

## A Gene Mutated in Nephronophthisis and Retinitis Pigmentosa Encodes a Novel Protein, Nephroretinin, Conserved in Evolution

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Nephronophthisis (NPHP) comprises a group of autosomal recessive cystic kidney diseases, which constitute the most frequent genetic cause for end-stage renal failure in children and young adults. The most prominent histologic feature of NPHP consists of development of renal fibrosis, which, in chronic renal failure of any origin, represents the pathogenic event correlated most strongly to loss of renal function. Four gene loci for NPHP have been mapped to chromosomes 2q13 (*NPHP1*), 9q22 (*NPHP2*), 3q22 (*NPHP3*), and 1p36 (*NPHP4*). At all four loci, linkage has also been demonstrated in families with the association of NPHP and retinitis pigmentosa, known as “Senior-Løken syndrome” (SLS). Identification of the gene for NPHP type 1 had revealed nephrocystin as a novel docking protein, providing new insights into mechanisms of cell-cell and cell-matrix signaling. We here report identification of the gene (*NPHP4*) causing NPHP type 4, by use of high-resolution haplotype analysis and by demonstration of nine likely loss-of-function mutations in six affected families. *NPHP4* encodes a novel protein, nephroretinin, that is conserved in evolution—for example, in the nematode *Caenorhabditis elegans*. In addition, we demonstrate two loss-of-function mutations of *NPHP4* in patients from two families with SLS. Thus, we have identified a novel gene with critical roles in renal tissue architecture and ophthalmic function.

### Introduction

Nephronophthisis (NPHP), an autosomal recessive cystic kidney disease, constitutes the most frequent genetic cause of end-stage renal disease (ESRD) in children and young adults, necessitating renal replacement therapy for survival (Smith and Graham 1945; Fanconi et al. 1951; Hildebrandt 1999). Three distinct gene loci for NPHP—*NPHP1* (MIM 256100), *NPHP2* (MIM 602088), and *NPHP3* (MIM 604387)—have been mapped to chromosomes 2q13 (Antignac et al. 1993; Hildebrandt et al. 1993), 9q22 (Haider et al. 1998), and 3q22 (Omran et al. 2000), respectively. These disease variants share renal histology of interstitial infiltrations, renal tubular cell atrophy with cyst development, and renal interstitial fibrosis (Waldherr et al. 1982). The variants can be distinguished clinically by age at onset of ESRD. The most prominent histologic feature of NPHP is renal fibrosis, which, in chronic renal failure, regardless of origin, represents the pathogenic event that is correlated most strongly to loss

of renal function (Zeisberg et al. 2001). Therefore, NPHP has been considered a model disease for the development of renal fibrosis.

We have previously identified, by positional cloning, the gene (*NPHP1*) for NPHP type 1 (Hildebrandt et al. 1997). Its gene product, nephrocystin, represents a novel docking protein that interacts with the signaling proteins p130Cas, tensin, focal adhesion kinase 2, and filamins A and B, which are involved in cell-cell and cell-matrix signaling of renal epithelial cells (Donaldson et al. 2000, 2002; Hildebrandt and Otto 2000; Benzing et al. 2001). The association between NPHP and autosomal recessive retinitis pigmentosa (RP) has been described as the so-called “Senior-Løken syndrome” (SLS [MIM 266900]) (Løken et al. 1961; Senior et al. 1961). In families with SLS, linkage has been demonstrated to the loci for *NPHP1* and *NPHP3* (Caridi et al. 1998; Omran et al. 2002). Very recently, we have localized a new gene locus (*NPHP4* [MIM 606966]) for NPHP type 4 (Schuermann et al. 2002), and we have demonstrated linkage to this locus in a large kindred with SLS.

