Human Adolescent Nephronophthisis: Gene Locus Synteny with Polycystic Kidney Disease in Pcy Mice

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Abstract. In a large Venezuelan kindred, a new type of nephronophthisis was recently identified: Adolescent nephronophthisis (NPH3) is a late-onset recessive renal cystic disorder of the nephronophthisis/medullary cystic group of diseases causing end-stage renal disease at a median age of 19 yr. With the use of a homozygosity mapping strategy, the gene (NPHP3) was previously localized to chromosome 3q22 within a critical interval of 2.4 cM. In the current study, the NPHP3 genetic region was cloned and seven genes, eight expressed sequence-tagged sites, and seven microsatellites were physically localized within the critical disease interval. By human-mouse synteny analysis based on expressed genes, synteny between the human NPHP3 locus on chromosome 3q and the pcy locus on mouse chromosome 9 was clearly demonstrated, thus providing the first evidence of synteny between a human and a spontaneous murine renal cystic disease. By fluorescence in situ hybridization the chromosomal assignment of NPHP3 to chromosome 3q21-q22 was refined. Renal pathology in NPH3 was found to consist of tubular basement membranes changes, tubular atrophy and dilation, and sclerosing tubulointerstitial nephropathy. This pathology clearly resembled findings observed in the recessive pcy mouse model of late-onset polycystic kidney disease. In analogy to pcy, renal cyst development at the corticomedullary junction was found to be an early sign of the disease. Through cloning of the NPH3 critical region and mapping of expressed genes, synteny between human NPH3 and murine pcy was established, thus generating the hypothesis that both diseases are caused by recessive mutations of homologous genes.

Nephronophthisis (NPH), an autosomal-recessive cystic kidney disorder, is a major genetic cause of chronic renal disease in children (1). Recently, we identified adolescent nephronophthisis (NPH3) as a novel type of NPH in a large consanguineous Venezuelan kindred. Applying a homozygosity mapping strategy, we localized the responsible gene (NPHP3) to a critical interval of 2.4 cM on chromosome 3q (2). Juvenile nephronophthisis (NPH1) and adolescent nephronophthisis (NPH3) share the same characteristic renal morphology, consisting of tubular basement membrane changes, tubular atrophy and dilation, sclerosing tubulointerstitial nephropathy, and cysts located predominantly at the corticomedullary junction. However, a gene locus for NPH1 (median age at end-stage renal disease [ESRD], 13.1 yr) was localized to 2q12 to q13, and the responsible gene NPHP1 was found to be homozygously deleted in 70 to 80% of NPH1 patients (3,4). In contrast to NPH1, patients with NPH3 reach ESRD significantly later with a median age of 19 yr (range, 12 to 47 yr), thus rendering NPH3 the recessive renal cystic disease with the latest onset of ESRD (2). The best animal model for this disease seems to be the pcy mouse model for late-onset polycystic kidney disease, because inheritance is recessive, renal pathology resembles that of NPH3, and ESRD is reached in adult mice. To study whether both diseases might be homologous, we compared phenotypic and genetic findings in both diseases. By a physical mapping strategy, we cloned the NPHP3 region, mapped several expressed genes, and the responsible gene NPHP3 to chromosome 3q21-q22 was refined. Renal pathology in NPH3 was found to consist of tubular basement membranes changes, tubular atrophy and dilation, and sclerosing tubulointerstitial nephropathy. This pathology clearly resembled findings observed in the recessive pcy mouse model of late-onset polycystic kidney disease. In analogy to pcy, renal cyst development at the corticomedullary junction was found to be an early sign of the disease. Through cloning of the NPH3 critical region and mapping of expressed genes, synteny between human NPH3 and murine pcy was established, thus generating the hypothesis that both diseases are caused by recessive mutations of homologous genes.
gously affected haplotype enabled us to study the disease phenotype of NPH3 at different stages of the disease.

