Mutation Screening of ENAM, KLK4, MMP20 and FAM83H Genes among the Members of Five Iranian Families Affected with Autosomal Recessive Hypoplastic Amelogenesis Imperfecta

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Abstract

Amelogenesis Imperfectas (AIs) are clinically and genetically heterogeneous conditions characterized by a wide range of clinical features. These abnormalities of enamel formation are categorized into three main groups, hypoplastic, hypomaturatation and hypocalcified with different modes of inheritance such as autosomal recessive (AR), autosomal dominant (AD) and X-lined recessive (XLR). In spite of the fact that frequent studies have explained the histological features of AIs, our knowledge regarding the molecular etiology of the affected enamel is not adequate. Up to now, different loci have been suggested to associate with the causation of AIs. Several genetic mutations including enamelin (ENAM), amelogenin (AMELX), ameloblastin (AMBN), tuftelin (TUFT1), kallikrein 4 (KLK4), matrix metalloproteinase 20 (MMP20) and family with sequence similarity 83, member H (FAM83H) have been suggested to play critical roles in the pathogenesis of these disorders. Therefore, the aim of this investigation was to study of mutation screening in ENAM, KLK4, MMP20 and FAM83H genes, responsible for AIs development in five Iranian families in which the probands were diagnosed with autosomal recessive hypoplastic amelogenesis imperfecta (ARHAI). Genomic DNA was extracted from probands and exon/intron boundaries of afore-mentioned genes were amplified by PCR and subjected to direct sequencing. We could not detect any mutation in the studied genes however; two different novel polymorphisms including T18515C and G18504A were identified in the intron 4 of MMP20 in the probands of two families. Our findings support the notion that different genes may be involved in the development of amelogenesis imperfectas.

Keywords: Amelogenesis Imperfecta; ENAM; KLK4; MMP20; FAM83H

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Introduction

Amelogenesis imperfectas (AIs) represent a group of genetic disorders affecting tooth enamel formation in both deciduous and permanent dentition. According to the clinical appearance of the enamel and molecular studies, AIs show both clinical and genetic heterogeneity. The amelogenesis imperfecta enamel defects can be classified into three main groups: hypoplastic, hypomutation and hypocalcified. It is also subdivided into fourteen subtypes, based on the phenotypes and patterns of inheritance[1]. Three different Mendelian mode of inheritance including autosomal recessive, autosomal dominant and X-linked forms have been reported. The molecular basis responsible for AIs still is not well clarified; although more than 300 genes have been documented to be associated with tooth development [2]. Several investigations have been demonstrated that enamel formation is controlled by different biomolecules including enamelin (ENAM), dentine sialophospho-protein (DSP), amelogenin (AMELX), ameloblastin (AMBN), tuftelin (TUFTI), and different enzymes such as kallikrein 4 (12KLK4) and matrix metalloproteinase 20 (MMP20) [1,3].

Several mutation studies on ENAM, KLK4, MMP20 and FAM83H have been revealed the importance of these genes in the etiology of amelogenesis imperfectas [4]. The FAM83H gene contains 5 exons and is located on 8q24.3. It has been suggested that the role of FAM83H gene is in differentiation of preameloblasts into functional ameloblasts and also in calcification of enamel matrix [5]. So far, nine mutations in FAM83H have been reported to be involved in the pathogenesis of ADHCAI [6-10].

The KLK4 gene encodes a serine protease that associated with terminal extracellular degradation of matrix proteins during the maturation stage of enamel formation [11]. This enzyme degrades different types of proteins and is almost responsible for the complete elimination of enamel matrix proteins and biomineralization of enamel [12, 13]. Hart et al. reported a mutation (g.2142G>A) in KLK4 gene and they suggested that this genetic change lead to a truncated protein and could play in the pathogenesis of AIs. [14].

The MMP20 consists of 7 exons and is located on 11q22.3-q23. This gene encodes a proteolytic enzyme that has an important role in removal of the protein components and formation of fully mineralized mature enamel. The dysfunction of this gene causes autosomal recessive amelogenesis imperfecta [4, 15]. Kim et al. have reported homozygosity for a splice site mutation (g.30 561A→T; c.954-2A→T or IVS6-2A→T) in the MMP20 gene in two affected members of a family [16].

