Copy-number variation analysis in familial nonsyndromic congenital anomalies of the kidney and urinary tract: Evidence for the causative role of a transposable element-associated genomic rearrangement

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Abstract. Most congenital anomalies of the kidney and urinary tract (CAKUT) are sporadic, but familial occurrence has been described, suggesting a genetic contribution. Copy-number variations (CNVs) were detected in patients with CAKUT to identify possible novel genomic regions associated with CAKUT. CNVs were investigated in 7 children with CAKUT from three unrelated families using array comparative genomic hybridization: female monozygotic twins with bilateral duplex collecting system/vesicoureteral reflux (VUR)/unilateral renal hypodysplasia (URHD); two male siblings with VUR/URHD; 3 male second cousins, one with bilateral VUR/URHD, one with bilateral VUR and one with ureterovesical junction obstruction (UVJO). Five patients had a normal constitution of CNVs, one had a duplication of 0.2 Mb on the 5q-arm (5q23.3), probably unrelated to CAKUT, and one with UVJO had a 1.4 Mb deletion on the 17q-arm (17q12) which includes a known CAKUT gene, HNF1B. The phenotype of HNF1B deletion was extended including renal magnesium wasting. A higher load in TEs of the deleted region compared with the expected density in any random genomic region. Notably, the 5' breakpoint was mapped within a solo long terminal repeat (LTR) sequence. Moreover, highly similar members of solo LTR and mammalian interspersed repetitive (MIR) elements, as well as nucleotide sequence microhomology were detected at the breakpoint regions. In conclusion, the deletion detected in one patient suggests this genomic imbalance as causative for UVJO. A not very well known phenotype of HNF1B deletion resulting in both low urinary tract malformations and renal wasting of magnesium was described. The high load in TEs of the deleted region, the presence of highly similar elements, and the microhomology found at breakpoint regions may have contributed to the generation of the deletion. CNV analysis could reveal novel causative genomic regions in patients with CAKUT, and further studies in larger cohorts are needed.

Introduction

The prevalence of congenital anomalies of the kidney and urinary tract (CAKUT) is 17.7:1,000 live births (1). CAKUT constitute the major cause of chronic renal failure in childhood (2) and can occur in isolation (nonsyndromic CAKUT), or in association with other organ system malformations (2). While most cases are sporadic, a familial aggregation has been described in up to 15% of cases, pointing to a genetic contribution. In a recent study, 37 different heterozygous mutations (33 novel mutations) in 12 of 17 known CAKUT-causing genes were identified in 6.3% of 650 unrelated families with CAKUT. It thus appears that CAKUT is a genetically heterogeneous disease with diverse clinical phenotypes (3).

Submicroscopic chromosomal imbalances (deletions or duplications), known as copy-number variations (CNVs), have been used to identify novel genomic regions associated with CAKUT. CNVs were detected in 10.1% of 178 patients with CAKUT and were found to be inherited in 90% (9/10) of the families in which they were identified (4).

Transposable elements (TEs) can sculpt genome structure, with a profound contribution to genetic variation, through single nucleotide variations, CNVs (indels) or larger structural variations (5). Such TE-mediated rearrangements can be either active, as a direct result of retrotransposition events, or passive
due to their repetitive nature (6,7), causing diseases in some cases (7).

The aim of the present study was to investigate CNVs in a series of children with nonsyndromic upper and lower urinary tract anomalies, and to search for evidence of a possible causative role of a transposable element-associated genomic rearrangement. The novelty of the study is that it focused on first and second degree relatives with the same or similar CAKUT phenotype.