We here identified, by positional cloning, the gene (*NPHP4*) that causes NPHP type 4, through demonstration of nine likely loss-of-function mutations in six affected families. In addition, we detected two loss-of-function mutations in patients from two families with SLS. *NPHP4* is a novel gene that is unrelated to any

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known gene families. It encodes a novel protein, nephroretinin. *NPHP4*, like *NPHP1*, is unique to the human genome, is conserved in *Caenorhabditis elegans*, and exhibits a broad expression pattern. It is therefore likely that both gene products, nephroretinin and nephrocystin, interact within a novel shared pathogenic pathway. Thus, we have identified a novel gene with critical roles in renal tissue architecture and ophthalmic function.

## Subjects and Methods

### Pedigree and Diagnosis

We obtained blood samples and pedigrees after receiving informed consent from patients with NPHP and their parents. Diagnostic criteria were (1) development of ESRD in addition to a history of polyuria, polydipsia, and anemia; and (2) renal ultrasound compatible with NPHP. In all families except F461, the diagnosis of NPHP was confirmed by renal biopsy. ESRD developed within a range of 6–35 years of age with a median age of 22 years (table 1). In SLS, the renal symptoms are associated with RP. Clinical data for family F3 (with SLS) have been published elsewhere (Polak et al. 1983; Schuermann et al. 2002). All three affected siblings had RP suggestive of Leber amaurosis congenita. Ophthalmologic data for family F60 have been published elsewhere (Fillastre et al. 1976) and comprise the following observations: In one individual (Fillastre et al. 1976), there was amblyopia and rotary nystagmus with grossly impaired vision starting at age 8 mo, and, on funduscopy, there was retinochoroidal atrophy surrounded by pigment. In two individuals, there were abnormal ERG findings with diminished amplitude (Fillastre et al. 1976).

### Haplotype and Mutational Analysis

The “screening markers” used for haplotype analysis consisted of microsatellites markers *D1S2845*, *D1S2660*, *D1S2795*, *D1S2870*, *D1S2642*, *D1S214*, *D1S2663*, and *D1S1612* (in pter→cen orientation) (Dib et al. 1996). Novel microsatellite markers were generated by searching for di-, tri-, and tetranucleotide repeats, using the BLAST program on human genomic sequence in the interval between flanking markers *D1S2660* and *D1S2642*. Preparation of genomic DNA and haplotype analysis were performed as described elsewhere (Schuermann et al. 2002). Mutational analysis was performed using exon-flanking primers, as described elsewhere (Hildebrandt et al. 1997). Primer sequences can be obtained from the authors.

### Northern Blot Analysis

A multiple-tissue northern blot with human adult poly(A)<sup>+</sup> RNA (MTN7760-1; Clontech) was hybridized with an *NPHP4* DNA probe of 584 bp, derived from exon 30 (nucleotides 4141–4724; see fig. A [online only]) and generated by PCR amplification of human genomic DNA. The probe was labeled with [<sup>32</sup>P]dCTP by using Random Primers DNA Labeling System (Invitrogen). Hybridization was performed at 68°C by using ExpressHyb solution (Clontech). The final washing condition was 0.1 × saline sodium citrate, 0.1% SDS, at 55°C for 40 min.

## Results

We have previously mapped, by total-genome search for linkage, a gene locus (*NPHP4*) for NPHP type 4 within a 2.1-Mb interval delimited by flanking markers *D1S2660* and *D1S2642* (Schuermann et al. 2002).

**Table 1**

**Clinical Details and Mutations Detected in Families with NPHP4**

Family	Number of Affected Individuals	ESRD at Age <sup>a</sup> (years)	RP	Origin	Parental Consanguinity	Exon	Nucleotide Change <sup>b</sup>	Effect on Coding Sequence	Segregation <sup>c</sup>
F3 <sup>d</sup>	3	28, 30, 35	Yes	Turkey	Yes	18	C2335T	Q779X	Hom
F24	2	ND	No	Germany	No	17	G2260A	G754R	P
F30 <sup>d</sup>	3	18, 22, 22	No	Germany	Yes	23	IVS16-1 G→C	Splice site	M
F32	2	19, 20	No	India	Yes	11	TC1334-1335AA	Stop at codon L1121	Hom
F60	4	6, 10, 17, 22	Yes	France	Yes	16	C1972T	R658X	Hom
F444 <sup>d</sup>	2	23, 33	No	Finland	No	15	IVS15+1 G→A	Splice site	M
						24	IVS24+1 G→A	Splice site	P
F461 <sup>d</sup>	3	ND	No	France	No	16	C2044T	R682X	P
						19	C2542T	R848W	M
F622	2	8, 9	No	Afghanistan	Yes	18	G2368T	E790X	Hom

<sup>a</sup> ND = no data available.

