


Novel *NPHS2* variant in patients with familial steroid-resistant nephrotic syndrome with early onset, slow progression and dominant inheritance pattern

Maija Suvanto¹  · Jaakko Patrakka² · Timo Jahnukainen^{1,2} · Pia-Maria Sjöström³ · Matti Nuutinen⁴ · Pekka Arikoski⁵ · Janne Kataja⁶ · Marjo Kestilä⁷ · Hannu Jalanko¹

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Abstract

Background Steroid-resistant nephrotic syndrome (SRNS) is a common cause of end-stage renal disease in children but also occurs as an adult-onset condition. In a subset of SRNS patients, pathogenic variants are found in genes coding for podocyte foot process proteins. The aim of this study was to define the role of pathogenic variants in Finnish patients with familial and sporadic SRNS.

Methods We analyzed SRNS-related genes *NPHS1*, *NPHS2*, *NEPH1*, *ACTN4*, *TRPC6*, *INF2*, *WT1*, *CD2AP*, *LAMB2*, and *PLCE1* for disease-causing variants using direct sequencing of exons and intron/exon boundaries in all members of a family with dominant SRNS with early onset and slow progression to end-stage renal disease. We carried out a whole genome sequencing in two affected and

two healthy family members. The function of found podocin variant was studied using co-immunoprecipitation and immunohistochemistry. Podocyte gene sequences were analyzed in a cohort of Finnish non-familial SRNS patients.

Results A heterozygous de novo deletion, c.988_989delCT in *NPHS2*, was found in all affected family members and in none of their healthy relatives, non-familial patients or controls. No other SRNS-related gene variant, coding or non-coding co-segregated with the disease phenotype in the family. While the truncated podocin remained able to bind nephrin, the expression of nephrin was fragmented and podocin expression reduced. The gene analysis of the non-familial SRNS patients revealed few variants.

Conclusion The role of podocin variants in nephrotic syndrome may be more varied than previously thought.

✉ Maija Suvanto
maija.suvanto@helsinki.fi

¹ Children's Hospital, University of Helsinki and Helsinki University Hospital, Biomedicum 2U, Tukholmankatu 8, 00290 Helsinki, Finland

² Department of Medical Biochemistry and Biophysics, Division of Matrix Biology, Karolinska Institute, Stockholm, Sweden

³ Department of Pediatrics, District Hospital of Pietarsaari, Pietarsaari, Finland

⁴ Department of Pediatrics, University Hospital of Oulu, Oulu, Finland

⁵ Department of Pediatrics, Kuopio University Hospital and University of Eastern Finland, Kuopio, Finland

⁶ Department of Pediatrics, University Hospital of Turku, Turku, Finland

⁷ Department of Chronic Disease Prevention, National Institute for Health and Welfare, Helsinki, Finland

Keywords Podocin · Podocyte · Glomerulus

Introduction

Steroid-resistant nephrotic syndrome (SRNS) manifesting during the first year of life is a rare disorder often caused by variants in genes encoding glomerular podocyte proteins, especially nephrin (*NPHS1*) and podocin (*NPHS2*) [1–5]. SRNS starting later in childhood is in most cases multifactorial with unknown etiology, although the causative factors can include genetic variants. Familial forms of SRNS exist with recessive and dominant inheritance patterns. Variants in *NPHS1* and *NPHS2* are linked to the autosomal recessive inheritance, whereas variants in α -actinin-4 (*ACTN4*), transient receptor potential cation channel subfamily C member 6 (*TRPC6*) and inverted formin 2 (*INF2*), associate with the autosomal dominant

pattern [6]. Altogether, variants in over 30 genes have been associated with SRNS, most of them in exceedingly rare cases or in association with extra-renal symptoms [7].

In this work, we analyzed the genes encoding for the three interacting podocyte proteins nephrin, podocin and neph1 in a cohort of Finnish pediatric SRNS patients including a family with newborn onset of proteinuria and exceptionally slow progression to renal failure. We also analyzed the sequences of *ACTN4*, *TRPC6* and *INF2* in the affected family members and carried out whole genome sequencing in two affected and two healthy family members to discover variants in the additional SRNS-related genes *CD2AP*, *WT1* and *PLCE1* as well as find all non-exonic variants in these SRNS-related genes. The familial disorder followed dominant inheritance pattern and was associated with a heterozygous, de novo mutation in *NPHS2*, encoding podocin.

