CLINICAL STUDY

Improvement of molecular-genetic diagnostics of the most common skeletal dysplasias

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ABSTRACT
OBJECTIVES: The aim of this study was to take the molecular-genetic methods for detection of the most frequent mutations in patients suspected for achondroplasia (ACH) and hypochondroplasia (HCH) into the routine practice.

BACKGROUND: Both disorders are usually caused by de novo gain-of-function type mutations in FGFR3 gene encoding the fibroblast growth factor receptor 3, which plays an important role in the metabolism of connective tissues. More than 99 % of ACH cases are caused by the glycine-to-arginine substitution at codon 380 and about 70 % of HCH cases result from the asparagine-to-lysine/-serine/-threonine substitutions at codon 540 in the consequence of the four different possible nucleotide changes occurred at the same codon.

METHODS: Exons 10 and 13 of the FGFR3 gene were analysed by PCR-RFLP and sequencing analysis. The exon 13 sequencing was necessary for mutation type specification.

RESULTS: We confirmed the diagnosis of ACH due to 1138G→A transition in 7 patients and we identified 1620C→A transversion responsible for HCH in 2 patients.

CONCLUSION: Due to serious limitations in recently used methods, we had to modify the molecular-genetic diagnostics approach. We developed the reliable diagnostics and made it available for achondroplasia and hypochondroplasia suspected patients (Tab. 1, Ref. 5, Ref. 17). Text in PDF www.elis.sk.

KEY WORDS: achondroplasia, hypochondroplasia, FGFR3 gene, 1138 G-to-A transition, 1620 C-to-A transversion.

Introduction

Achondroplasia (ACH, MIM 100800) and Hypochondroplasia (HCH, MIM 146000) are the common types of skeletal dysplasias with the prevalence between 1 : 30 000 and 1 : 10 000 predominantly characterised by short stature and dwarfism. They are inherited in an autosomal-dominant manner. Clinical features of achondroplasia are associated with enlarged head, depressed nasal bridge, midfacial hypoplasia, proximal shortening of the extremities, short stubby hands, genu varum, limitation of elbow extension, megalecephaly, lordotic lumbar spine, prominent buttocks, protuberant abdomen and final adult height of approximately 125 cm for males and 120 cm for females (1, 2). Hypochondroplasia is characterised by clinical manifestations milder than in ACH with the average height of adults – 146 cm in males and 138 cm in females, macrocephaly, limitation of elbow extension, varus deformity of the lower extremities, lumbar lordosis, narrowing of the spinal canal, seldom spinal stenosis and mild to moderate mental retardation (3). Both disorders result from recurrent nonsense mutations in FGFR3 gene, which spans over 18 exons and lies within the 4p16.3 region (4). Gene encodes the member of the family of tyrosine kinase (TK) transmembrane receptors (FGFR1, FGFR2, FGFR3 and FGFR4). They bind fibroblast growth factors with low capacity and high affinity. Mutations cause receptor activation and inhibition of chondrocyte proliferation, what leads to abnormalities of epiphyseal plate function and bone tissues growth.

FGFR3 receptor is highly homologous to other members of TK family and consists of three extracellular immunoglobulin-like FGF-binding domains (Ig I–III), one hydrophobic transmembrane (TM) domain and two cytoplasmic tyrosine kinase sub-domains TK1 and TK2 responsible for the catalytic activity. Typical features for FGFRs are the unique patterns of expression during embryogenesis, suggesting their different functions of FGFs mediating (5).

More than 99 % of achondroplasia is caused by the heterozygous glycine-to-arginine substitution at codon 380 in the transmembrane domain. About 98 % of this substitution cases are based on the G-to-A transition and about 1 % of cases on the G-to-C transversion at nucleotide 1138 (6). All new mutations occur on the paternal allele, suggesting an increased mutability of FGFR3 during spermatogenesis and are associated with an increased paternal age (7). Homozygosity or compound heterozygosity states are often manifest with a more severe phenotype and are lethal in the perinatal period (8).

The most common mutation detected in about 65 % cases of hypochondroplasia is the N540K amino-acid substitution in the TK1 domain that can lead to the disruption of the α-helical structure
Molecular-genetic diagnostics was not available in Slovakia. All samples were sent to our Institute in the last three years. Before the year 2011, HCH was based on clinical and radiological examinations. Blood samples of patients with suspected diagnoses of ACH and HCH were sent.

**Material and methods**

Our study involved patients from all regions of Slovakia. Our sample included patients with syndromes compatible with achondroplasia, with different amino-acid substitutions that arise at the same codon (Fig. 1). At present, it is evident that HCH can be caused also by other substitutions like N328I, I538V, K650N/Q, but many of them are private mutations only with single family recurrence (12, 13, 14).

