Glomerular podocytes are highly specialized cells with a complex cytoarchitecture. Their most prominent features are interdigitated foot processes (FP) with filtration slits. These are bridged by the slit diaphragm, which plays a major role in establishing the selective permeability of the glomerular filtration barrier. Injury to podocytes leads to proteinuria, a hallmark of most glomerular diseases [30].

The ICGN mouse is a model of glomerular sclerosis (GS) that shows gross morphological changes in the podocyte FP, accompanied by proteinuria. The ICGN mouse is also a model of chronic kidney disease (CKD) that presents the common symptoms and pathological changes associated with a variety of kidney diseases, such as hyperlipidemia, anemia and systemic edema, and eventual end-stage renal failure [28, 29]. Previously, we identified a deletion mutation in the tensin2 (Tns2) gene (designated Tns2nph) via CRISPR/Cas9-mediated genome editing. Tns2nph/ΔC compound heterozygous KO mice displayed podocyte abnormalities and massive proteinuria similar to ICGN mice, indicating that these two mutations are allelic. Further, this result suggests that the SH2-PTB domain of Tns2 is required for podocyte integrity. Tns2 knockout in a mouse podocyte cell line significantly enhanced actin stress fiber formation and cell migration. Thus, this study provides evidence that alteration of actin remodeling resulting from Tns2 deficiency causes morphological changes in podocytes and subsequent proteinuria.

KEY WORDS: CRISPR/Cas9, glomerular sclerosis, podocyte, SH2-PTB domain, tensin2

PTPase domain at the N-terminus followed by a PTEN region and a Src homology 2 (SH2)/phosphotyrosine binding (PTB) domain at the C-terminus [19]. The SH2-PTB domain has been suggested to bind to the intracellular domain of integrin-family proteins, resulting in intracellular transmission of the integrin signal, leading to adjustment of cytoskeleton dynamics [3, 5]. However, little is known about the function of the SH2-PTB domain in vivo.

RNA-guided, nuclease-mediated genome editing, based on the CRISPR/Cas system, offers an efficient and convenient technique for genome editing [14]. In brief, Cas9, a nuclease guided by single-guide RNA (sgRNA), binds to a targeted genomic sequence next to the protospacer adjacent motif (PAM) and generates a double-strand break (DSB). The DSB is then repaired by nonhomologous end-joining (NHEJ), leading to insertion/deletion mutations [20].

To test whether Tns2 mutation might cause the GS phenotype and to clarify the biological role of the SH2-PTB domain, we created knockout mice carrying a Tns2 protein deletion in the SH2-PTB domain via CRISPR/Cas9-mediated mutagenesis.

MATERIALS AND METHODS

Ethical statement: All research was conducted according to the Regulations for the Care and Use of Laboratory Animals of Kitasato University and the National Center for Global Health and Medicine. The animal experimentation protocol was approved by the President of Kitasato University based on the judgment of the Institutional Animal Care and Use Committee of Kitasato University (Approval ID: No. 15–053). A humane end point was applied when mice with severe anemia became moribund.

Mouse: CRISPR/Cas9-mediated genome editing in mice was performed as described previously [27]. Briefly, sgRNA expression vector for the target sequence (AGAGACTCATTCCATCCA) coupled with a T7 promoter was synthesized in vitro using the MEGASHortscript kit (Thermo Fisher Scientific, Waltham, MA, U.S.A.). hCas9 mRNA from pX330 (https://www.addgene.org/42230/) was synthesized using the mMESSAGE mMACHINE T7 kit (Thermo Fisher Scientific) and was polyadenylated with the Poly (A) Tailing kit (Thermo Fisher Scientific). The purified hCas9 mRNAs (100 ng/μl) and sgRNAs (50 ng/μl) were co-injected into the cytoplasm of fertilized eggs derived from BDF1 females (Japan SLC, Hamamatsu, Japan). After the injected oocytes were cultured overnight in vitro, two-cell embryos were transferred into pseudo-pregnant female mice. Genomic DNA was isolated from the offspring from samples taken from the tail, using standard methods. The region around exon 22 of the Tns2 locus was amplified by PCR, using two sets of primers: Tns2 forward, GCCTCAGACTAGATTGGTTCACGAAT and Tns2 reverse, GAAATGGCGACACGTGTTCAGA. The amplification products were sequenced and compared to the wild-type. The resulting founder animals were crossed to FVB/N mice (CLEA Japan, Tokyo, Japan) and then backcrossed to the same for three generations. The Tns2ΔC KO heterozygote mice were bred to Tns2ΔC KO heterozygote and Tns2nph (FVB-Tns2nph) mice to produce compound heterozygotes (Tns2nph/Tns2ΔC) and homozygous knockouts (Tns2ΔC/Tns2ΔC), respectively. The nph genotype derived from the original ICGN mice was determined as described previously [32].