<sup>b</sup> All mutations were absent from 92–96 unaffected control individuals.

<sup>c</sup> M = maternal; P = paternal; Hom = homozygous mutation inherited from both parents.

<sup>d</sup> In these four families, linkage to *NPHP4* has been published elsewhere (Schuermann et al. 2002).

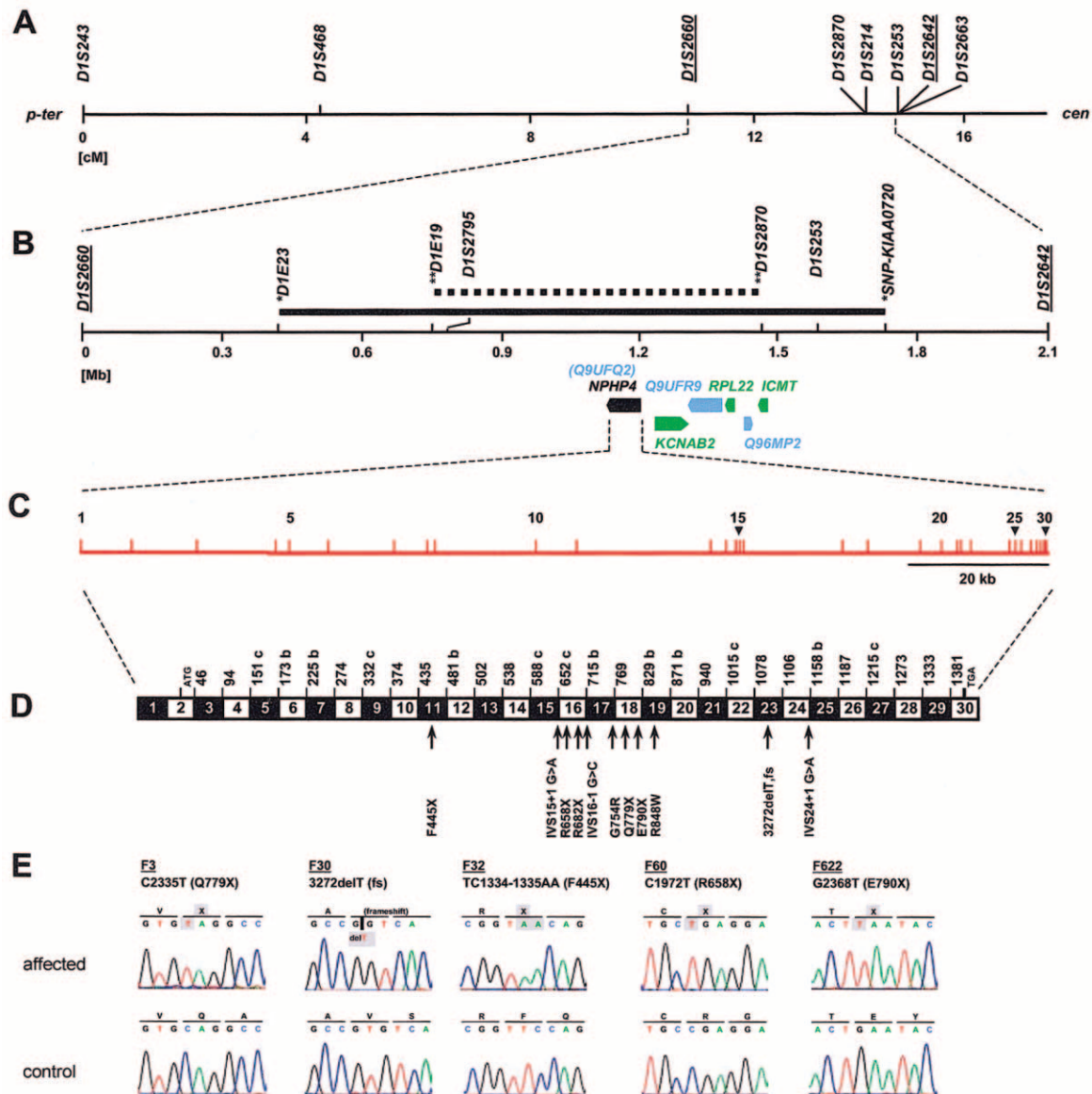
To establish compatibility with linkage to *NPHP4* in further kindreds, we selected 20 families with multiple affected children or parental consanguinity, in whom no mutation was present in the *NPHP1* gene. In 12 families, there was NPHP only, and, in 8 families, there was an association between NPHP and RP. Haplotype analysis using eight microsatellite markers covering the critical *NPHP4* region (Schuermann et al. 2002) was compatible with linkage to *NPHP4* in nine families. To further refine the critical genetic interval of 2.1 Mb, we performed high-resolution haplotype analysis in these nine families and in the seven families with linkage to *NPHP4* published elsewhere (Schuermann et al. 2002). In a total of three families, NPHP was associated with RP. These were two families (F18 and F60) from the new set of nine linked families and one family (F3) that has been described elsewhere (Schuermann et al. 2002). Eight published (Dib et al. 1996) and 38 newly generated microsatellite markers (table A [online only]) were used at an average marker density of one marker per 45 kb within the interval of flanking markers *D1S2660* and *D1S2642* (fig. 1). Haplotype analysis, by the criterion of minimization of recombinants, clearly revealed erroneous inversion of sequence between markers *D1S2795* and *D1S244* in human genomic sequence databases (see the Ensembl Genome Browser Web site) (data available from the authors); this inversion was not described in the recent high-resolution genetic map of the human genome (Kong et al. 2002). Using high-resolution haplotype data, we established the correct marker order at the *NPHP4* locus as pter-*D1S2660*-*D1S2795*-*D1S2633*-*D1S2870*-*D1S253*-*D1S2642*-*D1S214*-*D1S1612*-*D1S2663*-*D1S244*-cen (flanking markers to *NPHP4* are underlined) (Schuermann et al. 2002). A 22-kb sequence gap remaining in the interval *D1S2660*-*D1S2795* was filled by use of Celera human genomic sequence. In haplotype analysis, three consanguineous kindreds yielded new key recombinants by the criterion of homozygosity by descent (Lander and Botstein 1987) (fig. 1). We thus refined the *NPHP4* critical genetic interval to <1.2 Mb within secure borders that were based on a large kindred, and, in addition, to <700 kb within suggestive borders that were based on two small families (figs. 1, 2A, and 2B).

