Copy-Number Mutations on Chromosome 17q24.2-q24.3 in Congenital Generalized Hypertrichosis Terminalis with or without Gingival Hyperplasia

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Congenital generalized hypertrichosis terminalis (CGHT) is a rare condition characterized by universal excessive growth of pigmented terminal hairs and often accompanied with gingival hyperplasia. In the present study, we describe three Han Chinese families with autosomal-dominant CGHT and a sporadic case with extreme CGHT and gingival hyperplasia. We first did a genome-wide linkage scan in a large four-generation family. Our parametric multipoint linkage analysis revealed a genetic locus for CGHT on chromosome 17q24.2-q24.3. Further two-point linkage and haplotyping with microsatellite markers from the same chromosome region confirmed the genetic mapping and showed in all the families a microdeletion within the critical region that was present in all affected individuals but not in unaffected family members. We then carried out copy-number analysis with Affymetrix Genome-Wide Human SNP Array 6.0 and detected genomic microdeletions of different sizes and with different breakpoints in the three families. We validated these microdeletions by real-time quantitative PCR and confirmed their perfect cosegregation with the disease phenotype in the three families. In the sporadic case, however, we found a de novo microduplication. These copy-number variations (CNVs) shared a common genomic region in which CNV is not reported in the public database and was not detected in our 434 unrelated Han Chinese normal controls. Thus, pathogenic copy-number mutations on 17q24.2-q24.3 are responsible for CGHT with or without gingival hyperplasia. Our work identifies CGHT as a genomic disorder.

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We ascertained three Han Chinese families (SY, GD, and BJ) with autosomal-dominant CGHT and coarse face (Figures 1A–1D and 1F). Families SY, GD, and BJ have ten, nine, and three affected individuals in four, four, and two generations, respectively (Figure 1F). Sixteen affected individuals of the three families were available for physical examination. Universal overgrowth of the terminal hairs and coarse face were the common phenotypic features among the affected individuals of these families (Figures 1A–1D). Most affected individuals showed excessive growth of terminal hairs all over the body as compared to the normal hair pattern and growth in Han Chinese of the same age and sex. Hair overgrowth on the back and limbs was particularly remarkable (Figures 1A and 1C). Affected females in family SY presented with less marked hypertrichosis. All affected individuals showed bushy and dark eyebrows, a broad and flat nose (in affected adults) or a bulbous soft nose (in affected children), large ears with thick, long, and hairy lobes, and thickened lips (Figures 1B and 1D). Approximately half of the affected individuals had a relatively large head (Figure 1C). With the exception of the widely spaced teeth observed in the affected adult individuals of family GD, no affected individual had obvious dental abnormalities. All affected individuals showed normal gingival growth and mental development.

We collected blood samples from participating family members after obtaining informed consent and approval of the Peking Union Medical College Institutional Review Board. Genomic DNA was extracted by standard methods from 19 (9 affected, 6 unaffected, and 4 spouse), 11 (4 affected, 5 unaffected, and 2 spouse) and 5 (3 affected, 1 unaffected, and 1 spouse) members of families SY, GD, and BJ, respectively. DNA samples from the previously described Irish family with autosomal-dominant CGHT and coarse face were also included in linkage analysis.11 To identify candidate loci for CGHT, we first carried out a genome-wide linkage scan in the four-generation family SY (Figure 1F). DNA samples were genotyped on the Affymetrix GeneChip Human Mapping 50K Array Hind 240 containing 57,244 SNPs. Array experiments were carried out according to the protocol of the manufacturer. The
Affymetrix GeneChip Operating Software (GCOS) was used in image processing. Genotypes were called with the Affymetrix GeneChip Genotyping Analysis Software (GTYPE 4.0). Parametric multipoint linkage analysis was performed with the dChipLinkage software under the assumptions of autosomal-dominant inheritance with 99% penetrance, a disease allele frequency of 0.1%, and equal SNP allele frequency (50%). Mendelian errors and unlikely genotypes were deleted prior to linkage analysis. Our multipoint linkage analysis produced a maximum logarithm of the odds (LOD) score of 2.37 with markers on chromosome 17q24.2-q24.3 (Figure 1E). To confirm the genome-wide linkage scan and define the critical region, we then typed in the family seven microsatellite markers (D17S789, D17S795, D17S2182, D17S1786, D17S1295, D17S1534, and D17S1351) from the same chromosome region and performed two-point linkage analysis. Two-point LOD scores were calculated with the MLINK program of the LINKAGE package. The parameters used in linkage analysis were autosomal-dominant inheritance, complete penetrance, a mutation rate of zero, equal male-female recombination rate, equal microsatellite-allele frequency, and a disease-allele frequency of 0.01%. PCR primers used for genotyping are given in Table S1 available online. We obtained a maximum LOD score of 3.91 at \( q = 0 \) for two markers (D17S2182 and D17S1786) (Table S4), confirming the genetic linkage. Consistent with the genome scan, haplotype analysis in two recombinants (III-4 and IV-1 of family SY, Figure 1F) placed the critical region to an interval between markers D17S795 and D17S1351, representing a genomic extent of 4 Mb containing nine RefSeq genes. By typing the same microsatellite marker set, we detected haplotype sharing among affected members in families GD and BJ (Figure 1F). Notably, all affected individuals in the three Chinese CGHT families showed a null allele for markers D17S2182 and D17S1786 on the disease haplotypes, suggesting the possibility of a recurrent genomic microdeletion (Figure 1F). Our haplotype analysis with this microsatellite marker set excluded genetic linkage to 17q24.2-q24.3 in the Irish family. Further genome-wide linkage scan in this family yielded a maximum LOD score of 1.77 with SNP markers on a chromosome 6 region (data not shown), pointing to the presence of genetic heterogeneity in autosomal-dominant CGHT with coarse face.