Materials and methods

Patients

Two pediatric siblings with SRNS (Family AM, Fig. 1) were the starting point of this study. Seven other family members in three previous generations were recruited. Of these, three were inflicted with SRNS. In addition to Family AM, the study cohort included twelve pediatric patients with SRNS who were diagnosed at the University Hospitals in Finland after 1989. Nine of the twelve patients showed FSGS histology and MCNS. The clinical course of the disease of these patients was variable. Two had neurological symptoms.

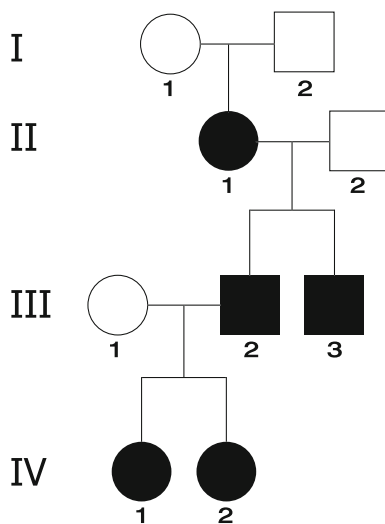


Fig. 1 The pedigree of the Family AM. All shown family members were included in this study. Filled symbols denote affected family members. The same family members carry the deletion c.988_989delCT, which appears to be a de novo variant in II-1. None of the healthy family members have this deletion

Clinical data from patients and their families were gathered from the hospital records. Blood samples were collected during routine checkups, or on request in their local health centers.

Genetic analyses

Genomic DNA was isolated from frozen or fresh peripheral blood samples using the conventional molecular biology technique (Puregene EPTM DNA Purification Kit, Gentra Systems). The analyses of protein coding regions of *NPHS1* (NM_004646), *NPHS2* (NM_014625), *NEPH1* (NM_018240), and *ACTN4* (NM_004924) as well as exons 2 and 4 of *INF2* (NM_022489) were performed using direct sequencing. Exons were amplified by PCR with flanking intronic primers. The reactions were performed in total volumes of 25 μ l as previously described [8]. Occasionally, the denaturation temperature was raised up to 98 $^{\circ}$ C, or betaine was added to the reaction mixture. PCR products were subjected to automated sequence analysis by BigDye-terminator chemistry (v.3.1) on Genetic Analyzer 3730 (Applied Biosystems). The sequences were analyzed with GeneComposer version 1.1.0.1051 (www.GeneComposer.com) or 4Peaks version 1.7.2 (by A. Griekspoor and Tom Groothuis, mekentsoj.com.).

When a nucleotide change resulting in an amino acid change was spotted, the significance of the change was studied by sequencing the same exon from 23 to 51 healthy controls except in the case of the deletion carried by Family AM which was checked from 101 healthy controls. In addition, the healthy members of Family AM were sequenced as controls.

In addition, Blueprint Genetics was commissioned to carry out a Nephrotic Syndrome sequencing panel of nine genes (<http://blueprintgenetics.com/nephrotic-syndrome-panel/>) to four healthy and three affected members of Family AM. While the panel included *NPHS1*, *NPHS2*, *ACTN4* and *TRPC6* that were previously sequenced it also contained all the rest of the *INF2* exons as well as genes *CD2AP*, *LAMB2*, *PLCE1* and *WT1*. The panel targets all exons, intron/exon boundaries and known mutations outside these regions.

The effects of amino acid changes were evaluated using online bioinformatics tools Predict Protein server [9], Prosite database [10] and PolyPhen [11].

Whole genome sequencing (WGS) was carried out in two affected (the first affected in the family, II-1 and her grand daughter, IV-2) and two healthy (I-1 and II-2) members of the family AM in the Finnish Institute of Molecular Medicine (www.fimm.fi, University of Helsinki, Biomedicum, Helsinki Finland). The aim was to look for sequence variants where the genotypes of the affected differ from the healthy in accordance with dominant

inheritance pattern. The analysis revealed over 100,000 different type of variants spread across all autosomal chromosomes and chromosome X. We concentrated on variants in the coding and non-coding areas of genes associated with SRNS in the literature. As the disease in Family AM shows no extra-renal expression we focused our efforts on genes causing isolated SRNS including *NPHS2*, *NPHS1*, *INF2*, *CD2AP*, *WT1*, *TRPC6*, *ACTN4*, *PLCE1*, *MYH9*, *Myo1E* and *ARHGAP24*. The WGS data analysis was conducted using Integrative Genome Viewer (IGV) (www.broadinstitute.org/igv/).