Within the 700-kb critical interval for *NPHP4*, there mapped three known genes (*KCNAB2*, *RPL22*, and *ICMT*) and three unknown genes (*Q9UFQ2*, *Q9UFR9*, and *Q96MP2*) (fig. 2B). In addition, in the interval between *Q9UFQ2* and flanking marker *D1E19* (fig. 2B), the program GENESCAN predicted ~40 nonannotated exons (see the Ensembl Genome Browser Web site) (data not shown). We performed mutational analysis in affected individuals from the 16 families compatible with

	F30 II-2		F30 II-3		F32 II-1		F60 II-1	
<i>p-ter</i>								
<i>D1S2845</i>	201	207	201	207	215	201	219	219
<i>D1S2660</i>	257	259	257	259	261	261	261	261
<i>D1S2660e</i>	166	166	166	166	174	153	166	166
<i>D1S2660i</i>	224	224	224	224	nd	nd	232	232
<i>D1S2660k</i>	149	149	149	149	145	145	149	149
<i>D1S2660h</i>	263	259	263	259	255	255	261	261
<i>D1S2660d</i>	128	136	128	136	134	136	128	128
<i>D1S2660c</i>	155	159	nd	nd	155	169	155	155
<i>D1S2660b</i>	156	154	156	154	158	154	156	156
* <i>D1E23</i>	175	171	175	171	175	175	175	175
<i>D1E22</i>	123	123	123	123	121	121	123	123
<i>D1S2660q</i>	149	149	149	149	149	149	149	149
** <i>D1E19</i>	266	266	266	266	266	272	nd	nd
<i>D1S2795</i>	219	219	219	219	217	217	217	217
<i>D1S2660t</i>	170	170	170	170	170	170	173	173
<i>D1E18</i>	212	212	212	212	212	212	208	208
<i>D1S2660p</i>	197	197	197	197	199	199	191	191
<i>D1S2660u</i>	180	180	180	180	180	180	180	180
<i>D1E17</i>	243	243	243	243	235	235	242	242
<i>D1S2660a</i>	nd	nd	117	117	127	127	115	115
<i>D1S2660r</i>	191	191	191	191	183	183	191	191
<i>D1E16</i>	189	189	189	189	198	198	189	189
<i>D1E15</i>	126	126	126	126	138	138	126	126
<i>D1E14</i>	127	127	127	127	123	123	127	127
<i>D1S2660m</i>	205	205	205	205	209	209	205	205
<i>D1E13</i>	166	166	166	166	174	174	174	174
<i>D1S2633g</i>	236	236	236	236	263	263	246	246
<i>D1S2633e</i>	206	206	206	206	206	206	206	206
<i>D1E12</i>	128	128	128	128	128	128	128	128
<i>D1S2633f</i>	165	165	165	165	173	173	165	165
<i>D1S2633c</i>	161	161	161	161	157	157	161	161
<i>D1E11</i>	142	142	nd	nd	142	142	140	140
<i>D1S2633a</i>	140	140	140	140	140	140	140	140
<i>D1E9</i>	184	184	184	184	184	184	184	184
<i>D1E8</i>	180	180	180	180	180	180	180	180
<i>D1E4</i>	148	148	nd	nd	148	148	148	148
** <i>D1S2870</i>	208	208	208	208	200	200	207	190
<i>D1S253</i>	2	2	2	2	nd	nd	nd	nd
<i>D1S2870c</i>	171	171	171	171	187	187	187	187
<i>D1E3</i>	127	127	127	127	131	131	131	131
*SNP-KIAA0720-Ex19	A	A	A	B	nd	nd	nd	nd
<i>D1S2642f</i>	138	138	138	144	138	138	144	148
<i>D1S2642b</i>	151	151	151	149	151	151	161	166
<i>D1S2642</i>	181	181	181	183	183	183	181	179
<i>D1S214</i>	122	122	122	142	138	138	142	142
<i>D1S2663</i>	201	199	201	193	189	189	197	195
<i>cen</i>								

**Figure 1** Haplotype results on chromosome 1p36 performed for refinement of the *NPHP4* locus in affected offspring from three consanguineous families with NPHP. Family, generation, and individual numbers are indicated above haplotypes. Paternal haplotypes are shown on blue background, and maternal haplotypes are shown on yellow background. A recombination in the maternal haplotype, which was directly observed in parent-to-child transmission in this pedigree, is shown on orange background. At left, 8 published microsatellites are given in italic, and 38 newly generated microsatellites are given in roman. Flanking markers to the published 2.1-Mb critical *NPHP4* interval are depicted in red. Informative alleles are underlined. Haplotypes homozygous in continuity are encased in boxes. Homozygosity mapping revealed a p-terminal recombinant for marker *D1E23* (*asterisks*) on the basis of heterozygosity in individuals F30 II-2 and F30 II-3 and an observed recombinant for marker SNP-KIAA0720-Ex19 (*asterisks*) in individual F30 II-3, thus refining the critical genetic region to a secure interval <1.2 Mb. On the basis of a significant LOD score yielded for F30 alone (Schuermann et al. 2002), this refinement yields secure borders. Further refinement was achieved through heterozygosity for markers *D1E19* and *D1S2870* (*double asterisks*) in individuals F32 II-1 and F60 II-1, respectively. This refines the critical genetic region to a suggestive interval <700 kb. Because of the presence of only two affected individuals in each family, this refinement is only suggestive. p-ter = Telomeric; cen = centromeric; nd = not done.

