Defects of CRB2 Cause Steroid-Resistant Nephrotic Syndrome

Lwaki Ebarasi,^{1,2,12} Shazia Ashraf,^{3,12} Agnieszka Bierzynska,⁴ Heon Yung Gee,³ Hugh J. McCarthy,⁴ Svjetlana Lovric,³ Carolin E. Sadowski,³ Werner Pabst,³ Virginia Vega-Warner,⁵ Humphrey Fang,³ Ania Koziell,⁶ Michael A. Simpson,⁷ Ismail Dursun,⁸ Erkin Serdaroglu,⁹ Shawn Levy,¹⁰ Moin A. Saleem,⁴ Friedhelm Hildebrandt,^{3,11,*} and Arindam Majumdar^{1,*}

Nephrotic syndrome (NS), the association of gross proteinuria, hypoalbuminaemia, edema, and hyperlipidemia, can be clinically divided into steroid-sensitive (SSNS) and steroid-resistant (SRNS) forms. SRNS regularly progresses to end-stage renal failure. By homozygosity mapping and whole exome sequencing, we here identify recessive mutations in Crumbs homolog 2 (CRB2) in four different families affected by SRNS. Previously, we established a requirement for zebrafish crb2b, a conserved regulator of epithelial polarity, in podocyte morphogenesis. By characterization of a loss-of-function mutation in zebrafish crb2b, we now show that zebrafish crb2b is required for podocyte foot process arborization, slit diaphragm formation, and proper nephrin trafficking. Furthermore, by complementation experiments in zebrafish, we demonstrate that CRB2 mutations result in loss of function and therefore constitute causative mutations leading to NS in humans. These results implicate defects in podocyte apico-basal polarity in the pathogenesis of NS.

Podocytes are highly specialized and polarized epithelial cells that are critical for renal glomerular filtration via their interdigitated foot processes connected by the slit diaphragm.¹ Accordingly, disruption of foot process organization inevitably results in nephrotic syndrome (NS).² Steroid-resistant NS (SRNS) leads to end-stage renal disease.^{3–5} We have recently shown in a cohort of families affected by SRNS that 33% of all cases are caused by mutation in 1 of 21 different genes described in Mendelian forms of SRNS.⁶ However, a large percentage of cases remain molecularly unsolved. To identify additional genes mutated in SRNS in humans, we obtained blood samples and pedigrees after acquiring informed consent from individuals with SRNS and their family members. Approval for human subject research was obtained from the institutional review boards at the University of Michigan and the Boston Children's Hospital. We performed homozygosity mapping (HM)⁷ followed by whole exome sequencing (WES) in these families affected by SRNS. In a family (A1968) of Turkish origin, two siblings had SRNS with renal histology of focal segmental glomerulosclerosis (FSGS) (Table 1). HM in both affected siblings yielded five regions of homozygosity by descent with a cumulative genomic length of ~106 Mb. None of the homozygous peaks coincided with any of seven common recessive causes of SRNS (Figure 1A), suggesting that genes known to be mutated in SRNS were not likely to be involved. By WES in one of the affected siblings from family A1968, we detected a homozygous missense mutation: c.1859G>C (p.Cys620Ser) in exon 7 of CRB2 (crumbs family member 2; RefSeq accession number NM_173689 [MIM 609720]) on chromosome 9 (Figures 1B–1F). This variant was the only homozygous variant remaining from the variant filtering process (Table S1 available online). The mutation alters an evolutionarily conserved cysteine residue within the tenth EGF-like repeat (Figures 1C-1F). It segregated with the affected status in this family and was absent from >190 ethnically matched healthy control individuals and from >6,500 European controls in the Exome Variant Server (Table 1).

By WES in another family (S1232) with an individual affected with SRNS, we identified compound heterozygous mutations: c.1882C>T (p.Arg628Cys) and c.3089_ 3104dup (p.Gly1036Alafs*43) in CRB2 (Figure 1F, Table 1). The heterozygous mutation c.1882C>T (p.Arg628Cys) altered an amino acid residue that was conserved from C. intestinalis to humans and was inherited from the mother (Figure 1G, Table 1). The other heterozygous mutation in this individual was a deleterious duplication of 16 bases c.3089_3104dup (p.Gly1036Alafs*43) in exon 10 of CRB2 (Figure 1F). This variant occurred de novo in the affected individual (Table 1). The duplication was confirmed by PCR amplification, cloning, and sequencing of the genomic DNA from the affected individual (Figure S1).