Measurement of urinary albumin excretion: Urine samples were collected by gentle manual compression of the abdomen. A 10-μl aliquot [containing 2% SDS, 5% β-mercaptoethanol, 10% glycerol, 60 mM Tris-HCl (pH 6.8), bromophenol blue and 5 μl of urine] was heated for 5 min at 95°C and subjected to 10% SDS-polyacrylamide gel electrophoresis. As a positive control, bovine serum albumin (BSA) was loaded simultaneously. The gel was fixed and stained with Coomassie brilliant blue (CBB; Wako, Osaka, Japan) according to the manufacturer’s instructions. CBB-stained urinary albumin was quantified using the image analysis program ImageJ (http://rsb.info.nih.gov/ij/).

Histology: Ten-week-old mice were sacrificed by an overdose of isoflurane, and whole kidneys were dissected out. The kidneys were embedded in OCT compound and frozen with liquid nitrogen. Four-micrometer-thick cryostat sections were cut, transferred to MAS-coated slides, air dried and stored at −80°C until use. For immunohistochemical analysis, the slides were washed in PBS, fixed with acetone at 4°C and incubated for 8 hr at 4°C with the primary antibody diluted in 1% BSA in PBS (rabbit anti-Tns2 central portion [26] 1:1,000; and rabbit anti-Tns2 C-terminal (SAB4200268, Sigma-Aldrich, MO, U.S.A.) 1:1,000). The slides were then washed in PBS and incubated with biotin-conjugated donkey anti-rabbit IgG as a secondary antibody (Histofine; Nichirei Biosciences, Tokyo, Japan) for 30 min at room temperature and treated with horseshadish peroxidase-conjugated streptavidin complex (Histofine; Nichirei Biosciences) for 3,3-diaminobenzidine staining. Lastly, slides were dehydrated and mounted. Periodic acid-Schiff (PAS) staining and ultrastructural analysis were performed using transmission electron microscopy (TEM) as described in our previous report [32].

Knockdown of Tns2 in the podocyte cell line: The conditionally immortalized mouse podocyte cell line MPC5 used in our study was a kind gift from Professor Peter Mundel [25]. Briefly, the podocytes were cultured at 33°C in RPMI1640 medium (Thermo Fisher Scientific) containing 10% fetal bovine serum (Thermo Fisher Scientific) and 100 U/ml recombinant mouse interferon-γ (Sigma-Aldrich). MPC5 cells were inoculated at a density of 5 × 10⁵ cells/well in 6-well plates. After 24 hr, cells at 70–80% confluence were transfected with Stealth Select RNAi (CCACUCUAAAGCAGCAGUACUCUA, UAGAGUACUCGGUCUUUGAGUGG, MSS209763, Thermo Fisher Scientific) (nucleotides 321–345 of Tns2 cDNA; Accession no. NM_153533.2) or MISSION siRNA universal negative control (Sigma-Aldrich) in serum-free medium using lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer’s instructions. To differentiate MPC5 cells after siRNA transfection, the cells were plated on type I collagen dishes and cultured with 1% FBS in a 5% CO₂ atmosphere at 37°C for 3 days. To confirm silenc-
Cell Adhesion assay: Adhesion assays with crystal violet staining were performed according to the method described in a previous report [10]. Tns2 knockout (KD) MPC5 cells and control cells were trypsinized and seeded on 6-well plates coated with collagen type IV (Sigma-Aldrich, 10 mg/ml), laminin (Sigma-Aldrich, 10 mg/ml), vitronectin (Sigma-Aldrich, 10 mg/ml) or fibronectin (Sigma-Aldrich, 10 mg/ml) at a density of 1.5 × 10^4 cells per well. After incubation for 1 hr at 37°C, non-adherent cells were removed by gentle washing with PBS, followed by fixation in 100% ethanol. Cells were stained in 0.1% crystal violet for 15 min at 25°C, washed in water and then counted under a microscope. All experiments were performed in triplicate wells for each condition, and data are expressed as means ± standard deviation.

Statistical analyses were performed using Student’s t-test and Dunn’s multiple comparison test. P-values <0.05 were considered significant.