linkage to *NPHP4*, examining all 79 exons of the three known and three unknown genes by direct sequencing of the forward strands of exon-PCR products. Although no mutations were detected in five of these genes, in *Q9UFQ2*, we detected 11 distinct mutations in 8 of the



**Figure 2** Positional cloning strategy for the *NPHP4* gene, on human chromosome 1p36. **A**, Genetic map position for microsatellites used in linkage mapping of *NPHP4* (see fig. 1). Sex-averaged genetic distance (in cM) from the Marshfield map was used. Published flanking markers are underlined (Schuermann et al. 2002). p-ter = Telomeric; cen = centromeric. **B**, Physical map distances of critical microsatellites relative to *D1S2660*. The secure 1.2-Mb critical interval (solid bar) and the 700-kb suggestive critical interval (stippled bar) are delimited by the newly identified secure flanking markers (asterisks) and suggestive flanking markers (double asterisks) defined by haplotype analysis (see fig. 1). Below the axis, known genes (green), predicted unknown genes (blue), and the *NPHP4* gene (also known as “*Q9UFQ2*”) are represented as arrows in the direction of transcription. **C**, Genomic organization of *NPHP4*, with exons indicated by vertical hatches and numbered. **D**, Exon structure of *NPHP4* cDNA. Blackened and unblackened boxes represent the 30 exons encoding nephroretinin. The number of the first codon of each exon is indicated; exons beginning with the second or third base of a codon are indicated by “b” or “c,” respectively. At bottom, locations of the 11 different mutations identified in eight kindreds with *NPHP4* mutations are shown. fs = Frameshift. **E**, *NPHP4* mutations occurring homozygously in affected individuals from five consanguineous families (underlined). Compound heterozygous mutations are not shown. Mutated nucleotides and altered amino acids are depicted on gray background.

16 families with NPHP (table 1). In families F3 and F60, NPHP is associated with RP. In the affected individuals from all eight families, mutations were shown to segregate from both parents (table 1). All of these mutations

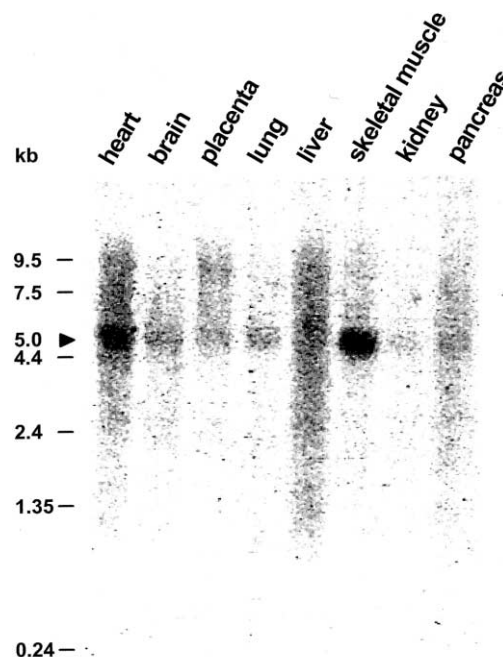
were absent from 92–96 unaffected control individuals from a similar ethnic background. Of the 11 mutations detected, 9 represent very likely loss-of-function mutations: 5 were stop-codon mutations, 1 was a frameshift