To discover additional mutations in CRB2, we then performed array-based multiplex barcoded PCR amplification

¹Department of Immunology, Genetics, and Pathology, Uppsala University, 751 85 Uppsala, Sweden; ²Department of Medical Biochemistry and Biophysics, Karolinska Institute, 171 77 Stockholm, Sweden; ³Division of Nephrology, Boston Children's Hospital, Harvard Medical School, Boston, MA 02115, USA; ⁴Children's and Academic Renal Unit, University of Bristol, Bristol BS1 5NB, UK; ⁵Department of Pediatrics, University of Michigan, Ann Arbor, MI 48109, USA; ⁶Department of Experimental Immunobiology, Medical and Molecular Genetics, King's College London, 8th Floor Tower Wing, Guy's Hospital, Great Maze Pond, London SE1 9RT, UK; ⁷Medical and Molecular Genetics, King's College London, 8th Floor Tower Wing, Guy's Hospital, Great Maze Pond, London SE1 9RT, UK; 8Departments of Pediatric Nephrology and Rheumatology, Erciyes University, Kayseri 38039, Turkey; 9Department of Pediatric Nephrology, Dr. Behcet Uz Children Hospital, Izmir 35210, Turkey; ¹⁰HudsonAlpha Institute for Biotechnology, 601 Genome Way, Huntsville, AL 35806, USA; ¹¹Howard Hughes Medical Institute, Chevy Chase, MD 20815, USA ¹²These authors contributed equally to this work

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^{*}Correspondence: friedhelm.hildebrandt@childrens.harvard.edu (F.H.), arindam.majumdar@igp.uu.se (A.M.)

Individual Ethnic Origin Nucleotide Alteration(s) in Conservation(s) Amino Acid Sequence A1968-22 Turkey yes c.1859G>C p.Cys620Ser 7 (HOM, M, P) C.elegans A1968-22 Turkey yes c.1882G>C p.Cys620Ser 7 (HOM, M, P) C.elegans Europe no c.1882C>T p.Arg628Cys 7 (het, M), 10 C. elegans Turkey ws c.3089_3104dup p.Arg628Cys 7 (het, de novo in child) C.intestinalis, NA											
Turkey yes c.1859G>C p.Cys620Ser 7 (HOM, M, P) C. elegans Europe no c.1882C>T p.Arg628Cys 7 (het, M), 10 C. intestinalis, NA Turkey vos c.3089_3104dup) (p.Gly1036Alafs*43) (het, de novo in child) C. intestinalis, NA	Family and Individual	Ethnic Origin	Parental Consanguinity	Nucleotide Alteration(s) ^a	Alteration(s) in Coding Sequence	Exon (Segregation)	Amino Acid Sequence Conservation ^b	PolyPhen-2 Age at Kidney Histology Score Onset Disease (at Age)	Age at Onset	Kidney Disease	Age at Kidney Histology Onset Disease (at Age)
Europenoc.1882C>Tp.Arg628Cys7 (het, M), 10C. intestinalis, NA(c.3089_3104dup)(p.Gly1036Alafs*43)(het, de novo in child)Turbovvocc.1886C>Cof Crosc200cor7 (HOM M. D)C alacano		Turkey	yes	c.1859G>C	p.Cys620Ser	7 (HOM, M, P)	C. elegans	0.989	6 years, SRNS, 4 years SRNS	SRNS, SRNS	FSGS (6 years), FSGS (4 years)
Turbov vas c 1886C-C v Cveč 200a D C alazans	S1232	Europe	ou	c.1882C>T (c.3089_3104dup)	p.Arg628Cys (p.Gly1036Alafs*43)	7 (het, M), 10 (het, de novo in child)	C. intestinalis, NA	0.549, NA	9 mo	SRNS	ND
10000 y 10000 y 10000 y 10000 y 10000 y 10000 y 10000	A3893-21	Turkey	yes	c.1886G>C	p.Cys629Ser	7 (HOM, M, P)	C. elegans	0.997	3 years	SRNS	FSGS (3 years)
A2222-21 Western Europe yes c.3746G>A p.Arg1249Gln 13 (HOM, M, P) C. elegans 0.998	A2222-21	Western Europe	yes	c.3746G>A	p.Arg1249Gln	13 (HOM, M, P)	C. elegans	0.998	QN	SRNS	SRNS FSGS (ND)

and next-generation sequencing⁸ in an additional 1,010 families with SRNS. In an individual from Turkey with SRNS (A3893-21), we detected a homozygous missense mutation (c.1886G>C [p.Cys629Ser]) in exon 7 of *CRB2* (Figure 1F, Table 1). In another individual from an unrelated family (A2222-21), we identified a third homozygous missense mutation: c.3746G>A (p.Arg1249Gln) in *CRB2* (Figure 1F, Table 1). The missense mutation c.1886G>C (p.Cys629Ser) also alters a conserved cysteine within the tenth EGF-like repeat, whereas c.3746G>A (p.Arg1249Gln) changes a conserved arginine in the cytoplasmic tail of CRB2 (Figure 1F). Renal biopsy revealed FSGS in four of the five individuals (Table 1).