Materials and Methods

Physical Mapping

A set of 19 yeast artificial chromosomes (YAC) potentially residing at human chromosome 3q21 to q22 was derived from the information available in the public database of the Whitehead Institute (http://www-genome.wi.mit.edu). YAC clones were obtained from Centre d’Etude du Polymorphisme Humain (CEPH; Paris, France). Several polymorphic markers, sequence-tagged site (STS) markers, and expressed sequence-tagged site (EST) markers, presumably residing in the region of interest, were analyzed by PCR amplification individually for positivity with YAC clones by STS content mapping. Information for the examined markers concerning PCR primers, conditions, and location were retrieved through public databases (Whitehead [http://www-genome.wi.mit.edu], Cooperative Human Linkage Center [http://www.chlc.org], Genome database [http://www.gdb.org], Gene Map 99 [http://www.ncbi.nlm.nih.gov/genemap]). Inter-Alu PCR was performed using YAC clones 951a4 and 858b8 as templates (5). PCR products were used as probes for hybridization to screen the Peter de Jong P1 artificial chromosome (PAC) library (6). Bacterial artificial chromosome (BAC) libraries CEPH B and Research Genetics RB (Research Genetics, Huntsville, AL) were screened for STS content. BAC/PAC clones were tested by STS content mapping with the above markers. BAC/PAC ends were sequenced, and PCR primers for clone-end markers were generated. For STS content mapping, one individual colony was picked for each YAC, PAC, or BAC clone. One colony was suspended in 200 μl of sterile water. Ten μl of each sample were dried in a microtiter plate, followed by one negative control (water for template) and one positive control (10 to 30 ng of genomic DNA of a healthy individual). PCR was performed with dry template, 6 to 18 pmol of primers, 0.2 mM control (10 to 30 ng of genomic DNA of a healthy individual). PCR followed by one negative control (water for template) and one positive control. PCR products were used as probes for hybridization to screen the Petcr de Jong P1 artificial chromosome (PAC) library (6). Bacterial artificial chromosome (BAC) libraries CEPH B and Research Genetics RB (Research Genetics, Huntsville, AL) were screened for STS content. BAC/PAC clones were tested by STS content mapping with the above markers. BAC/PAC ends were sequenced, and PCR primers for clone-end markers were generated. For STS content mapping, one individual colony was picked for each YAC, PAC, or BAC clone. One colony was suspended in 200 μl of sterile water. Ten μl of each sample were dried in a microtiter plate, followed by one negative control (water for template) and one positive control (10 to 30 ng of genomic DNA of a healthy individual). PCR was performed with dry template, 6 to 18 pmol of primers, 0.2 mM each dATP, dCTP, dGTP, and dTTP, 10 mM Tris-HCl (pH 7.3), 50 mM KCl, 0.001% gelatin (w/vol), and 0.3 U of Thermus aquaticus DNA polymerase (Perkin-Elmer Cetus, Norwalk, CT). Amplification was carried out with denaturation at 94°C for 30 s, annealing at 53 to 60°C for 30 s, extension at 72°C for 60 s, and 32 cycles. PCR products were separated by electrophoresis in a 1% agarose gel. The gels were stained with ethidium bromide and photographed. Each result was confirmed at least once. BAC/PAC ends were directly sequenced using the Big dye sequencing system (ABI-377, Perkin-Elmer Cetus), and PCR primers for clone-end markers were generated.

Human-Mouse Synteny Analysis

On the basis of the physical mapping data obtained with the studies described above and on the basis of physical mapping data from the Mouse Genome Project (data obtained from http://www.jackson.org), we performed a human-mouse synteny analysis using for DNA sequence comparison the BLASTN subroutine (7). The pcy locus has been mapped to mouse chromosome 9 and is flanked by markers D9Mit16 and D9Mit24 (8). Physical data for the centromeric part of this critical region are available from the Whitehead YAC-contig WC9.35 (http://www-genome.wi.mit.edu). We used all mouse STS and EST sequences of that contig and performed BLAST-N analysis. If no sequence similarity to any human EST/gene was obtained, we conducted a BLAST-N analysis to identify a murine EST cluster. This EST cluster was again used for BLAST analysis to identify homologous human EST or genes.