To confirm whether the CGHT phenotype is associated with a recurrent microdeletion on 17q24.2-q24.3, we performed CNV analysis with the Affymetrix Genome-Wide Human SNP Array 6.0 containing more than 906,600 SNPs and more than 946,000 copy-number probes. Genomic DNA samples were genotyped with the SNP Array 6.0 in accordance with the manufacturer’s protocols. The Affymetrix Genotyping Console 3.0 software was used for genotype calling, genotyping quality control, and CNV identification. Copy-number-state calls were determined with the Canary algorithm embedded in the Affymetrix Genotyping Console 3.0 package. We analyzed one affected individual per family and detected genomic microdeletions of different sizes and with different breakpoints in all the three familial cases (Figure 2A). To validate

**Figure 2. Identification of Inherited Microdeletions on Chromosome 17q24.2-q24.3 in Three CGHT Families**

(A) Copy-number state of a 5 Mb genomic region on chromosome 17q24.2-q24.3 in three affected individuals, one per family. (B) Real-time quantitative PCR (qPCR) assays of 11 amplicons validating the microdeletions. The deletion is seen as a ~0.5-fold normalized relative copy number (RCN). Error bars represent SD. (C) Amplification of deletion junctions showing segregation of the microdeletions with the phenotypes in families SY, GD, and BJ. The sizes of long-range PCR fragments are shown. (D) Sequence analysis of the deletion junctions. The colored sequences represent the deletion junctions. The precise positions of the proximal and distal breakpoints are indicated in red and blue, respectively.
these nonrecurrent microdeletions by an alternative method, we also performed multiple qPCR assays to determine the relative copy number (RCN) of the target genomic regions. We designed 11 qPCR assays by using the Primer Express v2.0 software (Applied Biosystems). Information about the primer sequences and amplicon position is given in Table S2. The qPCR was carried out as described previously.16 The quantification of the target sequences was normalized to an assay from chromosome 21, and the RCN was determined on the basis of the comparative ΔΔCt method with a normal control DNA as the calibrator.16 The experiments were repeated three times. A ~0.5-fold RCN and a ~1.5-fold RCN were used for deletion and duplication, respectively. We detected an RCN of 0.5 for the corresponding qPCR targets in all analyzed individuals and thus provided independent confirmation of all the three microdeletions (Figure 2B).

To amplify the deletion junctions, we designed long-range PCR primers from the sequences flanking the breakpoints on the basis of the SNP Array 6.0 genomic coordinates. Information about the primer sequences is given in Table S3. With use of the long-range PCR assays, we detected the corresponding microdeletions in all affected individuals of the three study families, but not in the unaffected family members (Figure 2C). Sequence analysis of the PCR fragments revealed the breakpoint positions and defined the precise sizes of the three nonrecurrent microdeletions to be 964,182 bp, 555,432 bp, and 677,661 bp in affected cases of families SY, GD, and BJ, respectively (Figure 2D). For affected cases of families SY, GD, and BJ, the proximal and distal breakpoints were located in unique and L2 repeat, L1PA16 repeat and L1MC4 repeat, respectively (Figure 2D). Microhomology between the proximal and distal breakpoints was not present (Figure 2D). Thus, the microdeletions in our three CGHT families most likely occurred through a nonhomologous end-joining mechanism.