CRB2 spans 22.49 kb on chromosome 9q33.4 (Figure 1C). The longest transcript of CRB2 (RefSeq NM_173689 [MIM 609720]) has 13 coding exons (Figure 1D). As a result of alternative splicing, CRB2 encodes two isoforms: isoform 1, a putative type I transmembrane protein of 1,285 amino acids (Figure 1E), and isoform 2, a secreted protein of 1,176 amino acids.9 CRB2 is known to contain 15 extracellular EGF-like domains and 3 extracellular laminin G-like domains (Figure 1E). Interestingly, three of the identified missense mutations (p.Cys620Ser, p.Arg628Cys, and p.Cys629Ser) occur within exon 7 of CRB2, which encodes the extracellular tenth EGF-like domain of this protein. This suggests that the tenth EGF-like domain might play an important role in CRB2 function in podocytes. Interestingly, many other disease-associated missense mutations affect amino acids in the well-conserved EGF-like repeats and laminin A domains of the paralog CRB1, implying an important function for the extracellular region of CRB1 in human retinal dystrophies.^{10,11}

We performed immunofluoresence staining in rat kidneys and demonstrated that CRB2 is expressed in podocytes in adult rat glomeruli (Figure 2). CRB2-positive staining was seen in cells positive for the podocyte markers WT1, GLEPP1, SYNAPTOPODIN, and PODOCALYXIN (Figure 2). CRB2 colocalizes most tightly with GLEPP1 among podocytic markers used in the immunofluorescence, consistent with the localization of CRB2 at the slit diaphragms of podocytes (Figure 2C).

In an earlier study, we reported that morpholinoinduced knockdown of zebrafish crb2b resulted in podocyte foot process defects with ensuing proteinuria.¹² To genetically define crb2b function in podocyte differentiation, we now obtained a stable heritable loss-of-function mutation in crb2b. The crb2b mutant allele was caused by a retroviral murine leukemia virus (MLV) insertion in the *crb2b* locus and is transmitted to offspring as a recessive mutation in Mendelian ratios (see Supplemental Methods).^{13,14} crb2b^{-/-} homozygous embryos are indistinguishable from $crb2b^{+/-}$ sibs up to 4 days postfertilization (dpf), after which they show pronephric cysts and pericardial edema, both indicators of kidney dysfunction (Figures 3A and 3B). By 5 dpf, $crb2b^{-/-}$ embryos have smaller eyes, consistant with requirement in photoreceptor differentiation.^{15,16} The pronephric and eye phenotypes are due to

specific loss of *crb2b* gene function, as shown by the fact that both can be rescued by injection of full-length zebra-fish *Crb2b* mRNA (Figure 3C).

Histological sectioning showed glomerular morphogenesis defects in $crb2b^{-/-}$ homozygotes (Figure 3D). We next performed electron microscopic analysis of *crb2b^{-/-}* mutant pronephric glomeruli to assess podocyte structure. Ultrastructurally, the $crb2b^{-/-}$ homozygotes show disruption of the regular array of patterned podocyte foot processes, which represents the disapearance of slit diaphragms (Figures 3E–3G). Interestingly, $crb2b^{-/-}$ foot processes contain vesicular-like structures not observed in control $crb2b^{+/-}$ sibs. In addition, the apical membranes of $crb2b^{-/-}$ podocytes show membrane projections that reach into Bowman's space (Figures 3E, 3F, and S2). In $crb2b^{-/-}$ glomeruli, the glomerular basement membrane (GBM) is present but capillary endothelia lack membrane fenestrations (Figure 3F). In control phenotypically wildtype 5 dpf $crb2b^{wt}$ embryos, we counted 2.67 \pm 0.71 fenestrations/ μ m (n = 5 capillary loops from 3 glomeruli). However, in 5 dpf $crb2b^{-/-}$ embryos, we found no glomerular capillary endothelial fenestrations at all (Figure 3G). In order to determine whether glomerular filtration function was affected in $crb2b^{-/-}$ mutants, we performed a dye filtration assay in living 4.5 dpf larvae. Both 500 kDa FITC-labeled and 10 kDa rhodamine-labeled dextrans colocalized within the pronephric proximal tubules, indicating compromised size selectivity in the glomerular filtration barrier (Figure 3H). We conclude that *crb2b* is genetically required for correct foot process arborization and podocyte morphological differentiation.

