesion, leading to glomerular dysfunction. Integrin αβ1 is a major cell surface receptor on immune cells such as lymphocytes. Moreover, Jurkat and Raji cells derived from human lymphocytes adhere to laminin β1 via integrin αβ1 (data not shown). The nephritogenic immune responses are driven by CD4+ T cells (44). CD4+ T cells spontaneously adhere within uninflamed glomerular capillaries (45), and the accumulation of immune cells such as CD4+ T cells, neutrophils, and macrophages is often observed in inflamed glomeruli (44). In addition to VCAM-1 on glomerular endothelial cells, laminin β1 may also serve as an adhesive ligand for inflammatory cells in nephritis.

Increased laminin deposition in the GBM has been reported in a C5b6a3−/− mouse model of Alport syndrome (46, 47). LAMB1 and LAMA2 are normally found in the mesangial matrix, but ectopic deposition is detected in Alport GBM (46). Furthermore, increased laminin α1 mRNA was detected in Alport glomeruli (47). The increased deposition of laminin was also reported in patients with Alport syndrome (48). These increases may be due to increased production or altered turnover of extracellular matrix molecules triggered by instability caused by the collagen abnormality. We hypothesize that the mechanism of GBM laminin anomalies observed in Alport syndrome is different from that predicted here due to variations in the LEa-LF-LEb domains of LAMB2.

The variants in the LEa-LF-LEb domains found in isolated nephropathy are not common, but the allele frequencies of some variants are around 1% in specific ethnic populations. For example, the frequency of c.2095G>C (p.G699R; rs28364667) is 0.012 in the East Asian population and 0.007 in Japanese databases. Variant c.3071C>T (p.R1078C) is observed at 0.0098 in the South Asian population (gnomAD, ref. 20). Many Mendelian disorders, such as Pierson syndrome, are caused by highly penetrant rare variants. Recent empirical evidence has shown that variants at low frequency or rare variants are associated with complex diseases. The recessive nature of these variants and the residual function of the mutant proteins make it difficult to determine their pathogenicity. Mouse models with altered LEa-LF-LEb domains to mimic the defect or exome- or whole genome–based variant analysis in large cohorts, including several target populations, will be required to further uncover the contribution of disease-related variants in this region to the pathogenesis of proteinuric kidney disease.

Methods

Antibodies and reagents. Monoclonal antibodies against human integrin α1 (P1E6), α2 (P1B5), α3 (P4G9), α4 (P1D6), α5 (P5G10), and α6 (P3G8) (Developmental Studies Hybridoma Bank) were purified on Protein G Sepharose (Cytiva) from conditioned media of hybridoma cells purchased from Developmental Studies Hybridoma Bank. Anti-integrin α1 (FB12), β1 (65E6), and β2 (25E11) antibodies were purchased from Merck. Anti-β1–actin antibody conjugated with peroxidase was from Medical & Biological Laboratories Co. Ltd. Alexa Fluor 647–conjugated rat monoclonal antibody against mouse integrin α1 (R1-2) and isotype control rat IgG2b (RTK4530) were purchased from BioLegend. Anti-podocin polyclonal antibody (catalog ab50339) was from Abcam. Human recombinant laminin-111 (LMI111), laminin-511 (LM511), and laminin-521 (LM521) were purchased from BioLamina. Mouse EHS laminin was from BD Biosciences and Takako Sasaki (Oita University School of Medicine, Oita, Japan). Biotinylated heparin with an average mass of 12.5 kDa was from Celsus Laboratories, Inc.