Fluorescence In Situ Hybridization

Metaphases were harvested after direct preparations of short-term cultures (24 to 48 h) of phytoseomagglutinin-stimulated normal peripheral blood lymphocytes. Methods of cell culture, chromosome preparation, and staining by a modified GAG banding technique have been described previously (9,10). Chromosomal bands were classified according to the International System for Cytogenetic Nomenclature (ISCN, 1995). After fixation in methanol:acetic acid 3:1, the cells were air dried on glass slides and frozen at −20°C for at least 1 d. PAC/BAC DNA of clones RB815E10, 874N07, 544P15, 967G23, 156M15, and 286D11, which are located within the critical genetic NPHP3 region (for exact location of the probes, see Figure 1) was labeled with Texas Red (Molecular Probes, Göttingen, Germany) using a nick translation kit (Vysis, Bergisch Gladbach, Germany). Ten ng of labeled DNA was dissolved in 10 μl of hybridization solution consisting of 50% formamide, 10% dextran sulfate, 1 × SSC, 500 ng of Cot-1 DNA, and 5 μg of sonicated salmon sperm DNA and then denatured at 95°C for 5 min. Slides with metaphase spreads of normal human controls were denatured in 70% formamide/2 × SSC at 72°C for 3 min on a heating plate and subsequently passed through a graded ethanol series. The slides were counterstained with 4,6-diamino-2-phenylindol (2 mg/ml) and mounted in anti-fade solution (90% glycerol, 10% phosphate-buffered saline, 1 mg/ml phenylenediamine). Metaphases were analyzed using Zeiss Axioskop epifluorescence microscope equipped with filter combination I according to Pinkel and a cooled charge-coupled device camera (Metasystems, Alltussheim, Germany) was used. TexasRed and 4,6-diamino-2-phenylindol fluorescence were recorded separately as gray-scale images by changing the excitation wave length only while the beam splitter and emission filter remained in position. The images were then pseudocolored and were merged using a digital image analysis software program (ISIS, Metasystems).

Clinical Studies

In the large Venezuelan kindred with NPH3 (2), blood samples from 13 probands who had not been studied previously were obtained on the basis of informed consent. Haplotype analysis with microsatellites from the NPHP3 region was performed as described previously (2). All probands were studied by measurement of serum creatinine, hemoglobin level, and urinalysis. Renal ultrasound was performed in probands ID202 and ID213 carrying the homozygously affected haplotype of NPH3, thereby cloning the critical region of NPH3, thereby cloning the critical genetic region (Figure 1, a and b). The critical disease interval was flanked by the centromeric marker D3S1292 and the telomeric marker D3S1238. The maximum physical distance for the two markers that flank the NPHP3 region as defined by the combined length of clones 951a4 and 858b8 is 3.3 Mb. We thereby restricted the physical interval for NPHP3 to less than 3.3 Mb (Figure 1b). We physically localized seven expressed genes (ACPP [phosphatase, prostate-specific acid], TOPB1
[topoisomerase (DNA) II binding protein], EDF1 [endothelial differentiation-related factor 1], TF [transferrin], DGKZ [diacylglycerol kinase-zeta], SLC21a2 [prostaglandin transporter, PGT], RYK [receptor-like tyrosine kinase]), eight EST, seven microsatellites, and three STS markers within the critical disease interval for which previously no physical location was known.

**Human-Mouse Synteny Analysis**

Human-mouse synteny analysis was performed on the basis of expressed genes localized to the YAC/PAC/BAC contig generated. Four murine EST residing on Whitehead YAC contig WC9.35 within the pcy critical disease interval showed a very high degree of sequence similarity with human genes/EST of the critical region of NPHP3 (see Figure 1d). The order of the four genes/EST was found to be conserved between human and mouse. This high degree of continuous sequence similarity gives evidence that human NPHP3 and murine pcy map to syntenic genetic regions (Figure 1). For results of BLAST analysis, see Table 1.

**Fluorescence In Situ Hybridization**

Fluorescence in situ hybridization (FISH) analysis with clones RB815E10, 874N07, 544P15, 967G23, 156M15, and 286D11 that are located within the critical disease interval of NPHP3 (Figure 1) revealed positive signals only for human

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**Table 1. Results of sequence similarity analysis using murine EST and genes located in the vicinity of the pcy locus**

<table>
<thead>
<tr>
<th>Mouse EST or Gene</th>
<th>Homologous Human EST or Gene</th>
<th>Sequence Identity</th>
</tr>
</thead>
<tbody>
<tr>
<td>D9Mit24, trf (transferrin)</td>
<td>TF (transferrin)</td>
<td>94/115 (81%)</td>
</tr>
<tr>
<td>04.MMHAP88FRB7.seq</td>
<td>SLC21a2 (prostaglandin transporter)</td>
<td>111/127 (87%)</td>
</tr>
<tr>
<td>D18389, EST-cluster Mm. 1687</td>
<td>TOPBP1</td>
<td>280/327 (85%)</td>
</tr>
<tr>
<td>R74726</td>
<td>EST cluster Hs. 103379 (WI-15706)</td>
<td>169/191 (88%)</td>
</tr>
</tbody>
</table>

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*a* EST, expressed sequence-tagged site.