mutation, and 3 were obligatory splice-consensus mutations (table 1 and fig. 2D). We thus identified *Q9UFQ2* as the gene that causes NPHP type 4. The gene was termed “*NPHP4*,” and the respective gene product was called “nephroretinin” for its role in NPHP and RP. In the five consanguineous families—F3, F30, F32, F60, and F622—all mutations occurred in the homozygous state and represented stop-codon mutations and one frame-shift mutation, truncating the protein in exons 18, 23, 11, 16, and 18, respectively (table 1 and figs. 2D and 2E). In the three nonconsanguineous families, we found six distinct compound heterozygous mutations. Four represented stop-codon or obligatory splice-consensus mutations, truncating the gene product in exons 15, 16, 17, and 24. The missense mutations R848W and G754R affect amino acid residues conserved in mouse and cow. No mutations were detected in eight families.

*NPHP4* expression studies by northern blot analysis revealed a 5.0-kb transcript expressed strongly in human skeletal muscle, weakly in kidney, and in six additional tissues studied (fig. 3). Northern dot blot analysis (data not shown) confirmed a widespread expression pattern in human adult and fetal tissues, including testis. This broad expression pattern, with strong expression in skeletal muscle and testis, corresponds well with the expression pattern described for the *NPHP1* gene (Otto et al. 2000).

Human genomic sequence of *NPHP4* was assembled using the *Homo sapiens* chromosome 1 working draft sequence segment NT\_028054, which predicted 25 exons. Five additional 5' exons were identified using additional working draft sequence—in particular, the mRNA KIAA0673 and 57 human ESTs from UniGene cluster Hs.106487. The genomic structure shown in figures 2C, 2D, and A was confirmed by human/mouse total-genomic-sequence comparison. The *NPHP4* gene contains 30 exons encoding 1,426 amino acids and extends over 130 kb, with splice sites that confirm to the canonical consensus gt-ag. An exception was found in intron 24, with gc-ag splicing, which occurs in 0.5% of mammalian splice sites (Burset et al. 2001). A polymorphism is known to be present at the intron 20 splice acceptor (tg for ag). The presence of exon 20 is supported by three human EST clones. Ten different splice variants have been suggested for KIAA0673 (see the AceView Web site).

The *NPHP4* cDNA (fig. A) and deduced nephroretinin protein sequences were found to be novel, without any sequence similarity to known human cDNA or protein sequences. Therefore, *NPHP4* encodes a hitherto unknown protein. As shown for the *NPHP1* gene product nephrocystin (Hildebrandt et al. 1997; Otto et al. 2000), there was, however, strong sequence conservation for nephroretinin in evolution, with 23% amino acid identity in a protein of *C. elegans* (fig. B [online only]). Translated



**Figure 3** Northern blot analysis of the *NPHP4* expression pattern. A multiple-tissue northern blot with human adult poly(A)<sup>+</sup> RNA was hybridized with a 584-bp *NPHP4* human DNA probe. Expression of a 5.0-kb transcript (arrowhead) is apparent in all tissues studied, with highest expression being in skeletal muscle.

ESTs also demonstrated evolutionary conservation in mouse, cow, pig, zebrafish, *Xenopus laevis*, *Ascaris suum*, and *Halocynthia roretzi*. Sequence identity of the murine ortholog was 78% (fig. B). Analysis of nephroretinin amino acid sequence (see the ExPASy Molecular Biology Server Web site) provided no signal sequence, conserved domains, or predicted transmembrane regions. Instead, in the N-terminal half, there was a putative nuclear localization signal, a glutamate-rich domain, and a proline-rich domain. The latter two domains have also been found in nephrocystin (Otto et al. 2000). No sequence similarity to nephrocystin was present. In addition, two serine-rich sequences and a C-terminal endoplasmic reticulum membrane domain, were found in human and murine nephroretinin sequences. Encoded by exons 15 and 16, there was, in nephroretinin, a domain of unknown function (DUF339) with evolutionary conservation dating back to prokaryotes and a 63-amino-acid stretch with 30% sequence identity to a gas-vesicle protein of *Halobacterium salinarium* (fig. B).