Plasmids

Wild-type (WT) and c.988_989delCT-mutated *NPHS2* sequences were inserted into pCMV-Myc-vector (Clontech, Palo Alto, CA, USA) to give them N-terminal Myc-tag. The WT-*NPHS1* was inserted into pcDNA3.1-vector (Invitrogen, Carlsbad, CA, USA). Plasmids were amplified using JM109 competent *E. coli* cells, QIAGEN Plasmid Maxi Kit (QIAGEN inc. Valencia, CA, USA) and the protocol provided by the kit.

Co-immunoprecipitation of nephrin and podocin

HEK293FT cells were transiently transfected with two plasmids: one carrying WT-nephrin, the other carrying either WT-podocin or c.988_989delCT-mutated podocin. The transfection was carried out using FuGENE 6-protocol (Roche, Basel, Switzerland). After incubating for 24 h, the cells were washed in PBS and lysed in non-denaturing lysis buffer containing 20 mM TrisHCl pH8, 139 mM NaCl, 10 % glycerol, 1 % NP40, 2 mM EDTA, and phosphatase and proteinase inhibitors. Cell lysates were incubated with c-Myc (A-14) antibody (Santa Cruz Biotechnology Inc., Santa Cruz, CA, USA) over night at +4 °C on gentle agitation. After this, 50 µl 50 % rec Protein G-Sepharose beads (Invitrogen, Carlsbad, CA, USA) were added to the antibody-protein complex mix and incubated over night at

+4 °C on gentle agitation to bind the antibody to the beads. Beads were washed with the lysis buffer and fractionated by SDS-PAGE. Western blot analysis was performed using nephrin antibody (Santa Cruz Biotechnology Inc., Santa Cruz, CA, USA) followed by the incubation with horseradish peroxidase-coupled anti-goat antibody. Immobilized antibodies were detected using chemiluminescence.

Immunofluorescence staining

Antibodies used for immunofluorescence staining of cryosections of the kidney tissue samples were goat polyclonal antibody against the C terminus of the human podocin C-18 (sc-22296, Santa Cruz Biotechnology, Santa Cruz, CA), goat polyclonal antibody against the N terminus of the human podocin N-21 (sc-22294, Santa Cruz Biotechnology, Santa Cruz, CA), and rabbit polyclonal antibody against human nephrin (UP3) (by courtesy of Karl Tryggvason lab, Karolinska Institute, Stockholm, Sweden). The staining was carried out in the traditional method using snap-frozen, acetone-fixed cryosections (5 µm) of kidney tissue. Samples included thin needle biopsy of affected family member (IV-1 in Fig. 1), nephrectomised tissue from congenital nephrotic syndrome of the Finnish type (CNF) patient and normal human kidney from adult donors as control. Light microscopy was performed with a standard Leica DM RX light microscope equipped with an Olympus DP70 digital camera.

The binding of the antibodies C-18 and N-21 to truncated podocin was confirmed using Western blot analysis (Fig. 2a).

Results

Clinical findings in the familial SRNS

Clinical findings of the affected members of Family AM are cataloged in Table 1. In four of the five affected family members, mild proteinuria appeared during early

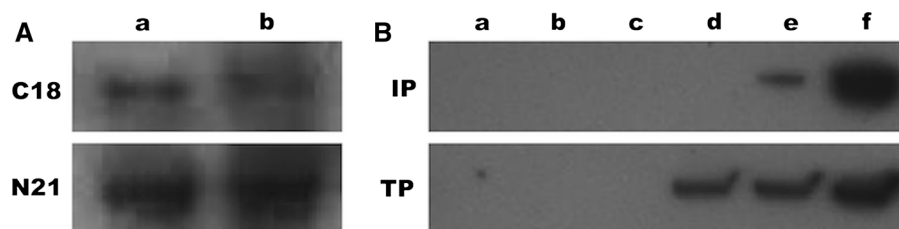


Fig. 2 Analysis of wild-type and c.988_989delCT podocin of patient IV-1. **a** Binding of antibodies recognizing the N terminus (N21) and C terminus (C18) of podocin to WT and c.988_989delCT podocin in transfected HEK293FT lysates. **a** WT-podocin. **b** c.988_989delCT podocin. **b** Co-Immunoprecipitation of podocin and nephrin visualized with α -nephrin antibody. **IP** Immunoprecipitation. **TP** Total