We also recruited a 31-year-old Chinese man (KK) with extreme CGHT, coarse face, and gingival hyperplasia (Figure 3A), a phenotype closely resembling that of Julia Pastrana. KK was recorded as one of the world’s hairiest men and has 96% of his body covered with thick, jet-black, and long (>5 cm) terminal hairs. He has a large head completely out of proportion to the rest of his body (Figure 3A). Oral examination revealed an abnormal dental occlusion and a delayed tooth eruption. He had surgical treatment for deformed nose and hyperplastic gingivae. His parents were apparently normal. We extracted genomic DNA from KK and his parents. Our high-resolution CNV screening with the SNP Array 6.0 revealed a genomic microduplication on chromosome 17q24.2-q24.3 that is 1,429,960 bp in size (CN_761503 to SNP_A-8412453) (Figure 3B). This microduplication was only present in KK and not in his parents (Figure 3B), indicating that the rearrangement is de novo in origin. We also validated the microduplication with the multiple qPCR assays (Figure 3C) and confirmed its de novo nature by an assay of the amplicon 7 (Figure 3D).

We used two-color interphase fluorescence in situ hybridization (FISH) to determine the orientation of the microduplication found in KK. The bacterial artificial chromosome (BAC) clones CTD-2331H1, RP11-648L7, and CTD-2309E11 were selected to cover the some 17q24.2-q24.3 in KK (A) Photograph of KK showing extreme hair overgrowth. (B) Copy-number state of a 5 Mb genomic region on chromosome 17q24.2-q24.3 showing the presence of a microduplication in KK (green) but not in his parents (blue and red). (C) Eleven qPCR assays validating the microduplication in KK. The duplication is seen as a ~1.5-fold RCN. Error bars represent SD. (D) The qPCR assay of the amplicon 7 confirming the de novo nature of the microduplication in KK. Error bars represent SD. (E) Two-color FISH of a representative interphase nucleus showing a “one fused red - one normal green” signal pattern. BAC probes are CTD-2331H1 (red), the most proximal fully duplicated BAC clone, and CTD-2309E11 (green). (F) Two-color FISH of a representative interphase nucleus showing a green-fused red-green signal order. BAC probes are CTD-2331H1 (red) and RP11-648L7 (green), the central duplicated BAC clone.
and the central duplicated BAC clone RP11-648L7 (green), 69 of the 106 counted interphase nuclei revealed a green-fused red-green signal order (Figure 3F). These two-color FISH signal patterns are consistent with an inverted duplication (Figure 4).

The four CNVs identified in our familial and sporadic CGHT cases shared a common overlapping genomic region that contains four genes, \( \text{ABCA5} \) (MIM 612503), \( \text{ABCA6} \) (MIM 612504), \( \text{ABCA10} \) (MIM 612508), and \( \text{MAP2K6} \) (Figure 4). CNV of this genomic region was not detected in 434 unrelated Han Chinese normal controls. We isolated human hair follicles from a scalp skin tissue donated by a healthy individual after a cosmetic surgery and extracted total RNA. With use of RT-PCR, we could detect mRNA expression of all four genes (Figure S1). Information about the primer sequences is given in Table S5.

The CGHT-coarse face-gingival hyperplasia syndrome was first reported in the years before the discovery of Mendel’s laws of heredity.\(^8,9\) However, not much has changed since then and the rare condition is still on the list of Mendelian phenotypes for which the underlying molecular basis remains unknown. We have identified a genetic locus for CGHT and found four overlapping 17q24.2-q24.3 CNVs (Figure 4), including three nonrecurrent microdeletions that show perfect cosegregation with the disease phenotype in three Chinese families with CGHT and coarse face, and one de novo inverted microduplication in a sporadic case with extreme CGHT, coarse face, and gingival hyperplasia. CNV of the common minimal region delineated by these four genomic rearrangements is not reported in the public Database of Genomic Variants and was not detected in our unrelated normal controls. Together, our results strongly support a causal relationship between the CNVs on 17q24.2-q24.3 and the CGHT with coarse face or with coarse face and gingival hyperplasia. Our finding of the copy-number mutations on the same chromosome region also suggested that the autosomal-dominant CGHT with coarse face could represent a variant of the CGHT-coarse face-gingival hyperplasia syndrome. Our work has established this rare syndrome as a genomic disorder.