Materials and methods

Patients. Three unrelated families, each with at least two members diagnosed with nonsyndromic CAKUT in the Pediatric Nephrology Department of the University Hospital of Ioannina (Greece) were invited to participate in the study. The first family consisted of female monozygotic twins with CAKUT. One twin had a bilateral completely duplex collecting system, bilateral vesicoureteral reflux (VUR) grade III-IV in the lower pole, and right renal hypoplasia (RHD) (relative function 38%). The second twin had a left incompletely duplicated system (the two ureters joined just before entering the bladder), bilateral VUR grade II-III, and left RHD (relative function 38%). The second family consisted of two male siblings one of which had bilateral VUR grade III and left RHD (relative function 26%) and the second bilateral VUR grade III-IV. In the third family three males, second cousins, had CAKUT. One had bilateral VUR grade V (normal urethra) and right RHD (relative function 28%), one had bilateral VUR grade III, and the third had severe ureterovesical junction obstruction (UVJO), requiring surgical intervention. The patients of the third family were members of a cohort of gypsies with a high rate of inbreeding (first cousin marriages). All the patients had normal laboratory findings, apart from the 15-year-old patient with UVJO from the gypsy family who had persistent hypomagnesemia (serum magnesium levels: 1.02-1.17 mEq/l, normal range 1.3 -2.1 mEq/l), hypermagnesuria [fractional magnesium excretion (FeMg): 5.0-5.5%, and hypocalciuria (fractional calcium excretion: 0.18-0.2%) [in patients with hypomagnesemia, hypermagnesuria has been defined as FeMg >4 (8), and hypocalciuria as fractional calcium excretion <1%] (9)]. Serum creatinine, calcium, phosphate, albumin, parathyroid hormone, 25-hydroxyvitamin D levels, and electrolytes were within normal limits.

The CAKUT in these children were investigated and characterized by urinary tract ultrasound, voiding cystourethrography, dimercaptosuccinic acid (DMSA) scan, and technetium-99 m mercaptoacetyltriglycine (MAG3) scan, as indicated. VUR was classified into grades I-V according to the International Reflux Classification (10). RHD was diagnosed in kidneys with reduced renal length and regular outline, with or without cortical hyperechogenicity and loss of corticomedullary differentiation on ultrasound, and with persistent (for ≥6 months), general reduction in 99mTc-DMSA uptake (relative function <45%).

CNV detection. For the genetic study, peripheral venous blood was collected for genomic DNA extraction, according to the manufacturer's protocol using QIAamp DNA Mini Kit (Qiagen, Hilden, Germany), at the time of routine blood testing of the patients. DNA samples were investigated for the presence of CNVs, using array-comparative genomic hybridization (CGH). Array-CGH was performed by Cytochip ISCA array (Blue Gnome-version 1.0) with 180,000 oligos, analyzed using buildGRCh37 (hg19).

The following resources were used for the analysis of the cases: Ensemble (http://www.ensembl.org), Database of Genomic Variants (http://projects.tcag.ca/variation/), UCSC Genome Bioinformatics Site (http://genome.ucsc.edu/), and Online Mendelian Inheritance In Man (http://ncbi.nlm.nih.gov/omim).

CNVs >0.2 Mb were considered significant. CNVs that did not contain genes, or were <0.2 Mb (unless containing a gene of known pathogenic significance) were considered of normal constitution.

Bioinformatics. TE sequences coverage was measured using the in silico program RepeatMasker (http://www.repeatmasker.org/), and UCSC genome browser to extract nucleotide sequences from the human genome (GRCh38). Sequence features were analyzed and visualized in the UCSC browser. Nucleotide sequence alignments were performed using the Blast algorithm (http://blast.ncbi.nlm.nih.gov/Blast.cgi). Breakpoint regions (200 bp stretches surrounding the breakpoints) were analyzed using the Blast algorithm for the identification of sequence homology.

Results

Of the 7 study children with familial nonsyndromic CAKUT, 5 had a normal constitution of CNVs (well-established polymorphism variants).

One sibling from the second family, with bilateral VUR grade III and left RHD had a 0.2 Mb duplication on the long arm of chromosome 5 (5q-arm) at chromosomal band 5q23.3. The duplication partially disrupts the Chondroitin Sulfate Sulphate 3 (CHSY3) gene (Fig. 1A). His brother, with a similar CAKUT (bilateral VUR grade III-IV), did not harbor the same CNV.