protein. **a** Lysates of HEK 293FT. **b** Lysates of HEK 293FT cells transfected with WT-podocin. **c** Lysates of HEK 293FT cells transfected with c.988_989delCT podocin. **d** Lysates of HEK 293FT cells transfected with nephrin. **e** Lysates of HEK 293FT cells transfected with both WT-podocin and nephrin. **f** Lysates of HEK 293FT cells transfected with both mutated podocin and nephrin

Table 1 Clinical features of SRNS in the affected members of Family AM

Patient	Age of onset (years)	Proteinuria (g/l)	Hematuria	Serum albumin (g/dl)	Creatinine [mg/dl (umol/l)]	Histology	Dialysis started (years)	Transplant (years)
II-1	18 ^a	2–3 > 8 ^b	Microscopic ^c	N/A	0.62 (55) > 0.59 (52) ^b	MCNS > FSGS ^b	40	41
III-2	2.5	1–3 > 14 ^d	No	1.3 ^e	0.7 (62) ^e	MCNS ^e	25	31
III-3	1.8	0.2 > 14.5 ^f	No	1.7 ^g	1.0 (90) ^g	FSGS ^g	n/a ^h	n/a ^h
IV-1	0	0.53 > 2.1–3.6 ⁱ	Microscopic	2.1 – > 1.5–2.5 ⁱ	0.24 (21) > 0.25 (22) ⁱ	MCNS ^j	n/a ^h	n/a ^h
IV-2	0	2.26 > 0.21 ^k	No	2.8 –> 1.5 ^k	0.43 (38) > 0.16 (14) ^k	n/a	n/a ^h	n/a ^h

^a Proteinuria was detected in a healthy control. As the condition was at that point symptomless, the precise age of onset cannot be determined

^b Values from onset to age 25 years

^c Finding at the age of onset

^d Values from onset to age of 17 years

^e Value at 17 years

^f Values from onset to age of 22 years

^g Value at 22 years

^h The patient had not progressed to ESRD at the time of the latest checkup

ⁱ Values from onset to age of 12 years

^j Value at 9 years

^k Values from onset to age of 3.5 years

childhood. The fifth affected (II-1 in Fig. 1) was diagnosed in a routine checkup in the age of 18 years, as her proteinuria had been symptomless. In two of the five affected (II-1, III-2), the proteinuria slowly increased leading to ESRD in adulthood. The kidney function of third affected (III-3) has been worsening but he has yet to reach ESRD, and the other two affected (IV-1, IV-2) are still children. In biopsy samples taken during childhood or when the proteinuria was still mild (II-1, III-2, IV-1), the histology of the affected shows only MCNS while biopsies taken after the kidney function has started to deteriorate show FSGS (II-1, III-3). All patients were given corticosteroid medication with little effect. Other members of Family AM (I-1, I-2, II-2, III-1) are healthy with no signs of kidney disease.

NPHS2 in the familial NS

Genetic analysis was performed on five cases and four healthy family members (Fig. 1). Sequencing of the podocin gene (*NPHS2*) revealed a heterozygous de novo deletion in exon 8 (c.988_989delCT) in all affected cases and in none of the healthy family members or in the unrelated controls. To see if a second mutation would lie outside the analyzed exons and exon/intron boundaries, NS panel of nine genes was commissioned. In *NPHS2*, an intronic variant (rs115778946) was discovered in all three affected family members included in the analysis and in

one healthy family member. However, the first affected family member (II-1) has inherited this invariant from her mother and passes it on to her children together with c.988_989delCT. The father of her affected children (II-2) and the mother of her affected grandchildren (III-1) do not carry it. Thus, it appears that both found variants lie in the same allele of the gene and all affected have one allele of *NPHS2* with no disease-causing variants. The WGS of two affected and non-affected family members was carried out to reveal variants that are unreachable for the gene panel. Only a few non-coding variants were found and these did not follow such inheritance pattern that could explain the phenotype.

c.988_989delCT results in a 343 aa polypeptide instead of the 382 aa wild type. The Prosite database calculated that the mutated sequence lacked one predicted N-glycosylation site (from aa 355: NRTQ), two protein kinase C phosphorylation sites (from aa 332: TEK and from aa 354: SNR) and a cAMP- and cGMP-dependant protein kinase phosphorylation site (from aa 377: KKDS).