Transwell migration assays: Tns2 KD MPC5 cells and control cells were plated in 8.0-µm pore size transwell inserts at a density of 1.5 × 10^4 cells per well according to the method described in a previous report [13]. After 24 hr, the cells on the upper side of the insert were removed by scraping, and the cells that had migrated through were fixed on the lower side of the membrane with 100% ethanol, stained with crystal violet and quantified by counting the number of cells in 20 separate fields. All experiments were performed in triplicate wells for each condition, and the data are expressed as means ± standard deviation. Statistical significance was determined using Dunn’s multiple comparison test, with P-values <0.05 considered significant.

Phalloidin staining of Tns2 KD cells: Tns2 KD MPC5 cells and control cells were seeded onto 24-well collagen I-coated culture glass. To observe actin reorganization, wound gaps were made in the MPC5 monolayers by scratching using a 200-µl tip. Images were captured after 8 hr. Cells were rinsed twice with PBS, fixed with fresh methanol-free 3.7% PFA for 10 min, permeabilized with 0.1% Triton-X for 5 min and incubated with Alexa Fluor 594 Phalloidin (Thermo Fisher Scientific). Actin stress fibers were identified and quantified using the EVOS FL cell imaging system (Thermo Fisher Scientific). All experiments were performed in quintuplicate wells for each condition. Data are expressed as means ± standard deviation. Statistical significance was determined using Dunn’s multiple comparison test, and P-values <0.05 were considered significant.
values < 0.05 were considered significant.

RESULTS

Generation of Tns2 SH2-PTB domain-KO mice: To generate mice carrying mutations that disrupt the Tns2 C-terminal SH2-PTB domain, we designed a sgRNA targeting exon 22 of Tns2. Exon 22 encodes the latter half of SH2, which binds specifically to many intracellular signal-transducing proteins (Fig. 1A). To verify genetic modification at the target locus, a region of genomic DNA including Tns2 exon 22 was amplified by PCR and subjected to sequencing analysis. After co-microinjection of sgRNA/Cas9 mRNA into fertilized eggs, five types of progeny were born (Fig. 1A, founder mouse nos. M1–M5). M3, M4 and M5 mice had 5-bp deletions or a 1-bp insertion close to the PAM sequence, which might lead to frame-shift mutations and subsequent protein deletion in both the SH2 and PTB domains (designated as Tns2ΔC). All nucleotide changes in these mice were transmitted to the next generation by mating with the FVB strain.

Loss of Tns2 results in proteinuria and FP effacement of podocytes: Because the original ICGN mouse is a spontaneous mutant derived from a closed colony of ICR mice, there is no control strain. The FVB/N strain has been identified as susceptible to the development of GS and CKD [33]. Thus, we previously created FVB.ICGN-Tns2nph (FVB-Tns2 nph) congenic mice. Next, genetic complementation testing was undertaken using FVB-Tns2ΔC and FVB-Tns2nph. Tns2ΔC heterozygous mice (M3 strain) were bred to Tns2nph homozygotes to produce compound heterozygotes (Tns2nph/ΔC) in the FVB strain background. To determine whether expression of the Tns2 protein remained at a normal level and only the SH2-PTB domain had been deleted, we conducted immunohistochemical analyses on kidney sections of compound heterozygotes (Tns2nph/ΔC) compared with FVB-Tns2nph homozygotes and age-matched wild-type (WT) controls (Tns2ΔC/ΔC). We used two antibodies recognizing the central portion of Tns2 and the C-terminal PTB domain, respectively (Fig. 2A). The use of the two antibodies revealed that Tns2 protein expression was lost in FVB-Tns2nph mice, which is in agreement with an earlier report [35]. In contrast, Tns2 protein expression and localization in glomeruli was normal in the compound heterozygotes. However, Tns2 could be detected by Ab1 but not by Ab2, indicating that the SH2-PTB domain had been deleted. Scale bars=10 µm.

In the compound heterozygotes, proteinuria was detectable at 4 weeks of age using SDS-PAGE, and subsequent CBB staining demonstrated remarkable proteinuria (Fig. 3B). In contrast, no urinary albumin excretion was detectable in FVB-Tns2ΔC/+ mice (Fig. 3A and 3B). Histological analy-
sis with PAS staining revealed that almost all glomeruli in the compound heterozygote mice showed entire expansion of the mesangial matrix at 12 weeks of age (Fig. 4, upper panel). Ultrastructural analysis revealed fused podocyte FPs, loss of slit diaphragms and GBM thickening in all glomeruli in the compound heterozygotes at 12 weeks of age (Fig. 4, lower panel). In addition, homozygous Tns2ΔC/Tns2ΔC mice displayed massive proteinuria similar to that seen in ICGN mice (Fig. 3B). In contrast, compound heterozygotes derived from M1 and M2 founders did not show any abnormalities (data not shown). Thus, the Tns2nph mutation was confirmed to be responsible for GS in ICGN mice. Further, this result suggests that the SH2-PTB domain of Tns2 is required for podocyte integrity. In addition, Tns2ΔC homozygote mice did not show any of these other phenotypes, indicating a selective role for Tns2 in kidney function.