Construction of expression vectors. cDNA clones encoding full-length mouse laminin β1 (provided by Albert E. Chung, University of Pittsburgh, Pittsburgh, Pennsylvania) and rat laminin β1 (generated in-house) were used to construct expression vectors. The fragments encoding laminin β1 LN-LEa domains (B1N) and short arm (B1SA) were amplified using primer sets described in Supplemental Table 3 and subcloned into a human IgG1 Fc expression vector. The EcoRI-XbaI fragment encoding the laminin β1 LN-LEa domains (B2N) was subcloned into a human IgG1 Fc expression vector. To construct the β1 short arm expression vector, the fragment encoding laminin β1 LF-LEb domains was amplified using primer sets described in Supplemental Table 3 and subcloned into the XbaI site of the B2N_Fc expression vector. The R469Q, G699R, and R1078C human laminin β1 variants were separately engineered into analogous sites of the rat cDNA by site directed mutant PCR using primer sets described in Supplemental Table 3. Laminin β1 carrying the C321R variant was prepared in our previous study (28).
Preparation of serum-free conditioned media and cell lysates. HEK293 cells were obtained from the American Type Culture Collection. The cells were maintained in DMEM containing 10% fetal calf serum. To express WT and mutant proteins, the expression vectors were transiently transfected into the cells in 24-well plates using Lipofectamine 2000 (Thermo Fisher Scientific). The next day, the growth media were replaced with 500 μL of serum-free medium. After 3 days’ culture, the conditioned media were harvested and clarified by centrifugation twice at 100 g for 5 minutes and 9730 g for 10 minutes. Then 20 μL of conditioned media were applied for immunoblotting. The transfectants were also lysed with 500 μL of 10 mM Tris-HCl (pH 7.5), 150 mM NaCl, 10 mM EDTA, 1% NP-40, and protease inhibitor cocktail (Merck). The cell lysates were clarified by centrifugation once at 9730 g for 10 minutes. Then 20 μL of cell lysate was applied for immunoblotting.

**Immunoblotting.** Cell lysates and conditioned media were separated on 5%–20% gradient gels (ATTO) under reducing conditions and transferred to a PVDF membrane (Merck). Conditioned media and lysates were immunoblotted with anti-human IgG, Fc antibody (Jackson ImmunoResearch catalog 109-066-098). The bound antibodies were visualized with the ECL Western blotting detection system (Cytiva). The images of bands were captured with WSE-6100 Lumino Graph I (ATTO).

**Expression and purification of recombinant proteins.** HEK293 cells transfected with the expression vectors were selected using 100 μg/mL Zeocin (Thermo Fisher Scientific). All further cell culture was carried out in the presence of the antibiotic. The selected cells were grown to confluence in culture dishes with DMEM containing 10% FBS. The confluent cells were incubated in serum-free DMEM for 4 days. The conditioned media were harvested and clarified by centrifugation twice at 100 g for 5 minutes and 9730 g for 10 minutes. The recombinant proteins were purified from the conditioned culture media by Protein A Sepharose (Cytiva). The eluted fractions were pooled and dialyzed against PBS(-). The purified proteins were separated by SDS-PAGE using 5%–20% gels under reducing conditions. The separated proteins were stained with Coomassie Brilliant Blue (nacalai tesque).

**Solid phase binding assays.** Heparin binding assays were carried out with recombinant proteins coated onto high-protein-binding capacity 96-well microtiter plates (IWAKI). The plates were blocked with PBS(-) containing 1% BSA for 1 hour at room temperature. A total of 10 μg/mL of biotinylated heparin in 10 mM Tris-HCl (pH 7.5) containing 0, 150, and 500 mM NaCl or 10 mM BisTris-HCl (pH 6.1 and 7.1) containing 150 mM NaCl was added into the wells. After incubation for 1 hour at room temperature, the bound heparin was detected by addition of streptavidin-conjugated horseradish peroxidase, followed by addition of 1 mg/mL o-phenylenediamine and 0.012% H2O2. The absorbance was measured at 450 nm with Multiskan GO microplate spectrophotometer (Thermo Fisher Scientific). For laminin-111 binding assay, 96-well microtiter plates were coated with 10 μg/mL of EHS laminin-111. After blocking with PBS(-) containing 1% BSA, 10 μg/mL of Fc-tagged recombinant proteins in the binding buffer (10 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM CaCl2, 1 mM MgCl2 containing 0.05% Tween 20) were incubated at room temperature for 1 hour. The bound recombinant proteins were detected with a biotinylated anti-human IgG Fc antibody (Jackson ImmunoResearch) and quantified as described above.