One male teenager, from the gypsy family, with UVJO, was found to have a deletion approximately 1.4 Mb in size on the long arm of chromosome 17 (17q-arm) at chromosomal band 17q12. This deletion consists of two OMIM disease genes, the Acetyl-CoaCarboxylase-Alfa (ACACA, OMIM#200350), and the Hepatocyte Nuclear Factor-1-Beta (HNF1B, OMIM#189907) (Fig. 1B). To gain insight into the possible molecular mechanism underlying the deletion detected, we measured the load in TE sequences within the deleted region. A higher coverage (51.33%) was found than expected when inspecting any random genome region (Fig. 2A). Notably, the 5’ breakpoint was mapped in a solo long terminal repeat (LTR) sequence (MLT1U2), sharing high similarity (76%) to the solo LTR sequence MamRep1527 located at the 3’ breakpoint region. Moreover, a significant load in highly similar (70-89%) mammalian interspersed repetitive (MIR) element sequences, namely MIRb and MIRc, was found in the 20kb-region bilateral to the 5’ and 3’ breakpoints, amounting to 11.87 and 5.25%, respectively (Fig. 2B). Finally, a 47% nucleotide sequence homology was identified in a stretch of 200 bp surrounding the breakpoints, while
no microhomology was found in the breakpoint junctions (Fig. 2C and D).

CNVs were not studied in the parents of the affected children, but all the parents were investigated by urinary tract US and had no CAKUT phenotype.

Discussion

This study investigated the presence of CNVs in a series of 7 children with familial nonsyndromic CAKUT. Its novelty is the focus on first and second degree relatives with the same or similar nonsyndromic CAKUT phenotype, and on evidence for the causative role of a transposable element-associated genomic rearrangement.

CNV associated with CAKUT was detected in one of the 7 patients (14%). The number of patients investigated was very small, but the rate of CNVs was in agreement with previous studies in which CNVs were identified in 16.6% (11) and 10.1% (4) in large numbers of patients with nonsyndromic CAKUT (522 and 178 respectively). Sanna-Cherchi and colleagues (11) studied patients with congenital solitary kidney and renal hypoplasia, while Caruana and colleagues (4) included a wide range of CAKUT, finding a high incidence of CNVs in cases of multicystic dysplastic kidney (MCDK) (30%), and posterior urethral valves (24%). Similarly, Westland and colleagues (12), using CNV analysis in 80 patients with a solitary functioning kidney, found genomic imbalances in 11 of 80 (14%). Weber and colleagues (2) reported a similar CNV frequency (10%) in 30 children with syndromic CAKUT.

The 0.2 Mb duplication found in one sibling of the second family is not considered causative for his CAKUT phenotype, since it does not include any genes known to be associated with CAKUT (Fig. 1A), and his brother with the same CAKUT phenotype had a normal constitution of CNVs.

One child in the gypsy family, with UVJO, had a deletion approximately 1.4 Mb in size on the long arm of chromosome 17q12. The deletion consists of two OMIM disease genes, the Acetyl-CoA Carboxylase-Alpha (ACACA, OMIM#200350), and the Hepatocyte Nuclear Factor-1beta (HNF1B, OMIM#189907) (Fig. 1B). The same CNV including the HNF1B gene has also been reported recently by Caruana and colleagues (4) in a female infant with MCDK. The HNF1B gene has been associated with renal cyst, diabetes mellitus and genital malformation (RCAD syndrome: OMIM 137920), where a 75 bp deletion in exon 2 causes a loss-of-function mutation (13). HNF1B mutations are well described in patients with upper urinary tract malformations. Nakayama and colleagues (14) identified HNF1B mutations in 5 of 50 patients (10%) with CAKUT (3 with MCDK and 2 with RHD), where de novo heterozygous complete deletions of HNF1B were found in patients with MCDK. CNV analysis showed 1.4 Mb deletion of chromosome 7, involving the whole HNF1B gene with breakpoints in flanking segmental duplications. Thomas and colleagues (15) reported that among 50 North American Caucasian children with RHD, 4 (8%) carried mutations in HNF1B gene. In our study, a CNV involving the whole HNF1B gene was found in one patient with a lower urinary tract malformation (UVJO). Considering that the HNF1B is expressed in the ureteric bud...
derivatives (16,17), this finding is not surprising. The association of HNF1B alteration with the UVJO phenotype has not been reported as often as its association with upper urinary tract malformations. Adalat et al (18) reported a mutation in the HNF1B gene in one patient with hydrenephrosis/hydroureter. Specifically, they found heterozygous mutations in HNF1B gene in 23% of 91 cases with renal malformation. The range of phenotypes included large bright kidneys, MCDK, and hydrenephrosis/hydroureter. One patient has been reported with complete prune-belly syndrome associated with a complete HNF1B gene deletion (19).