*b* All human homologous EST and genes that were identified are located within the critical region of NPHP3 (see Figure 1), demonstrating synteny of the human NPHP3 and the murine pcy locus.
chromosome 3q21 to q22. To illustrate these data, the results from FISH analysis using PAC 286D11 is shown in Figure 2.

Clinical Studies

From the 13 probands studied, we identified 2 probands with a homozygously affected haplotype diagnostic for NPH3 (Figure 3). Laboratory investigations and haplotypes for the other 11 probands were normal. Proband ID202 was a 12-yr-old girl without any clinical complaints. Laboratory investigations including maximum urine concentration capacity were normal. Ultrasound of her kidneys revealed multiple small cysts measuring approximately 0.5 cm in diameter at the corticomedullary junction. The male proband ID213 presented with polyuria and loss of appetite at the age of 27 yr. His serum creatinine was 18.8 mg/dl. Within 1 mo of clinical presentation, renal replacement therapy had to be implemented. Ultrasound examination of the kidney revealed multiple cysts in both kidneys. The cysts were predominantly located at the corticomedullary junction but also elsewhere in the kidney (Figure 4, a and b). Histology showed typical findings for NPH3 consisting of tubular basement membranes changes with segments of thickening, thinning, folding and a multilayered appearance, tubular atrophy and dilation, mononuclear infiltrates and diffuse interstitial fibrosis, and concentric periglomerular fibrosis with thickening of Bowman’s capsule (Figure 4d). Ultrasound findings and histologic findings in patient ID213 were compared with findings from macroscopic and microscopic anatomy in the pcy mouse (Figure 4).

Discussion

NPH3 was only recently identified as a novel type of NPH in a large Venezuelan inbred kindred (2). ESRD is reached in NPH3 at a median age of 19 yr, indicating that this disease has a late onset of ESRD, when compared with other recessive renal cystic diseases such as infantile and juvenile NPH and autosomal-recessive polycystic kidney disease (11–13). We previously localized the gene locus (NPHP3) of NPH3 to a 2.4-cM interval on chromosome 3 applying a homozygosity mapping strategy in a large Venezuelan kindred (2).

To identify the responsible gene NPHP3, we cloned the critical NPHP3 region in a complete YAC and partial PAC/BAC contig and physically localized several genes/EST within the region of interest. The maximum length of the critical NPHP3 region is less than 3.3 Mb and corresponds to the combined length of clones 951a4 and 858b8. By FISH analysis using PAC/BAC clones evenly distributed within the critical region of NPHP3, we cytogenetically localized the gene locus to chromosome 3q21 to q22 (Figure 2). On the basis of these human physical mapping data of the critical NPHP3 region, a human-to-mouse synteny analysis was performed and allowed definition of synteny for NPHP3 and a genetic region on mouse chromosome 9 (Figure 1). How far the area of synteny reaches centromeric toward human chromosome 3q21 or murine chromosome 9tel, respectively, cannot be discerned because no additional mouse physical data are available for that region. It is interesting that the genetic region on mouse chromosome 9 contains the polycystic kidney disease (pcy) locus, which was previously mapped to mouse chromosome 9 flanked by D9Mit16 and D9Mit24 (8). Thus, we demonstrate synteny between the human NPHP3 and the murine cystic kidney disease locus pcy (Figure 1). To our knowledge, this is the first report of synteny between a human and a spontaneous mouse cystic kidney disease locus. Among recessive mouse models of renal cystic disease, pcy is the mouse model with the most slowly progressing adult-type form of polycystic kidney disease (14,15), thus resembling the late manifestation of NPH3. Disease traits in pcy and NPH3 share the same autosomal-recessive inheritance pattern. By back-crossing the affected pcy gene to different mouse strains, a variation of the severity...
of the pcy phenotype in relation to different genetic backgrounds was noted (16). Subsequently, two modifier loci on mouse chromosomes 4 and 16 were identified by linkage analysis (17). The presence of modifier genes could also serve to explain the wide age range (12 to 47 yr) of ESRD observed in NPH3. Comparison of NPH3 and pcy revealed several phenotypic similarities, including (1) mode of inheritance, (2) late disease manifestation, (3) renal cyst location at the corticomедullary junction, (4) lack of extrarenal disease manifestations such as hepatic disease, and (5) histology (see also Table 2). There was, however, a dissimilarity with respect to renal enlargement and cerebral vascular aneurysms present in pcy but not in NPH3.