Co-immunoprecipitation of normal and c.988_989delCT-podocin and nephrin showed that the truncated podocin bound to nephrin similar to the normal podocin or slightly more strongly (Fig. 2b). Immunohistochemistry for nephrin in biopsy sample of IV-1 showed that while the protein was expressed in kidney glomeruli, the staining pattern was uneven and discontinuous. The

staining of podocin was done using two antibodies, one staining the N terminus of the protein and the other the C terminus carrying the variant. The epitope recognized by the C-terminal antibody C-18 is partially disrupted by the c.988_989delCT variant, and thus the ability of C-18 to bind mutated podocin was tested with Western blot analysis from the transfected HEK293 lysates. Figure 2 A shows that the C-18 binding of c988_989delCT podocin is similar to WT-podocin. The immunostaining of podocin shows that while the C-terminal antibody stains slightly less than the N-terminal antibody in c.988_989delCT mutated glomeruli, the staining of both is greatly reduced when compared to the glomeruli of normal kidney tissue (Fig. 3).

Other podocyte genes in the familial NS

Sequencing of the podocyte protein coding genes *NPHS1*, *Neph1* and *ACTN4* uncovered only known polymorphisms. Similarly, none of the patients had variants leading to aa

changes in *INF2* exons 2 and 4, where majority of the mutations leading to dominant FSGS are clustered [12].

The NS panel of Blueprint Genetics did not find any variant in any gene that would have followed the disease phenotype. The WGS revealed one variant that was shared by the two analyzed affected family members (II-1 and IV-2) but not the two healthy ones. Direct sequencing of this non-coding variant, a nucleotide deletion (rs76500597) in *WT1*, in rest of the family members revealed no shared genotype between the affected individuals: one affected did not carry the deletion while two healthy family members did (one as a homozygote). All sequence variants of Family AM in SRNS-associated genes can be viewed in Table 2.

Genes in non-familial cases of SRNS

One of the twelve sporadic SRNS cases in the study had aa changing variants in *NPHS2*, p.P341R in one allele and p.R229Q in two alleles. SRNS was diagnosed in this