The minimal disease genomic region of the four copy-number mutations contains three genes of the ATP-binding cassette transporter subfamily A, \( \text{ABCA5} \), \( \text{ABCA6} \), and \( \text{ABCA10} \), which functions to maintain lipid homeostasis.\(^17\) In a large-scale survey of CNVs in normal Han Chinese population, Lin and colleagues found that 4 of the 300 individuals had a 203 kb deletion affecting these three genes.\(^18\) These make \( \text{MAP2K6} \), the gene producing the human mitogen-activated protein kinase kinase 6, the leading candidate from the overlapping region. Activating mutations in components of the RAS-MAPK pathway can result in a group of phenotypically overlapping genetic syndromes with several dysmorphic features common to the patients reported herein.\(^19\) Costello syndrome (MIM 218040), for example, is caused by mutations in \( \text{HRAS} \) (MIM 190020), an important gene involved in the RAS-MAPK pathway, and often shows coarse face with thick lips, large-lobed ears, gingival hypertrophy, and relative macrocephaly.\(^19,20\) Moreover, it has been demonstrated that a gain-of-function mutation in \( \text{SOS1} \) (MIM 182530) activates the RAS-MAPK pathway and thus cause gingival overgrowth in hereditary gingival fibromatosis (MIM 135300).\(^21,22\) Notably, a hallmark of the cardiofaciocutaneous syndrome (MIM 115150) caused by mutations in \( \text{BRAF} \) (MIM 164757), \( \text{KRAS} \) (MIM 190070), \( \text{MAP2K1} \) (MIM 176872), or \( \text{MAP2K2} \) (MIM 601263) is sparse and curly hair but not the thick and long hair.

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**Figure 4. Schematic Diagram of the 17q24.2-q24.3 Region with a Summary of the Copy-Number Mutations Identified in the Present Study**

Positions of the RefSeq genes, microsatellite markers, BAC clones, and qPCR amplicons (vertical arrows 1–11) are shown. Open bars represent the sizes of the three microdeletions in affected individuals of families SY, GD, and BJ. Solid bars display the inverted microduplication in KK. Color bars indicate the positions and sizes of the BAC clones used in the two-color interphase FISH.
seen in CGHT.\(^23\) In a 12-year-old girl with a de novo microdeletion of 17q24.2-q24.3 encompassing MAP2K6, neither CGH nor coarse face was observed.\(^24\) Additionally, knockout mice for Map2k6 lack abnormal hair growth.\(^25,26\) These facts might argue against the phenotypic contribution of the MAP2K6 gene in our three CGHT families.

Altered expression of the dosage-sensitive genes caused by copy-number mutations has traditionally been viewed as the general pathogenic mechanism for genomic disorders.\(^13,14\) Recently, it has been shown that some copy-number mutations can have a position effect, by disrupting regulatory elements or changing local chromatin structure, on expression of the key developmental regulator genes flanking the deleted or duplicated regions.\(^14\) Congenital hypertrichosis has been documented in several cases with chromosome 17q trisomy, suggesting that hair overgrowth can result from increased function of the gene(s) on chromosome 17q.\(^27\) Of note are two studies that have identified Sox9, a SRY-related HMG-box transcription factor, as an essential regulator for hair follicle stem cells.\(^28,29\) Mouse skin inactivated for the Sox9 gene develops alopecia.\(^28\) Conceivably, enhanced expression of Sox9 might lead to hair overgrowth. The above-mentioned microdeletion in the girl without CGH was \(-2.3\) Mb (chr17:63,260,444-65,593,531), containing 18 RefSeq genes, and had the distal breakpoint within KCNJ16 (MIM 605722).\(^24\) Moreover, a reciprocal translocation t(3;17)(p14.3;q24.3) has been reported in a sporadic case with Zimmermann-Laband syndrome (MIM 135500) presenting with hypertrichosis, coarse face, and gingival hyperplasia.\(^30\) The breakpoint at 17q24.3 was defined at the telomeric side of KCNJ2 (MIM 600681).\(^31\) The human SOX9 locus is located 2,578,690 bp and 1,940,977 bp apart from the MAP2K6 and KCNJ2 loci, respectively (Figure 4). Therefore, the copy-number mutations described in the present study might exert a long-range position effect by altering the local chromatin architecture, resulting in an increased expression of SOX9 in hair follicle stem cells and thus causing a universal hair overgrowth. Constitutive activation of MAP2K6 in mouse condrocytes has been shown to increase the transcriptional activity of Sox9.\(^32\) This observation might provide a molecular clue to understanding the pathogenesis of the extreme hypertrichosis in KK. Clearly, further studies are needed to elucidate the molecular mechanisms by which the copy-number mutations on 17q24.2-q24.3 produce the phenotypes.

Supplemental Data

Supplemental Data include five tables and one figure and can be found with this article online at http://www.ajhg.org/.

Acknowledgments

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Web Resources

The URLs for data presented herein are as follows:

Database of Genomic Variants, http://projects.tcag.ca/variation
UCSC Genome Browser, http://genome.ucsc.edu/cgi-bin/hgGateway

References