Because Crb proteins are required for epithelial apical basal differentiation, we examined whether cell polarity might be affected in $crb2b^{-/-}$ podocytes. Phalloidin labels the F-actin network of podocyte foot processes. In phenotypically wild-type *crb2b*^{wt} 4.5 dpf larvae, phalloidin labeled the basal F-actin rich podocyte processes that cover the outer aspect of glomerular capillaries. We found that in $crb2b^{-/-}$, phalloidin is basally concentrated and seen outlining large fused capillaries, indicating that capillary morphogenesis is affected (Figure 4A). Podocyte apical membranes are rich in podocalyxin.¹⁷ In both *crb2b*^{wt} sibs and $crb2b^{-/-}$ embryos, α -Pdxl2¹⁸ staining is present in podocyte membranes, indicating the presence of apical membranes. However, in $crb2b^{-/-}$ podocytes, ectopic α -Pdxl2 membrane extensions are seen in the Bowman's space (insets, Figures 4B, 4C, and 4F), suggesting apical membrane defects. Nephrin is a transmembrane protein component of the podocyte slit diaphragms and basally localized in *crb2b*^{wt} podocytes.^{19–21} In contrast, we found apical α -nephrin¹² localization in $crb2b^{-/-}$ podocytes, indicating defects in nephrin trafficking (insets, Figures 4D and 4G). ZO-1 is a tight junction protein and also found in podocyte slit diaphragms.^{22,23} In *crb2b*^{-/-} podocytes, α -ZO-1 staining was found to be generally reduced (Figure 4E). These results indicate that apical membrane differentiation and protein trafficking of slit components are affected in *crb2b* mutants.

We employed the zebrafish $crb2b^{-/-}$ mutant to test the functional consequences of CRB2 mutations identified in the human families. The human CRB2 open reading frame (RefSeq NM_173689 [MIM 609720]) was synthesized and cloned into pcDNA 3.1 by Genescript. Mutations were introduced into the human CRB2 open reading frame by site-directed mutagenesis. In $crb2b^{+/-}$ $d \times Q$ incrosses, crb2b-/- embryos were generated in Mendelian ratios (Figure 5). However, when an in vitro synthesized mRNA encoding the human wild-type CRB2 was injected, only 9% of the resulting embyros were phenotypically *crb2b* mutant, demonstrating rescue and functional conservation of the human and zebrafish CRB2 genes. Injection of mRNA harboring the human CRB2 mutation c.1859G>C (p.Cys620Ser); $CRB2^{C620S}$ into $crb2b^{+/-}$, $\delta \times$ incrosses resulted in 19% $crb2b^{-/-}$, suggesting that mutation c.1859G>C (p.Cys620Ser) disrupts CRB2 ability to rescue and represents a loss-of-function mutation (Figure 5). The CRB2 protein harboring p.Cys629Ser showed an intermediate level of rescue compared to wild-type CRB2, suggesting a milder loss of protein function compared to p.Cys620Ser.

In this report, we show that heritable mutations in the gene encoding human polarity complex protein CRB2 cause monogenic SRNS in humans. In addition, by testing for phenotypic complementation in the zebrafish $crb2b^{-/-}$ mutant, we were able to demonstrate that these mutations resulted in loss of function and were probably pathogenic alterations in human CRB2. The discovery that *CRB2* mutations cause a recessive Mendelian form of SRNS suggests that the misregulation of podocyte apical basal polarity is an important causative factor in primary FSGS. Foot process arborization, cytoskeletal architecture, trafficking, and membrane biogenesis take part in the regulation of apical basal polarity. Our findings raise the possibility that genes encoding other polarity complex members could also be mutated in heritable and sporadic forms of NS.

Supplemental Data

Supplemental Data include one table and two figures and can be found with this article online at http://dx.doi.org/10.1016/j. ajhg.2014.11.014.