Our patient had also persistent hypomagnesemia, hypermagnesuria and hypocaliuria. Adalat et al (18) found that 44% of the 18 HNF1B mutation carriers had hypomagnesemia. They detected highly conserved HNF1-recognition sites in FXYD2, and demonstrated HNF1B-mediated transactivation of FXYD2, a gene that can cause autosomal dominant hypomagnesemia, hypermagnesuria and hypocaliuria when mutated (20). Our findings, even though not novel, depict a phenotype of HNF1B gene alteration, which includes both lower urinary tract malformations and renal magnesium wasting. Therefore, our patient's phenotypic features both overlap and expand on...
reported cases of nonsyndromic CAKUT. Given, however, that the two other family members (cousins) with CAKUT did not have the 17q12 deletion, we suggest that either the CAKUT in this family is unrelated or that the UVJO and the VUR in the cousins are due to a separate undetected genetic defect. Considering that these patients had a highly consanguineous background, a possibly recessive cause of CAKUT cannot be excluded. It should be pointed out that the etiology of the majority of CAKUT cases remains unknown. The genomic imbalance, such as copy number variants, genomic, or de novo mutations, can explain up to one-third of all CAKUT cases. Moreover, findings from several studies suggest a contribution of epigenetic and environmental factors on the pathogenesis of CAKUT, supporting the theory of its multifactorial character (21).

Congenital diseases, such as CAKUT, have developmental origin (22), and TE and their control mechanisms regulate development (23). It is known that TEs are involved in recombination events, representing a major source of structural variation in the genomic landscape (24). Even if deletions of chromosome 17q12 spanning HNF1B gene are one of the most commonly identified mutations associated with CAKUT (21), their molecular characterization, and, especially, the coverage in TE sequences within or at the boundaries of such deletions has not been previously reported. The novelty of our study was the bioinformatic analysis of a CAKUT-causative 1.4 Mb deletion on the 17q-arm based on the in silico program RepeatMasker, which effectively identifies TE sequences within any given genomic region. An additional advantage of such approach comes from the use of the alignment heuristic program cross_match for the performance of sequence comparisons, providing a high sensitivity. Our analysis revealed: (a) a high coverage in TE sequences within the 17q12-deleted region (Fig. 2A), (b) a MLT1J2 solo LTR sequence at the 5΄ breakpoint (Fig. 2B), (c) the presence of 4.6- and 2-fold higher load of MIR sequences on either side of the 5΄ and 3΄ breakpoint regions respectively, compared to their expected genomic percentages (25) (Fig. 2B), and (d) 47% nucleotide sequence microhomology in a 200 bp region surrounding the breakpoints (Fig. 2C). Based on these findings, it is tempting to suggest that TEs may have served as a substrate for the generation of the deletion. We believe that the deletion detected may have originated from the mispairing of TE sequences, possibly via non-allelic homologous recombination. Given that TEs can form non B-DNA structures promoting the formation of DNA nicks, double strand breaks or stall replication forks (26), we cannot exclude the involvement of microhomology-mediated replication-dependent recombination mechanisms, as previously reported in human Xq isochromosome formation (27).

In conclusion, among 7 children with familial nonsyndromic CAKUT, one with UVJO was found to have a significant CNV, suggesting this genomic imbalance as causative of the anomaly, since it includes a known CAKUT gene, HNF1B. The phenotype of the HNF1B deletion was extended, and included both lower urinary tract malformation and renal magnesium wasting. Moreover, we determined the topological features, in terms of nucleotide sequence identity, of the deleted genomic region. Overall, our findings provide evidence of a correlation between a TE-associated genomic rearrangement and CAKUT. Future studies will shed more light for the role of TEs as causative factors of pathogenic variants. Finally, CNV analysis could reveal novel causative genomic regions in patients with CAKUT as well as in other multifactorial diseases, and further studies in larger cohorts are needed.

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