**Cell attachment assays.** For adhesion assays, 96-well microtiter plates (IWAKI) were coated with recombinant proteins and blocked with PBS(-) containing 1% BSA. The dissociated HEK293 cells were suspended in serum-free DMEM and plated at 2 × 10^4 cells/50 μL/well. After incubation at 37°C for 0.5 or 1 hour, the attached cells were stained with Diff-Quik (SYSMEX). The stained cells were counted using image cytometer BZ-X800 (Keyence). To identify the receptors for laminin β1, 10 μg/mL of monoclonal antibodies against different integrins were preincubated individually with cells in a volume of 50 μL of serum-free medium (2 × 10^4 cells/well) at room temperature for 10 minutes. The preincubated cells were transferred to plates coated with laminin β1, LN-LEa domains fused with Fc-tag (B2N_Fc) and then incubated further at 37°C for 1 hour. After incubation, the attached cells were quantified as described above.

**Staining of renal biopsy specimens.** Kidney biopsy specimens were fixed in 10% neutral buffered formalin and embedded in paraffin. The tissues were sectioned and stained with periodic acid–methenamine silver stain–Masson’s trichrome and periodic acid–Schiff.

**Electron microscopy.** After fixation in glutaraldehyde fixative (1.6% v/v paraformaldehyde, 2.5% v/v glutaraldehyde, 0.2 M sodium cacodylate buffer pH 7.4, 5 mM CaCl2, 10 mM MgCl2), tissues were embedded in epon resin and sectioned. Imaging was performed using a HITACHI H-7100.

**Immunohistochemistry.** C57BL/6J mice (72 weeks old, male) were purchased from Charles River Laboratories Japan. Mice were sacrificed under deep anesthesia with Thiopental. Mouse kidneys were frozen in Tissue-Tek optimum cutting temperature compound (Sakura Finetek) and were cut at 7 μm in a cryostat. After
blocking with 10% normal goat serum, the sections were incubated with rat monoclonal antibody against mouse integrin αv and rabbit polyclonal antibody against podocin. The integrin αv antibody conjugated with Alexa Fluor 647 was used. Rabbit IgG was detected with secondary antibody conjugated with Alexa Fluor 594 (Thermo Fisher Scientific catalog A11037). After several washes, sections were mounted in 90% glycerol containing 0.1× PBS and 1 mg/mL p-phenylenediamine. Images were captured using a BZ-X800 (Keyence).

Statistics. Comparisons between groups were analyzed using 1-way ANOVA with Tukey’s multiple-comparison test. Statistical analyses were performed in R Studio (Version 1.3.1093). Significance, set at $P < 0.05$, is depicted as follows in figures: *$P < 0.001$, **$P < 0.05$.

Study approval. Informed consent was obtained from all the members of a family according to the guidelines of the Ethics Committee of Yamagata University (protocol number 2018-190) and the University of New Mexico School of Medicine. Clinical data of patients, including laboratory data, imaging data, and kidney biopsy, were obtained for diagnostic purposes and with informed consent. Animal studies were approved by the Animal Research Committee of Tokyo University of Pharmacy and Life Sciences (F20-33).

Author contributions

TH, YH, and YK were responsible for the overall design of the study. KT, SU, HM, MM, YY, KH, MN, M Hisano, YA, and KM performed the experiments and analyzed the data. JHM prepared cDNA clone encoding full-length rat laminin β2. TH, YH, M Hattori, HL, JHM, and YK reviewed the results and wrote the manuscript. All authors offered constructive comments.

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