Because all reported patients with NPH3 presented already in ESRD, only little is known about the disease before renal failure. Therefore, we screened new relatives of the large Venezuelan kindred with NPH3, reported previously (2). We identified a 12-yr-old girl with no clinical symptoms, who carries a homozygously affected haplotype for markers from the NPHP3 region (Figure 3). This enabled us to study the disease phenotype of NPH3 before renal failure. Laboratory investigations including urine concentration capacity were normal. Because in ID202 renal sonography showed cysts at the corticomедullary junction, renal cyst development at the corticomедullary junction seems to be an early sign of NPH3 in this proband. The other proband who was diagnosed as having NPH3 was 27 yr old and presented with clinical and laboratory findings of chronic renal failure (Figure 3). Ultrasound of his kidneys showed multiple cysts throughout the entire kidney but predominantly located at the corticomедullary junction (Figure 4). In pcy, cysts tend to develop primarily at a similar site in the kidney (Figure 4). Takahashi et al. (15) noted in DBA/2-pcy/pcy mice by 4 wk of age a well-delineated cyst in the outer portion of the inner medulla in almost every sample examined, and in the latest stages of the disease (30 wk and older) the whole kidney was enlarged and dominated by cysts. Therefore, early cyst development seems to be an early sign of NPH3 and pcy. It is interesting that histopathologic changes observed in NPH3 and pcy show similarities (Figure 4, d and e), consisting of predominant dilatation of distal tubules and collecting ducts, sporadic dilatation of Bowman’s capsule, and interstitial inflammatory infiltrates causing a chronic sclerosing tubulointerstitial nephropathy (2,15). In addition, tubular basement membrane changes with local attenuation, thickening, and multilayering, which are supposed to be characteristic findings in adolescent and juvenile NPH, can also be observed in pcy mice (2,15,18,19).

On the basis of genetic and phenotypic findings, NPH3 and murine pcy most likely are caused by recessive mutations of homologous genes. This will help to accelerate gene identification of NPH3 and pcy because the search for the responsible genes should be focused to the critical genetic region that overlaps in both diseases, which is flanked by D3S1292 and SGC3327 (TF). Considering all EST and genes localized within this interval, only EDF1 and TOPB1 show a broad expression pattern, including kidney similar to the expression pattern of NPHP1 (20), thus rendering them potential candi-
Table 2. Comparison of clinical, pathological, and genetic findings of adolescent NPH and polycystic kidney disease in mice (pcy)*

<table>
<thead>
<tr>
<th>Inheritance</th>
<th>Adolescent NPH</th>
<th>Polycystic Kidney Disease (pcy)</th>
</tr>
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<tbody>
<tr>
<td>Manifestation of renal failure</td>
<td>Autosomal recessive</td>
<td>Autosomal recessive</td>
</tr>
<tr>
<td>Range of age at renal failure</td>
<td>Late (median age, 19 yr)</td>
<td>Late (adult mice)</td>
</tr>
<tr>
<td>Renal cyst location</td>
<td>Wide range (12 to 47 yr)</td>
<td>Wide range depending on genetic background</td>
</tr>
<tr>
<td>Kidney size</td>
<td>Predominantly at the corticomedullary junction</td>
<td>In the beginning at the corticomedullary junction, later throughout the entire kidney</td>
</tr>
<tr>
<td>Extrarenal disease manifestation</td>
<td>Normal or reduced</td>
<td>Enlarged</td>
</tr>
<tr>
<td>Histology</td>
<td>Tubular basement membrane changes, tubular dilation and atrophy, sclerosing tubulointerstitial nephropathy</td>
<td>Tubular basement membrane changes, tubular dilation and atrophy, sclerosing tubulointerstitial nephropathy</td>
</tr>
</tbody>
</table>

* NPH, nephronophthisis.

date genes (http://www.ncbi.nlm.nih.gov/UniGene). If homology of NPH3 and pcy is confirmed by gene identification, this will, in addition, have clinical implications, because intervention studies in pcy mice have demonstrated beneficial effects by modification of protein intake and administration of methylprednisolone (21–24).

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**References**


Access to UpToDate on-line is available for additional clinical information at http://www.jasn.org/