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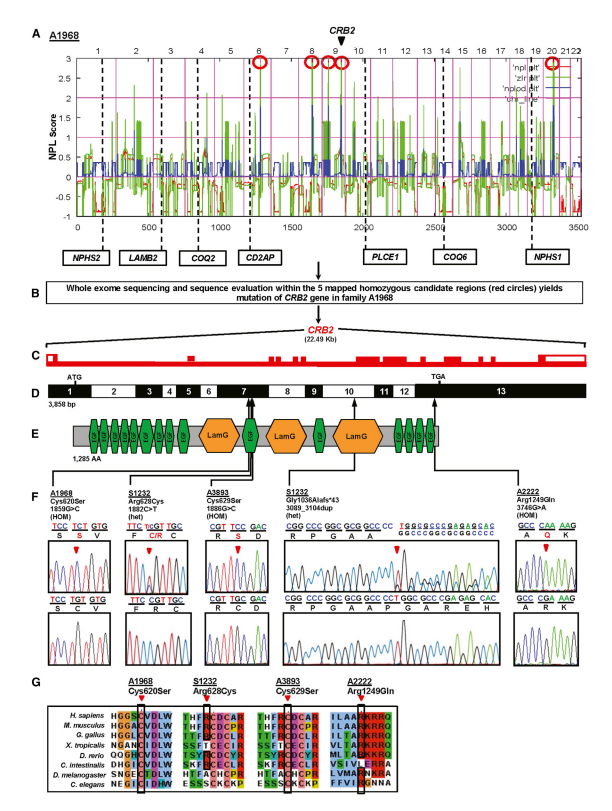


Figure 1. Homozygosity Mapping and WES Identifies *CRB2* Mutations as Causing Steroid-Resistant Nephrotic Syndrome in Humans (A) Nonparametric LOD score (NPL) profile across the human genome in two sibs with SRNS of consanguineous family A1968. SNP mapping was performed with the Affymetrix 250 *Styl* array. SNP positions on human chromosomes are concatenated from p-ter (left) to q-ter (right) on the x axis. Genetic distance is given in cM. Five maximum NPL peaks (red circles) indicate candidate regions of homozygosity by descent. Note that none of the peaks overlap with any of the seven known recessive NS loci.

(B) WES of one of the affected siblings from family A1968 and sequence evaluation within the five mapped homozygous candidate regions (red circles in A) yields mutation of *CRB2* in A1968.

(C) The CRB2 gene extends over 22.49 kb and contains 13 exons (vertical hatches).

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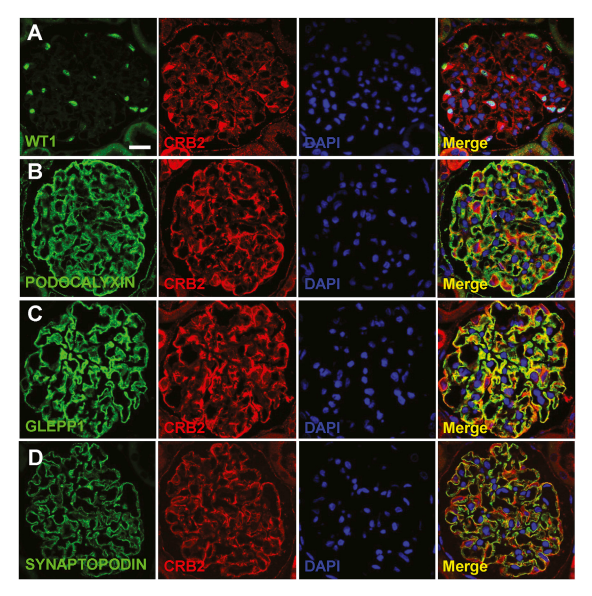


Figure 2. Localization of CRB2 in Adult Rat Kidney

(A) Coimmunofluorescence of CRB2 (Abgent) with WT1 (Santa Cruz Biotech). CRB2 localizes to podocytes, the nuclei of which are marked by WT1.

(B–D) Communofluorescence of CRB2 with podocytic markers PODOCALYXIN (B), GLEPP1 (C), and SYNAPTOPODIN (D) (American Research Products). CRB2 colocalizes most tightly with GLEPP1 among podocytic markers used in immunofluorescence. Note that PODOCALYXIN and GLEPP1 mark the apical podocyte foot process domain, and GLEPP1 is next to the slit membrane adherens junctions. SYNAPTOPODIN marks podocyte processes distal of the slit membrane.

Scale bar represents 10 μm. PODOCALYXIN and GLEPP1 antibodies were kindly provided by Roger C. Wiggins at the University of Michigan.

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(G) The conservation across evolution of altered amino acid residues is shown for all four missense variants (p.Cys620Ser, p.Arg628Cys, p.Cys629Ser, and p.Arg1249Gln).

⁽D) Exon structure of human *CRB2* cDNA. Positions of start codon (ATG) at nt + 1 and of stop codon (TGA) are indicated. For the mutations detected (see F), arrows indicate positions in relation to exons and protein domains (see E).

⁽E) Domain structure of the CRB2 protein. 15 EGF-like; calcium-binding domains (green) and 3 Laminin G-like domains (orange) are predicted.

⁽F) *CRB2* mutations detected in four families affected by SRNS. Family number and predicted translational change are indicated (see Table 1). Sequence traces are shown for homozygous mutations above normal controls, and mutated nucleotides are indicated by arrowheads. "HOM" denotes homozygous and "het" denotes heterozygous mutations.