Though molecular-genetic and biochemical base of ACH and HCH is known for more than 20 years, no effective therapy for bone growth correction can be offered to patients. Nowadays, some new strategies for therapy are in development. Both disorders result from an increased signal transduction from the mutant receptor. Therefore, therapeutic strategies are focused on reducing these signals. The first tested therapeutics, like kinase inhibitors, directly targeting FGFR3 are in the preclinical phase of testing. For example binding of C-type natriuretic peptide to its receptor, natriuretic-peptide receptor B (NPR-B), inhibits signal transduction pathways downstream of FGFR3 (15, 16).

**Patients**

Our study involved patients from all regions of Slovakia. Blood samples of patients with suspected diagnoses of ACH and HCH based on clinical and radiological examinations were sent to our Institute in period of last three years. Before the year 2011, molecular-genetic diagnostics was not available in Slovakia. All subjects signed an informed written consent. DNA was extracted from peripheral blood cells or from amniotic fluid cells using MN NucleoSpin Blood Mini according to the manufacturer’s protocol (Macherey-Nagel).

**Mutation analyses**

Genomic DNA was amplified using primer sets common for both methods, for PCR-RFLP and sequencing analyses. Primers flanking the intron-exon boundaries were designed using Primer 3 software v. 0.4.0 and checked by SNPCheck v3 and NCBI/Blast softwares. PCR amplification of FGFR3 exons 10 and 13 was performed under the following conditions. Exon 10 was amplified using 5X Phusion® GC buffer (Thermo Fisher Scientific), 0.2 mM dNTPs, 0.75 M Betaine, 0.5 μM of each primer and 0.02 U/μl of Phusion® High-Fidelity DNA polymerase in a reaction volume of 20 μl. Exon 13 was amplified using 2X PCR Master Mix (Thermo Fisher Scientific), 3% DMSO and 0.3 μM of each primer in a reaction volume of 25 μl. Primer sequences, amplification conditions for Mastercycler EPPGradient (Eppendorf®) and product lengths were summarised in Table 1.

In accordance with the known facts that mutations create novel restriction sites, screening for G380R mutations in exon 10 was performed by digestion with 3U of restriction enzymes – SfiI and MspI (1). 1138G>A transition creates the novel restriction site for SfiI (recognition sequence 5’-CTGAC-3’) and 1138G>C transversion for MspI (recognition sequence 5’-CGG-3’). The digested products were separated on the Spreadex® EL 800 Mini gel (Elchrom Scientific) with working separation range of 60–800bp and optimal range of 200–500bp under conditions: 110V on 20 °C for 2 hours and 45 minutes. Visualisation was made by solution of GelRed™ Nucleic Acid Gel Stain and NaCl.

We found that all mutations in codon 540 abolish a restriction site for enzyme, which recognizes non-mutated standard sequence. PCR product of exon 13 was at first digested with 4U of BstI (recognition sequence 5’-ACCTGC(N)4-3’) to distinguish between pathological and normal allele. The digested products were separated on a 2% agarose gel. BstI is an isoschizomer of BspMI and cleaves target sequences more efficiently. It belongs to a group of enzymes, which requires two copies of its recognition sequence for effective cleavage and furthermore needs to interact with the DNA >6 bp upstream and >10 bp downstream of the recognition site. Therefore, after the first mutation screening by PCR-RFLP, it was necessary to make a sequencing analysis. For verification of detected G380R mutations and for identification of 540 codon mutations, a sequencing analysis has to be performed. PCR products were enzymatically purified using thermostable alkaline phosphatase FastAP™ and Exonuclease I (Thermo Fisher Scientific) and directly sequenced. Sequencing analysis was

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**Table 1. Primers for amplification and sequencing of genomic DNA and conditions for PCR amplification.**

<table>
<thead>
<tr>
<th>Primer location</th>
<th>Primer sequence 5′→3′</th>
<th>Amplification conditions</th>
<th>Product size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FGFR3ex10F</td>
<td>CAGGCCAGGCCCCTCACGCCC</td>
<td>98 °C x 1 min [98 °C x 10 s; 72 °C x 30 s] x 27 cycles</td>
<td>908bp</td>
</tr>
<tr>
<td>FGFR3ex10R</td>
<td>GACGGAAGAGTGTCACCACA</td>
<td>92 °C x 7 min</td>
<td></td>
</tr>
<tr>
<td>FGFR3ex13F</td>
<td>CCGTGAAGATGCTGAAGGCT</td>
<td>95 ± °C x 4 min [95 °C x 30 s; 59 °C x 30 s; 72 °C x 30 s] x 30 cycles</td>
<td>395bp</td>
</tr>
<tr>
<td>FGFR3ex13R</td>
<td>CTTGCAGGTGTCAAGGAGGCT</td>
<td>72 °C x 7 min</td>
<td></td>
</tr>
</tbody>
</table>
performed with ABI 3100 Genetic Analyser using BigDye® Terminator v3.1 Cycle Sequencing kit, following the manufacturer’s instructions. Data were analysed by Chromas 2.2 (Technelysium Pty Ltd., Australia) and Vector NTI 11.5 (Informax) and compared to reference sequence from GenBank (NM_000142.4).