## Discussion

Our conclusion that we have identified the gene causing NPHP type 4 is based on the identification, in eight families with NPHP, of nine distinct truncating muta-

tions and two missense mutations, none of which occurred in >92 unaffected control individuals. The absence of mutations in eight additional families with NPHP is not surprising, since linkage in kindreds with only two affected individuals can be fortuitous and since NPHP in these families may be caused by mutations in the *NPHP3* gene, which has not yet been identified (Omran et al. 2000). This finding may also indicate further genetic locus heterogeneity for NPHP. Here we also demonstrate the presence of two homozygous truncating mutations in two families with SLS (F3 and F60). Interestingly, a small percentage of patients also exhibit SLS in families with *NPHP1* mutations (Caridi et al. 1998) and in families with linkage to *NPHP3* (Omran et al. 2002). For all three genes, no distinction can be made on the basis of allelic differences between the NPHP phenotypes with and without RP. Therefore, it seems likely that modifier genes may be responsible for the occurrence of RP in NPHP types 1, 3, and 4. *NPHP4*, like *NPHP1*, is unique to the human genome, is conserved in *C. elegans*, and exhibits a broad expression pattern. It is possible that their gene products, nephroretinin and nephrocystin, respectively, interact within a novel shared pathogenic pathway. Since identification of the *NPHP1* gene (Hildebrandt et al. 1997) had revealed nephrocystin as a novel docking protein that interacts with p130Cas (Donaldson et al. 2000; Hildebrandt and Otto 2000), tensin, focal adhesion kinase 2 (Benzing et al. 2001), and filamins A and B (Donaldson et al. 2002), which are involved in cell-cell and cell-matrix signaling, further studies will have to determine the functional role that nephroretinin plays within those interactions. Studies into the function of nephroretinin will provide new insights into disease mechanisms of renal fibrosis, cyst development, and visual function.

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## Electronic-Database Information

Accession numbers and URLs for data presented herein are as follows:

AceView, <http://www.ncbi.nlm.nih.gov/IEB/Research/Acembly/> (for genomic structure)  
 Ensembl Genome Browser, <http://www.ensembl.org/> (for genomic structure of *NPHP4*)

ExpASY Molecular Biology Server, <http://www.expasy.ch/> (for Tmpred)  
 Hits Home, <http://hits.isb-sib.ch/> (for MOTIF SCAN program)  
 Marshfield Medical Research Foundation Home Page, <http://research.marshfieldclinic.org/> (for map of microsatellite markers)  
 NCBI Blast Home Page, <http://www.ncbi.nlm.nih.gov/BLAST/> (for sequence comparisons through BLASTP/N/X programs)  
 Online Mendelian Inheritance in Man (OMIM), <http://www.ncbi.nlm.nih.gov/Omim/> (for *NPHP1* [MIM 256100], *NPHP2* [MIM 602088], *NPHP3* [MIM 604387], *NPHP4* [MIM 606966], and SLS [MIM 266900])  
 SMART, <http://smart.embl-heidelberg.de/> (for secondary-structure prediction)  
 Submit to GenBank, <http://www.ncbi.nlm.nih.gov/Genbank/> (for full-length *NPHP4* cDNAs in human, mouse, and *C. elegans* [accession numbers BankIt 486643, BankIt 486675, and Z81579] and gas-vesicle protein gvpL of *H. salinarium* [accession number P33964])

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## Erratum

In the November 2002 issue of the *Journal*, the article “A Gene Mutated in Nephronophthisis and Retinitis Pigmentosa Encodes a Novel Protein, Nephroretinin, Conserved in Evolution,” by Otto et al. (71:1161–1167), contained the following error in the Electronic-Database Information sec-

tion: Preliminary BankIt numbers should have been replaced by permanent GenBank numbers for full-length *NPHP4* cDNAs in human, mouse, and *Caenorhabditis elegans* (accession numbers AF537130, AF537131, and Z81579, respectively). The authors regret this error.