Loss of podocyte Tns2 results in increases in the formation of actin stress fibers and cell migration: Cell-cell contact and adherence of podocytes to the extracellular matrix of the GBM are crucial for podocyte function. It is well known that cell-to-cell and cell-to-extracellular matrix (ECM) adhesions affect morphological changes involved in cell migration. Integrins, a large family of cell adhesion proteins, mediate the adhesion of cells to the ECM and provide traction for cell motility. Many proteins present on the cytoplasmic side of focal adhesions, including those in the tensin family, are considered to link transmembrane receptors to the actin cytoskeleton [19]. The actin cytoskeleton is an essential structural and functional element that controls cell shape, cell motility and adhesion. When the extracellular environment is altered, these structures are disassembled and remodeled to meet the new requirements [12]. To examine whether Tns2 is involved in adhesion and migration in podocytes, MPC5 cells, a conditionally immortalized podocyte cell line, were transfected with siRNAs for the coding sequence of Tns2, and two assays were used to determine the effects of Tns2 KD on adhesion and migration. Adhesion assays were performed to investigate the effect of Tns2 KD on podocyte anchorage to the ECM. Reduced levels of Tns2 protein ideally should be verified by quantitative western blotting. However, because good Tns2 antibodies are not available, we used RT-qPCR to detect knockdown of Tns2 mRNA. In KD cells, Tns2 transcripts were significantly decreased compared to in control cells (Fig. 5A). We then examined the influence on adhesion to several types of ECM (collagen type IV, fibronectin, laminin and vitronectin). Figure 5B demonstrates that attachment of Tns2-KD podocytes did not differ from that of the negative control for any ECM. Thus, Tns2 suppression does not significantly affect the adherence of podocytes under these conditions. The Tns paralog, Tns1, has also been demonstrated to interact with and regulate the actin cytoskeleton or integrin [5].

To observe actin reorganization, we created wound gaps in cell monolayers by scratching a straight line with a 200-µl tip. Tns2 KD produced a tendency toward increased migration in the scratch assay (data not shown). We monitored the actin architecture in Tns2 KD podocytes migrating towards the center of the gap by staining with conjugated phalloidin-Alexa Fluor 594 (Fig. 5C). The degree of actin stress fiber formation was classified into three categories (high to low: a, b and c), depending on the thickness and length of the actin stress fibers in the cytoplasm. In Tns2-KD podocytes,
cells in category a (rich in stress fibers, with thick cables) increased, but the number of those in category b was similar to control cells. In contrast, cells in category c (with few stress fibers) decreased, suggesting that Tns2 suppression significantly enhanced actin stress fiber formation (Fig. 5C and 5D). Next, to quantify cell migration accurately, we performed a transwell migration assay, which is widely used for studying the motility of different types of cells. Cells that migrated across the transwell membrane were quantified by fixing and counting. In general, migrating cells have thicker stress fibers than non-motile cells. As expected, Tns2 KD podocytes, cells in the stress fiber-rich category, with thick cables, (type a) increased, whereas cells lacking stress fibers, without thick cables, (type c) decreased, suggesting that Tns2 suppression significantly enhanced actin stress fiber formation (Fig. 5C and 5D). Tns2-KD podocytes were compared with control cells in 5 separate fields (n=5).