**Results**

In the period of three years, we confirmed the diagnosis of ACH and HCH in 9 patients from 25 requested probands. 7 patients carried the G380R mutation in the position 1138G>A transition – leading to ACH phenotype and 2 carried the N540K mutation in the position 1620C>A transversion – leading to HCH phenotype.

**Mutation analyses for ACH**

Previous published methods for molecular-genetic diagnostics (17) was based on amplification of very short fragment (164bp) containing only one restriction site located in hot spot mutation codon. For the judgement of a digestion effectiveness it is recommended to set other digestion control sites except the site of interest. Therefore, we set up amplification of larger product to avoid the false-negative results. The 908bp PCR product is subsequently digested by two restriction enzymes. After SfiI digestion resulted in fragments of 322, 378, 208bp in wild-types and in fragments of 91, 231, 378, 208bp if the mutation 1138G>A is present. If the novel restriction site is create by the G-to-C transversion, the fragment sizes after MspI digestion will be 345, 206, 160, 77bp in wild type and 95, 370, 206, 160, 77bp in case of mutation presence (Fig. 2).

Sequencing of FGFR3 exon 10 was applied in aim to verify the RFLP analysis findings (Fig. 3).
Mutation analyses for HCH

In two patients, the restriction analysis of the 395bp amplified PCR product resulted in abolition of the digestion site for BfiI, demonstrating the presence of a closely unspecified mutation in codon 540 in heterozygous state. The fragment sizes after the digestion were 194, 146, 55bp in wild-type and 340, 55bp in case of mutation presence, due to the restriction site abolishment (Fig. 4).

Sequencing analysis had to be done in all cases, in which the 540 codon closely unspecified mutation was identified by PCR-RFLP to determine the mutation type (Fig. 5).

Discussion

Recently, it was supposed that there are only few causal mutations involved in phenotypes of achondroplasia and hypochondroplasia. According to present data, the N540K substitution should be a cause of HCH in about 65% of patients (8). For ACH, the assumption was much more rigid, because it was known that as much as 98% of positive cases carried the heterozygous amino-acid substitution of G380R. It indicates that this region is the most sensitive area for germline mutation in the entire FGFR3 gene (6). To date, more new causal mutations were identified. Only two new mutations have been found to be responsible for ACH and both of them lied within the transmembrane domain, as so as G380R. Seventeen new mutations have been found to lead to HCH (2). However, according to present knowledge, codons 380 and 540 are the most important mutation hot-spots.

We combined two methods PCR-RFLPs and sequencing analysis for diagnostics purposes.

We consider the sequencing analysis as the only one possible and reliable way to distinguish between mutations types that can occurred within the 540 codon in exon 13 of the FGFR3 gene. Despite the restriction enzyme advantage, PCR-RFLP method is in that case highly uninformative. Our findings obtained using these techniques are in agreement with previous reports. We identified 1620C>A transition in two patients with typical hypochondroplasia phenotype.

Every our patient with a confirmed diagnosis of achondroplasia had the same mutation type – 1138G>A transition, which is the single most important nucleotide in the human genome. This most frequent FGFR3 substitution (G380R) induces constitutive activation of the receptor producing by ligand binding. In consequence to this molecular phenomenon, a prolonged signalling at the cell surface is observed (1). All ACH patients had typical clinical findings with predomination of rhizomelic limbs shortening, macrocephaly and facial dysmorphism. We confirmed the supposed clinical diagnosis at early childhood period. Four of our patients were actually newborns, 2 were nurshing and the oldest one was at the age of 4 and this pointed to the importance of the prompt and early determination of diagnosis.

Unlike, clinical diagnosis is often not made during infancy and many affected children are present to specialists for failure to grow more later in childhood. Prompt and definitive diagnosis of both disorders could be made also prenatal and very early postnatal and will have important implications for prognosis, clinical management and genetic counselling of other family members. The combination of two molecular genetic methods allows a reliable diagnosis and becomes available for patients from Slovakia.

References


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