The First Iranian Case of Mucopolysaccharidosis IIIC: Use of Homozygosity Mapping in a Consanguineous Pedigree

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Mucopolysaccharidosis type IIIC (MPSIIIC) is a rare subtype of mucopolysaccharidosis disorder family caused by mutations in heparan-alpha-glucosaminide N-acetyltransferase (HGSNAT) gene. MPSIIIC is subdivided into four subtypes which have overlapping features, and are indistinguishable at clinical level. In populations with high consanguineous marriage rate, homozygosity mapping can be a good choice for finding a disease locus. Herein we report a female patient with a novel mutation in HGSNAT gene in Iranian population. Clinical diagnosis was accomplished based on clinical manifestations, and urine biochemical analysis. Homozygosity mapping was performed using SNP-array technology to narrow down the candidate locus. All coding exons of HGSNAT were scanned by direct DNA sequencing. We found a novel ins/del mutation as c.1357TA>C. This mutation is a frameshift which eventually leads to premature protein truncation. To the best of our knowledge this is the first case report of Sanfilippo type C in Iranian population. This result also supports the applicability of homozygosity mapping to the diagnosis of Sanfilippo subtype.

Keywords: Mucopolysaccharidosis type IIIC, HGSNAT, homozygosity mapping, SNP-array, mutation detection
shaped vertebral bodies are common symptoms in MPSIII subtypes.

Subtypes of MPSIII have different prevalence. IIIA and IIIB subtypes are the most common types, while subtypes IIIC and IIID are less prevalent (7).

MPS IIIA, also called Sanfilippo syndrome type A, is caused by deficiency of the heparan N-sulfatase enzyme. MPS IIIB or Sanfilippo syndrome type B, is due to a deficiency in α-N-acetylglucosaminidase. Both genes associated with MPS IIIA and MPS IIIB are located on the long arm of chromosome 17, 17q25 and 17q21, respectively.

Deficiency of N-acetylglucosamin 6-sulfatase causes MPS IIID. The gene that causes MPS IIID is located on the long arm of chromosome 12q14 (1).

The most rare form of MPSIII, MPS IIIC, is caused by deficiency in heparan acetyl-coA: alpha-glucosaminide N-acetyltransferase enzyme (HGSNAT), and breaks down a glycosaminoglycan (GAG) called heparan sulfate (7). The corresponding gene is located on chromosome 8p11.1.

The primary laboratory diagnosis of the MPS III relies on increased concentrations of GAG heparan sulfate in the urine. Specific enzyme assay on fibroblast or leukocyte sample is a reliable method for precise differential diagnosis of MPS III subtypes. However, this method is labor intensive and less available (1, 2).

Mutation detection in these genes is the most reliable diagnostic tool for detection of MPSIII subtypes. However, the large size and large number of coding exons make molecular diagnosis of MPSIII subtypes time and cost consuming.

Homozygosity mapping can be used for the tracking of mutated locus in consanguineous families.

Moreover, molecular detection of disease causing mutations is required for definite diagnosis, and also prenatal diagnosis.

In the present study, we report an Iranian case of Sanfilippo type IIIC diagnosed by genetic mapping and whole gene sequencing.

Case presentation
Clinical findings

The patient was a 6 years old girl, first child of family, suffering from aggressiveness, hyperactivity, and poor attention. The parents were healthy consanguineous (first cousins).

She was born at term with normal vaginal delivery, and good Apgar. Her height was 118 cm and head circumference was 52 cm at the time of evaluation.

She had a surgery for umbilical hernia. On physical examination, mild coarse facial features, nasal bridge repression, and filled eyebrows were prominent.

She had speech delay, hyperactivity, poor attention, and autistic behavior which became worth during the last two years.

In radiologic evaluation, lumbar spine showed ovoid shaped vertebral bodies (fig-1a). Both hands suffered from widening of metacarpals (fig-1b). Furthermore, brain MRI revealed mild ventriculomegaly, and cortical atrophy (fig-1c). Serum amino acid profile, metabolic screening results (performed by MS/MS), and urine organic acid profile were in normal range. Thin layer chromatography of urine showed elevated heparan sulfate levels which suggested MPSIII.