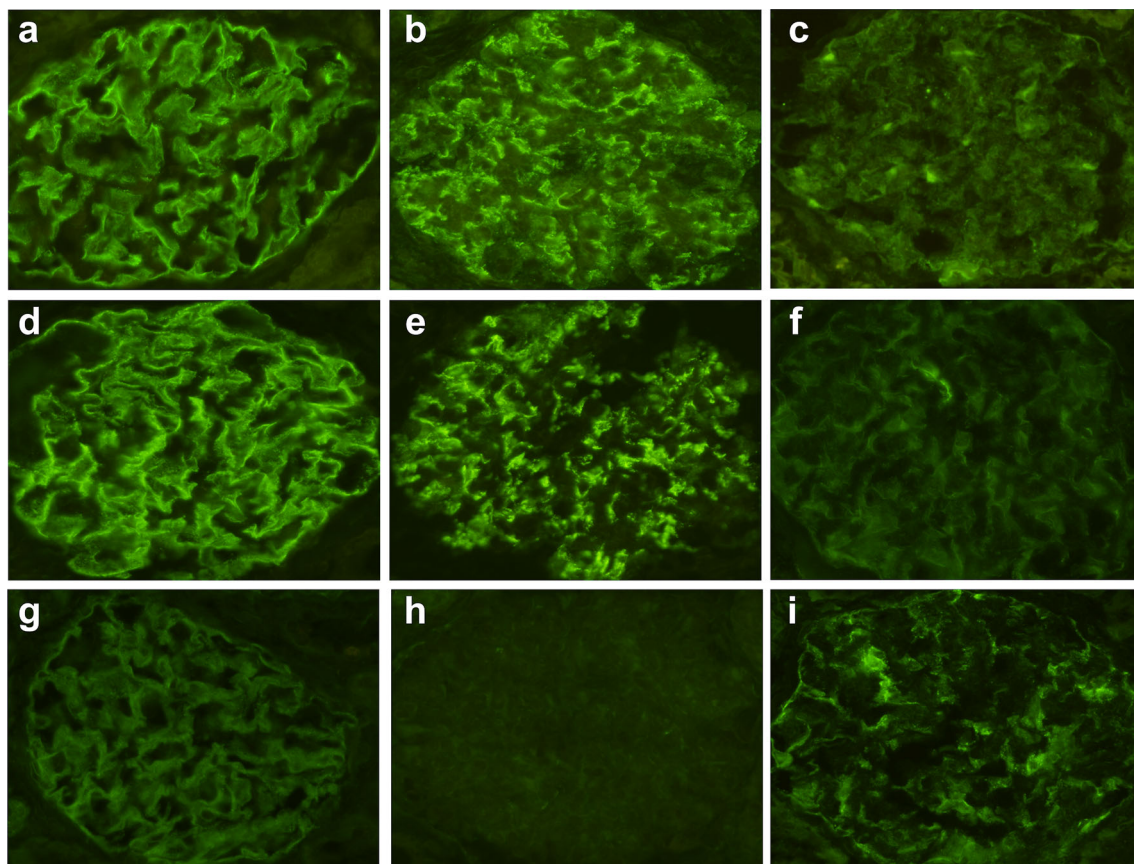


Fig. 3 Immunofluorescence staining of podocin and nephrin. Images magnified $\times 20$. **a** Podocin staining in normal human kidney by N21 antibody. **b** Podocin staining in CNF kidney by N21 antibody. **c** Podocin staining in c.988_989delCT-mutated kidney of patient IV-1 by N21 antibody. **d** Podocin staining in normal human kidney by C18

antibody. **e** Podocin staining in CNF kidney by C18 antibody. **f** Podocin staining in c.988_989delCT-mutated kidney of patient IV-1 by C18 antibody. **g** Nephrin staining in normal human kidney. **h** Nephrin staining in CNF. **i** Nephrin staining in c.988_989delCT-mutated kidney

Table 2 Genetic variants in known SRNS-causing genes in Family AM

Gene/variant	MAF	Variant type	Nucl. change	CDS pos.	AA change	Prot. pos.	Family AM									
							I-1	I-2	II-1	II-2	III-1	III-2	III-3	IV-1	IV-2	
<i>NPHS2</i>																
n/a	N/A	Deletion	CAG>C	988–989	L>Vfs*14	330	–	–	het	–	–	het	het	het	het	–
rs115778946	C = 0.0160	Intronic	T>C	N/A	N/A	N/A	het	–	het	–	–	n/a	het	n/a	het	–
<i>LAMB2</i>																
rs146522641	T = 0.0006	Missense	C>T	4163	R>Q	1388	–	–	–	–	het	n/a	–	n/a	–	–
rs112933248	n/a	Missense	T>C	4118	D>G	1373	–	–	–	het	–	n/a	het	n/a	–	–
rs34759087	T = 0.0593	Missense	C>T	2959	E>K	987	–	–	–	het	–	n/a	–	n/a	–	–
<i>PLCE1</i>																
rs17508082	A = 0.0172	Missense	T>A	481	S>T	161	–	het	–	–	het	n/a	–	n/a	het	–
rs2274224	C = 0.4862	Missense	G>C	4724	R>P	1575	het	het	het	–	–	n/a	het	n/a	het	–
rs3765524	T = 0.3079	Missense	C>T	5330	T>I	1777	het	het	het	–	–	n/a	het	n/a	het	–
rs17516758	A = 0.1068	Intronic	G>A	N/A	N/A	N/A	het	–	het	–	–	n/a	het	n/a	het	–
rs2274223	G = 0.2985	Missense	A>G	5780	H>R	1927	het	het	het	–	–	n/a	het	n/a	het	–
<i>WT1</i>																
rs76500597	– = 0.3123	Intronic; del	C>–	N/A	N/A	N/A	hom	–	het	het	–	het	het	het	het	–
<i>TRPC6</i>																
rs36111323	A = 0.0643	Missense	G>A	1211	A>V	404	het	–	het	–	–	n/a	het	n/a	–	–
rs117273916	A = 0.0008	Missense	G>A	172	R>W	58	–	het	het	–	–	het	–	–	het	–
rs3802829	A = 0.0677	Missense	G>A	43	P>S	15	het	–	–	–	–	n/a	–	n/a	–	–
<i>INF2</i>																
rs3840006	A = 0.3051	Intronic; ins	–>A	N/A	N/A	N/A	het	–	–	–	het	n/a	–	n/a	het	–
<i>NPHS1</i>																
rs77309273	A = 0.0198	Intronic	C>A	N/A	N/A	N/A	–	hom	het	–	–	n/a	–	n/a	–	–
rs113825926	A = 0.0038	Missense	G>A	881	T>I	294	–	–	–	–	het	n/a	–	n/a	–	–
rs34982899	C = 0.0108	Missense	G>C	791	P>R	264	–	–	–	het	–	n/a	–	n/a	–	–
rs3814995	T = 0.2919	Missense	C>T	349	E>K	117	–	hom	het	het	–	n/a	het	n/a	–	–