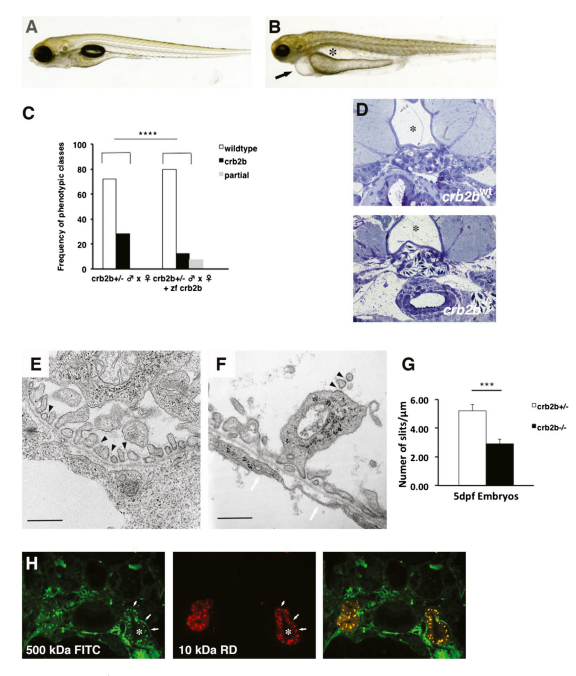


Figure 3. Zebrafish *crb2b*^{-/-} Mutants Have Morphologically Defective Podocytes

(A and B) Brightfield images of zebrafish 5 days postfertilization (dpf): $crb2b^{wt}$ (A) and $crb2b^{-/-}$ larvae (B). $crb2b^{-/-}$ homozygous mutants show reduced eye size, pericardial effusion (arrow), and pronephric cysts (asterisk).

(C) Rescue of the $crb2b^{-/-}$ eye, pronephric, and pericardial effusion phenotypes by injection of full-length zebrafish Crb2b mRNA. ($crb2b^{+/-} \delta \times \Im, n = 75$; $crb2b^{+/-} \delta \times \Im + zebrafish Crb2b$, n = 183 embryos, p < 0.0001). Complete rescue (black bars) and partial rescue (gray bars) frequencies are shown. Phenotypes were scored at 4.5 dpf.

(D) Transverse sections at the level of the glomerulus in $crb2b^{wt}$ and $crb2b^{-/-}$ 5 dpf larvae. In controls, the glomerulus is directly ventral to the notochord (asterisk) and dorsal aorta. In $crb2b^{wt}$, capillary loops are densely packed and covered with podocytes. In $crb2b^{-/-}$ glomeruli, capillary loops are fused together and podocytes are attached to the loops.

(E and F) Electron microsopic analysis of podocyte foot process organization in control $crb2b^{wt}$ (E) and $crb2b^{-/-}$ (F) homozygotes at 5 dpf. In $crb2b^{wt}$, slit diaphragms are visible (black arrowheads in E). The $crb2b^{-/-}$ mutant podocytes show disorganized foot process formation, apical membrane projections containing slit diaphragms (black arrowheads in F) in the urinary space, and a rarefaction of slit diaphragms. A glomerular basement membrane is visible. The endothelium lacks membrane fenestrations in $crb2b^{-/-}$ mutants (white arrowheads in F). Scale bars represent 500 nm.

(G) Quantification of slit diaphragm defects in *crb2b*^{-/-} mutants (p < 0.001, n = 3 regions/glomerulus from 3 different glomeruli). Data represent the mean \pm SEM.

(\dot{H}) Dye filtration assay shows that FITC-labeled 500 kDa (green) and rhodamine-labeled 10 kDa dextran (red) dyes injected into living 4.5 dpf *crb2b^{-/-}* mutants are both passed into and endocytosed (arrows) by the pronephric proximal tubules. The asterisk marks the tubule lumen.