Homozygosity mapping using SNP array

In order to localize the chromosomal location of the pathogenic variant, we genotyped samples from affected child and his unaffected sibling genome-wide by using Illumina Human CytoSNP-12 V2.1 bead chip array incorporating ∼330,000 genetic markers. Homozygosity mapping revealed the presence of a notable homozygous region in chr8p12-p21.11. This homozygosity box has 44.5 Mb length and harbors the most uncommon form of MPSIII (Sanfilippo syndrome) gene, HGSNAT (fig-2). Unaffected sibling did not show a run of homozygosity in this region.

DNA sequencing

To detect the responsible mutation in this
patient, all of HGSNAT exons were amplified by PCR. To detect the responsible mutation in this patient, all of HGSNAT exons were amplified by PCR. Sequences of used primers are available on request. All of PCR products were sequenced bidirectionally. Finally, c.1357TA>C was detected in homozygous state (fig-2). Patient’s parents were heterozygous for this mutation. This mutation was a frameshift mutation which causes reading frame change, and eventually leads to premature stop codon. This mRNA may be subject to ‘non-sense mediated mRNA decay”, and total loss of HGSNAT function.

**Discussion**

Sanfilippo syndrome type C is caused by acetyl coA-α-glucosaminide N-acetyl transferase deficiency. This enzyme is involved in glycosaminoglycan heparan sulphate degradation which its accumulation leads to cell cytotoxicity as the basis of disease pathophysiology.

The frequency of MPSIIIC varies among MPS patients population around the world. MPS IIIC accounts for approximately 18% of Korean patients with MPS, and 12.4% of the total MPS III in the Spanish population (8-12).

Most MPS III type C patients start to manifest disease symptoms in infancy or early childhood. Rarely some patients with MPS IIIC show adulthood onset of clinical symptoms (13). The genetic defect underlying MPS IIIC was identified only 5 years ago. Thus, a few documents are available about MPS IIIC syndrome (14-16).
To date, there is no any therapeutic approach for MPS IIIC. Medical interventions are limited to symptomatic medication such as flavonoid genistein, which decreases the pathological accumulation of GAGs. Furthermore, rhodamine B, an effective inhibitor of GAGs production in vitro and in vivo, may also be used to treat MPSIII subtypes (6).

Treatments based on deficient enzyme replacement do not show promising results for MPSIII disorders. This failure is due to inability of the enzyme to pass through the blood-brain barrier (2).

Recently, Clinical trials using direct injection of purified enzyme or viral vectors harboring enzyme coding sequence into the intracranial space showed promising positive results. The use of recombinant viruses that infect the central nervous system can be an attractive therapeutic option in the gene therapy studies for MPS IIIC in the near future (17-19).

Approximately 54 mutations have been reported in HGSNAT gene so far. These include 18 missense mutations, 13 splicing site mutations, 11 insertion and deletion mutations, 8 nonsense mutations and 4 polymorphisms (3, 20).

All MPSIII subtypes are autosomal recessive disorders, and therefore both enzyme coding alleles must be mutated to create the disease. Considering the low frequency of these disorders, most of MPSIII patients are expected to be result of consanguineous marriage. So most of these patients have identical by descent (IBD) disease alleles.

With that in mind, homozygosity mapping methods can be a promising approach to narrow down MPSIII subtype diagnosis. In next step, direct sequencing can be used for confirming diagnosis.

Here, we report a Sanfilippo type C case which was diagnosed using this strategy. A novel mutation designated as c.1357TA>C in coding sequence of HGSNAT gene was detected in this patient. This mutation results in frame shift, and produces a nonsense mutation residue downstream of mutation site which eventually leads to premature protein truncation and probably nonsense mediated mRNA decay. Therefore, the C.1357TA>C mutation is thought to be a disease causing mutation.

Patient’s parents were first cousin relatives, so it seems that the mutation identified in this family originated from a common ancestor. To the best of our knowledge, this is the first report of MPS III in Iranian population.

Conflict of interest
The authors declare that they have no competing interest.

References