MAF minor allele frequency, CDS coding sequence, AA amino acid, *het* heterozygous appearance of minor allele, *hom* homozygous appearance of the minor allele. Wild type is noted as –. N/A not applicable, n/a not available. Affected family members are marked with bolded identifier

patient at the age of six and she is now waiting for the kidney transplant at the age of 19 years.

In the *NPHS1* gene, six of the eleven patients had one or more of three known amino acid changing polymorphisms (p.E117K, p.R402Q, p.N1077S), which were also displayed in the healthy controls in similar frequencies. No patient carried amino acid changing variants in the *NEPH1* gene. A few polymorphisms, which did not lead to amino acid changes, were detected in all three genes.

Discussion

We describe a family with an exceptional clinical course of NS. In the affected family members, proteinuria started in the newborn period but remained moderate up to two decades of life until escalating in association with progressive renal failure. ESRD was reached at the age of 25–40 years. While the early kidney biopsies showed

MCNS, FSGS histology became evident after proteinuria worsened. All affected patients had a novel heterozygous deletion c.988_989delCT in the podocin gene (*NPHS2*), which was not present in their healthy family members, in controls or in a cohort of non-familial Finnish SRNS patients. No other mutation was found in *NPHS2* (including non-coding regions) or in other SRNS-related genes.

The *NPHS2* variants cause autosomal recessive form of SRNS [13, 14]. Patients with two *NPHS2* variants usually develop SRNS from birth to 6 years of life and reach ESRD before the end of the first decade [15]. However, phenotypic variability is associated with particular *NPHS2* variants. In patients with p.R229Q in one allele and another disease-causing variant in the other allele, SRNS often starts after the age of 17 years [13, 15, 16]. In cases where only one heterozygous alteration has been identified the disease has started later [2]. It has been assumed the second variant is intronic or in the promoter, and thus escapes detection in the sequencing of the exons.

This seems not to be the case in the family presented here. The disease follows dominant inheritance pattern in three consecutive generations. Undetected variant or variants would have to come into the family from three unrelated sources. The c.988_989delCT is de novo variant in II-1 that occurs in the family at the same time as the disease phenotype appears and then co-segregates perfectly through generations. This suggests that it has a significant role in the NS in the family.

In the literature, other small frameshift variations in *NPHS2* exon 8 have been described [2, 17–21]. Most of these appear as heterozygotes or compound heterozygotes and thus their specific function is difficult to determine. McCarthy et al. [20] reported two FSGS patients with heterozygous exon 8 variant F344fs as well as heterozygous R229Q, and the course of their disease shares similarities with the course of the disease in patients from Family AM with early onset (11 months and 2 years) and only moderate reduction in kidney function after over a decade long follow-up time (13 and 11 years). The effect of R229Q to the phenotype is unclear as its pathogenicity escapes precise definition. Interestingly, Bouchireb et al. [21] described a novel small deletion at almost the same location as c.988_989delCT. The deletion c.989_992del (p.Leu330ProfsX17) was found in one patient as a heterozygote and it is the only variant this patient is reported to have [Leiden Open Variation Database (LOVD 3.0: www.lovd.nl/NPHS2)]. As clinical data concerning the course of disease of this patient are not reported, it is impossible to ascertain if this patient expresses a similar phenotype to the patients in Family AM.