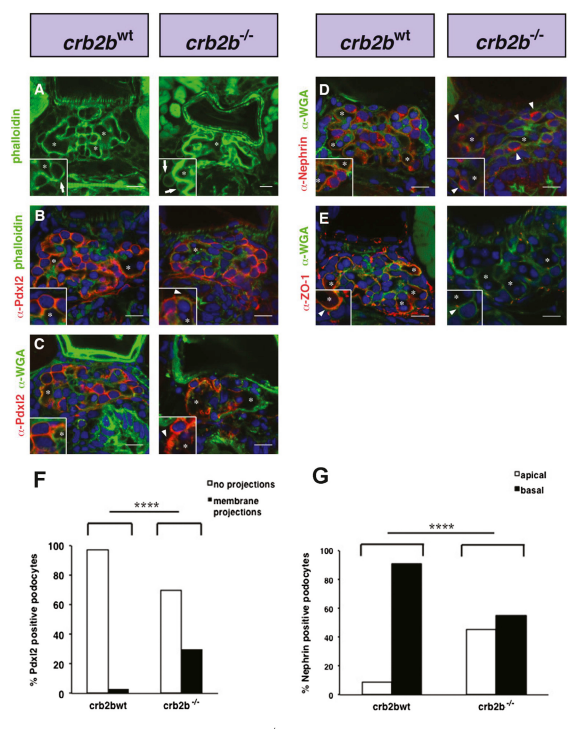


Figure 4. Apical Basal Polarity Is Affected in Zebrafish crb2b^{-/-} Glomerular Podocytes

(A) Phalloidin staining outlines the dense podocyte actin foot process network surrounding capillary lumens. In $crb2b^{-/-}$ mutants, capillary lumens are not compartmentalized but fused into larger vessels. Insets show enlarged images of podocytes. Asterisks mark glomerular capillary lumens.

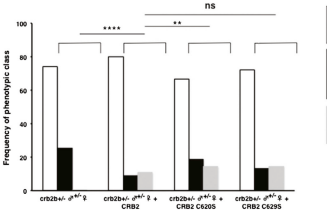
(B) In $crb2b^{wt}$, α -Pdxl2 localizes to apical podocyte membranes. In $crb2b^{-/-}$ mutant podocytes, α -Pdxl2 staining is found in ectopic apical projections (arrowheads).

(C) Glomerular basement membranes are visualized by α -wheat germ agglutinin (α -WGA) staining. Asterisks mark glomerular capillary lumens. Pdxl2 staining is again found in ectopic apical projections in $crb2b^{-/-}$ mutant podocytes (arrowheads).

(D) α-WGA shows glomerular basement membranes. α-Nephrin staining is basally localized in control podocytes but apically mislocalized in $crb2b^{-/-}$ podocytes (arrowheads). (E) α -ZO-1 (Zymed) podocyte staining lines the GBM in $crb2b^{\text{wt}}$ but is diminished in $crb2b^{-/-}$ mutants. Scale bars represent 10 μ m.

(F) Quantification of α -Pdxl2-positive membrane projections in $crb2b^{wt}$ and $crb2b^{-/-}$ podocytes. n = 37 (tallied from 6 embryos) $crb2b^{wt}$ control and n = 30 (tallied from 4 embryos) $crb2b^{-/-}$ podocytes.

(G) Quantification of α -Nephrin localization in $crb2b^{wt}$ and $crb2b^{-/-}$ podocytes. n = 106 (tallied from 9 embryos) control and n = 67 (tallied from 5 embryos) $crb2b^{-/-}$ podocytes. p < 0.0001.



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Web Resources

The URLs for data presented herein are as follows:

Ensembl Genome Browser, http://www.ensembl.org/index.html HomozygosityMapper software, http://www.homozygositymapper. org/

- NHLBI Exome Sequencing Project (ESP) Exome Variant Server, http://evs.gs.washington.edu/EVS/
- Online Mendelian Inheritance in Man (OMIM), http://www. omim.org/

PolyPhen-2, http://www.genetics.bwh.harvard.edu/pph2/

Renal Genes, http://www.renalgenes.org/

RefSeq, http://www.ncbi.nlm.nih.gov/RefSeq

UCSC Genome Browser, http://genome.ucsc.edu

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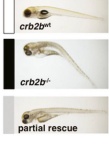


Figure 5. Functional Assay of Human CRB2 Mutations in Zebrafish $crb2b^{+/-}$ $d \times 9$ Incrosses

Phenotypic frequencies of crb2b^{wt} and $crb2b^{-/-}$ mutant embryos after injection of human CRB2 control mRNA and the mRNAs harboring the mutations c.1859G>C; *CRB2*^{C620S} and c.1882C>T; $CRB2^{C629S}$ ($crb2b^{+/-}$ $d \times 9$, n = 363; $crb2b^{+/-}$ $\delta \times$ $\Im + CRB2$, n = 167, p = 0.02; $crb2b^{+/-}$ $\delta \times$ $+ CRB2^{C620S}$, n = 117, p < 0.0001; $crb2b^{+/-}$ $\delta \times$ $+ CRB2^{C629S}$, n =283, p = 0.06; ns, no significant difference). Phenotypic classes of embryos recovered from rescue experiments. Partially rescued embryos have a straight body axis, phenotypically wild-type eyes, and lack pronephric cysts but still show some pericardial effusion. No rescue (black bars) and partial rescue (gray bars) frequencies are shown. Phenotypes were scored at 4.5 dpf.