**DISCUSSION**

In this study, to test whether Tns2nop might cause the mutant phenotype, we crossed np/hp mice with mice that carried a SH2-PTB domain deletion. Compound heterozygotes that inherited both Tns2nop and Tns2ΔC displayed marked GS and proteinuria, indicating that these two mutations are allelic, and confirming that Tns2 deficiency is responsible for the GS phenotype. Further, KD of Tns2 expression in the podocyte cell line increased both actin stress fiber formation and migration speed. There are several basic types of proteinuria, including glomerular, tubular, overflow and exercise-induced proteinuria. Glomerular proteinuria accounts for approximately 90% of all proteinuria [24]. Proteinuria arises due to injury of the glomerular filtration barrier. The currently available evidence suggests that podocytes act as the main component of this barrier, as mutations in a number of podocyte-specific genes have been identified to be responsible for GS [23, 36]. Thus, podocyte dysfunction is a common determining factor for progression toward many types of kidney diseases. A study of several inherited diseases in humans and of KO mouse models revealed that mutations in several podocyte genes (ACTN4, CD2AP, SYNPO, MYH9, ARHDGA and ARHGP24) lead to GS [1, 7, 9, 15–17]. These proteins are involved in actin organization in podocytes and the mutations results in FP effacement. Thus, it has become ever clearer that the precise organization and regulation of the actin cytoskeleton in podocytes is essential for the maintenance of normal structure and function and the actin cytoskeleton serves as the common final pathway organizing FP effacement, independent of the cause of podocyte damage [12, 22].

The function of Tns2 can be predicted based on its interactions with proteins of known function. Tensins are a
deficient mutants [31]. Although no cyte abnormalities and proteinuria similar to that of Tns2- [11]. Both α3 integrin- and β1 integrin-KO mice show podo-
N-terminal region of Tns1 interacts with actin at multiple
Tns3 and Tns4 [19]. All isoforms contain a PTB that allows
adhesions. Four members have been identified: Tns1, Tns2,
family of proteins that are localized to integrin-linked focal
adhesions. Four members have been identified: Tns1, Tns2,
Tns3 and Tns4 [19]. All isoforms contain a PTB that allows
them to interact with the cytoplasmic tail of β integrin.
The N-terminal region of Tns1 interacts with actin at multiple
sites, thereby linking the actin cytoskeleton to β integrin
[11]. Both α3 integrin- and β1 integrin-KO mice show podo-
cyte abnormalities and proteinuria similar to that of Tns2-
deficient mutants [31]. Although no Tns2 mutation has been found to be associated with human disease, expression of
TNS2 and TNS3 at the mRNA and protein levels was found
to be largely absent in a panel of diverse human cancer cell
lines [21]. The loss of Tns3 leads to greater tumor cell motili-
ity and consequent metastasis, similar to our in vitro results
[21]. Thus, it appears that Tns2 might anchor integrins to the
cytoskeleton or integrins to the ECM, rendering podocytes
stable. In contrast, deleted in liver cancer 1 (Dlc1) is a
recently identified tumor suppressor gene that is frequently
underexpressed in hepatocellular carcinomas (HCCs).
DLC1 encodes a Rho GTPase-activating protein domain
that exhibits growth-suppressive activity in HCC cell lines
[38]. Through its RhoGAP domain, DLC1 inhibits the activ-
ity of RhoA GTPase, which regulates the actin cytoskeleton
network [34]. It has been reported that the Tns2 SH2-PTB
domain binds to the DLC1 protein. Human DLC1 and TNS2
interact and co-localize to punctate structures at focal adhe-
sions, and their interaction is required for tumor suppressive
function [4]. In addition, Dlc1 KD or KO increases actin
stress fiber formation, similar to our result for Tns2 KD [2,
37]. Since Dlc1-KO embryos did not survive beyond 10.5
days post coitum [8], it is unclear whether DLC1 is essen-
tial to maintain podocyte viability and function. The mouse
podocyte mRNA expression database contains mRNA
expression data from FACS-sorted mouse podocytes, as
analyzed by RNA sequencing [18]. This database shows
that both Tns2 and Dlc1 mRNA are highly expressed in
podocytes. These results suggest that both proteins may play
roles in the regulation of actin reorganization in podocytes.
Our Tns2ΔC mice will contribute to determination of which
of the Tns2-integrin and the Tns2-DLC1 signaling axes is
essential for the precise organization and regulation of the
actin cytoskeleton in podocytes.
In conclusion, we show here that Tns2 regulates the podo-
cyte cytoskeleton. Further analysis of Tns2 should provide
a better understanding of the molecular mechanisms of
podocyte cytoskeleton regulation. Moreover, these studies
may lead to the development of podocyte-specific drugs for
restoration of the actin cytoskeleton in podocytes.

Fig. 6. Decreased Tns2 mRNA expression results in increased cell
migration. Transwell assay of Tns2-KD and control cells. Cells
were seeded on the upper side of the inserts. After incubation for
24 hr, migrated cells were fixed and stained with crystal violet
and then quantified by counting the number of cells in each well
(n=3). Representative images of the migration of control and Tns2
KD cells. Magnification, ×200. The number of migrated cells was
counted in each group. In Tns2-KD podocytes, the number of
migrating cells significantly increased.

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