The *NPHS2* deletion c.988_989delCT results in frameshift after amino acid 329. The altered protein gains a novel fourteen amino sequence before a stop codon appears truncating the protein to 343 amino acids instead of the 384 amino acids of the WT-podocin. The truncated podocin had quite normal or slightly elevated affinity to the major slit diaphragm protein nephrin but, in immunofluorescence staining of glomeruli of an affected family member, the expression of nephrin was affected both quantitatively and qualitatively. The Predict Protein revealed that the two- and three-dimensional structure of the podocin was not drastically altered by c.988_989delCT. The protein–protein interactions are often facilitated by amino acid residue proline and c.988_989delCT erases nine proline residues. The truncated podocin may have difficulty to interact with other proteins the same way the WT does.

It is probable that the c.988_989delCT is a gain-of-function variant. These have different mechanisms: hypermorphs increase the activity of the mutated protein, as is the case in *ACTN4* and its disease-causing variants, neomorphs give the mutated protein a new function separate from the WT-protein, and antimorphs tweak the

proteins' function to be antagonistic to the WT. The antimorphs are also known as the dominant negative mutations.

The immunofluorescent staining of podocin was notably reduced in glomeruli of the affected tissue compared to WT. This was seen with both N- and C-terminal antibodies. The amount of reduction leads to assume that it is not only due to c.988_989delCT podocin failing to express properly but that the expression of the WT copy is also hampered. This would suggest that the variant has a dominant negative effect, and it could also explain the altered staining of nephrin, as podocin plays a role in anchoring the protein to the cell membrane. However, the underlying mechanism is unknown and requires more studying and, what the effect of c.988_989delCT may be, its negative impact on the glomerular filtration barrier is likely to be a modest one, leading to the exceptionally slow progression of the disease.

In dominantly inherited SRNS, candidate genes include *ACTN4*, *TRPC6*, *INF2*, *WT1*, *CD2AP* and *ARHGAP* [22–27]. We explored the possibility that the causative mutation would be found in these. None of the variants in these genes followed the disease phenotype.

Besides *NPHS2*, we also included other genes that cause recessive form of NS to the study. No disease phenotype following variant, homozygous or compound heterozygous, was found in *NPHS1*, *PLCE1*, *LAMB2*, *Myo1E* or *MYH9*. As the disease in all affected members of Family AM was unambiguously limited to the kidney, we did not look for variants in genes causing disease characterized by extrarenal manifestations. As genome-wide sequencing was a method used in this study, it should be noted that the weakness of this method is its difficulty in identifying large deletions, insertions and complex rearrangements. It is possible these play a role in the genetic background of NS in this family.

The analysis of *NPHS1*, *NPHS2* and *NEPH1* in the cohort of Finnish pediatric patients with sporadic SRNS showed very little genetic variation. While *Neph1* is associated to proteinuria only in animal models [28] and *NPHS1* mutations are mainly found in congenital NS, previous studies have found frequency of disease-causing *NPHS2* variants in sporadic SRNS to be 10–28 % [3, 19]. In the cohort included to this study, there was only one patient with potentially disease-causing variants, homozygous p.R229Q and novel variant p.P341R in *NPHS2*. This is slightly less than anticipated. Frequencies of disease-causing variants in the genes behind SRNS are known to differ between populations [1, 2, 29]. SRNS is relatively rare in Finland and it is possible that lack of *NPHS2* variation is the reason behind this. However, the cohort size in this study is insufficient to draw conclusions, and larger scale study must be conducted to clarify the matter.

The analysis of NPHS2 variants in sporadic SRNS does, however, act as a type control for the familial SRNS discussed in this article, underlying the rareness of their variant.

This paper outlines an unusual phenotype of familial NS. Although the disease mechanism is not clear we offer evidence that suggests that a small heterozygous deletion in the *NPHS2* gene plays a significant role in its pathogenesis. It seems that not only the altered gene but also the nature of the alteration itself may have an effect on the character of the disease.

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Compliance with ethical standards

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Ethical approval All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee at which the studies were conducted (IRB: HUS 509/E7/05) with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual participants (or their parents) included in the study.

Conflict of interest The authors have declared that no conflict of interest exists.

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