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The American Journal of Human Genetics Supplemental Data

Defects of CRB2 Cause

Steroid-Resistant Nephrotic Syndrome

Lwaki Ebarasi, Shazia Ashraf, Agnieszka Bierzynska, Heon Yung Gee, Hugh J. McCarthy, Svjetlana Lovric, Carolin E. Sadowski, Werner Pabst, Virginia Vega-Warner, Humphrey Fang, Ania Koziell, Michael A. Simpson, Ismail Dursun, Ervin Serdaroglu, Shawn Levy, Moin A. Saleem, Friedhelm Hildebrandt, and Arindam Majumdar

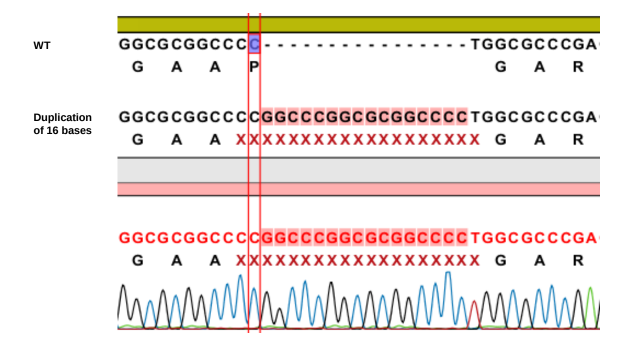


Figure S1. Amplification and cloning using the genomic DNA of the affected individual from family S1232 showed duplication of sixteen bases in exon 10 of CRB2.

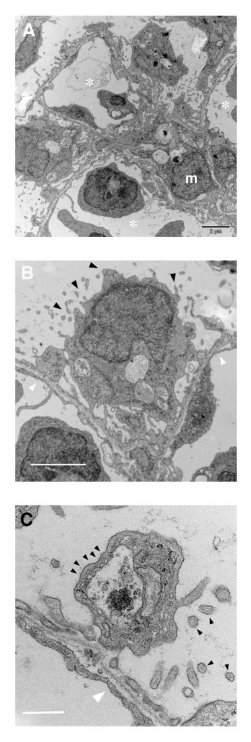


Figure S2. Electron microscopic analysis of crb2b-/- glomeruli and podocytes.

(A) Capillary loops in *crb2b^{-/-}* glomeruli are disorganized. Capillary endothelia lack fenestrations (capillary lumens are marked with asterisks). Mesangial cells (m) can be identified closedly associated with endothelia and surrounded by GBM. Scale bar, 2 mm.
(B) Representative *crb2b^{-/-}* podocyte shows apical membrane projections (black arrowheads) and disorganized foot process structure. Note the absence of endothelial fenestrations (white arrowhead). Scale bar, 2 mm.

(C) Detail of *crb2b^{-/-}* podocyte process. The disorganized process contains vesicular structures (black arrowheads) and apical membrane extensions. The GBM is visible and endothelial membranes lack fenestrations (white arrowhead). Scale bar, 500 nm.

Table S1. Filtering process for variants from normal reference sequence (VRS) following WES in one sibling from family A1968 affected with SRNS.

FAMILY	A1968
^a AFFECTED SIBLING SENT FOR WES	A1968-21
Consanguinity	Yes
^b # of homozygosity peaks	5
Cumulative Homozygosity by descent ^c [Mb]	106
^c Hypothesis from mapping: homozygous (Hom), heterozygous (het)	Hom
Total sequence reads (Mill.)	212
Matched Reads	98.2%
Total DIPs	79,344
Exonic DIPs	325
% exonic / total DIPs	0.40%
DIPS not SNP137	152
DIPS in linked region	24
DIPS after after inspection and not SNP138 (>1% MAF)	0
Sanger confirmation / Segregation	-
Total SNPs	319,298
Exonic SNPs	5,529
% exonic / total SNPs	1.70%
SNPs not SNP137	466
SNPs in linked region	132
SNPs after after inspection and not SNP138 (>1% MAF)	15
^d Sanger confirmation / Segregation	1
Causative gene	CRB2
Mutation effect on gene product	Cys620Ser (Hom)

^asee Table 1

^bsee Fig. 1

^cevaluation for homozygous variants was done in regions of homozygosity by descent for 2 affected sibs. ^dred numbers denote number of filtered-down variant(s) that contained the disease causing gene.

"-", not applicable; DIP, deletion/insertion polymorphism; SNPs, single nucleotide polymorphism; SRNS, Steroid Resistant Nephrotic Syndrome.