ENAM mutation causes soft and local hypoplastic amelogenesis imperfecta[17-21] and both autosomal dominant and recessive inheritance patterns have been identified in ENAM mutation[22-24]. Rajpar et al. reported a G-to-A transition (g.6395G > A) in the splice donor site in ENAM gene[18]. Mardha et al. identified a nonsense mutation(g.2382A > T) leading to lys53-to-ter (K53X) substitution[20]. Another study reported a g.8344 of G in exon 9 in a Japanese family that was related to hypoplastic type [19].

In this study, the affected members of five Iranian families, which were diagnosed with ARHPAI, were evaluated for mutations in the ENAM; KLK4; MMP20 and FAM83H genes.

Materials and Methods

Patients

This study was performed with the approval of the Institutional Review Board (IRB) and informed consent was obtained from each patient before genetic testing. All the patients were diagnosed at the Department of Pediatric Dentistry, Faculty of Dentistry, Tehran University of Medical Sciences (TUMS). The pedigree analysis was carried out by Cyrillic 2.1 software.

Molecular Analysis

From each case, five milliliters peripheral blood was collected in test tubes containing EDTA. Then, DNA was extracted using DNGPLUS kit (Cinnagen, Tehran, Iran). PCR amplification was typically carried out using primer pairs of exon-intron boundaries of ENAM, KLK4 and MMP20 genes as described previously [25]. Also, FAM83H primers were designed using Primer3 software (Table 1), 0.2U Taq DNA polymerase (Roche, Mannheim, Germany), 10pmole of each primer, 200 μM of each dNTPs, .0.67μl of 50mM MgCl2, 60ng DNA and 2.5 μl of PCR buffer in 25μl of PCR reactions. The PCR conditions included an initial denaturation step for 3 min at 95°C, 30 sec at 95°C, 45 sec at 64°C with a 1°C decrease every second cycle down to 55°C, then 55°C for 14 cycles, 1 min at 72°C for extension, and finally 10 min at 72°C. PCR products were separated on1.8% agarose gels and stained with ethidium bromide. Subsequently, for detection any mutation, PCR products were purified using Quick Spin PCR purification kit according to manufacturer’s procedure (Qiagen). Then, purified PCR products were subjected for direct PCR sequencing using forward and reverse
primers as shown in Table 1 (Gene Fanavaran, Iran). Sequence data searches were performed in non-redundant nucleic and protein databases BLAST (http://www.ncbi.nlm.nih.gov/BLAST).

Table 1. List of FAM83H primer sequences used in screening of mutations in studied genes

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence 5′→3′</th>
<th>PCR Product Size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAM83PF1</td>
<td>CTCGCCAGGAGCCCCCTTGTCTGTA</td>
<td>486</td>
</tr>
<tr>
<td>FAM83PR1</td>
<td>GGAAGGCCGACAGGAAGT</td>
<td></td>
</tr>
<tr>
<td>FAM83PF2</td>
<td>CCCCCTCTCCCCTCTTAACG</td>
<td>523</td>
</tr>
<tr>
<td>FAM83PR2</td>
<td>CGCCCAAGGTTAGTCAT</td>
<td></td>
</tr>
<tr>
<td>FAM83PF3</td>
<td>CTACACCACGCGTGACAGTG</td>
<td>652</td>
</tr>
<tr>
<td>FAM83PR3</td>
<td>CGAGCGGAAATGCTCTG</td>
<td></td>
</tr>
<tr>
<td>FAM83PF4</td>
<td>GCCCTCCCCACAAAGGTG</td>
<td>497</td>
</tr>
<tr>
<td>FAM83PR4</td>
<td>CGTCTGTGCAAAGGAGTCG</td>
<td></td>
</tr>
<tr>
<td>FAM83PF5</td>
<td>GGGCTTATCTCGACAGGAAG</td>
<td>566</td>
</tr>
<tr>
<td>FAM83PR5</td>
<td>GACTCCCCGGAGATGGTAAAG</td>
<td></td>
</tr>
<tr>
<td>FAM83PF6</td>
<td>CAGGATTTATCGACAGGAAGG</td>
<td>593</td>
</tr>
<tr>
<td>FAM83PR6</td>
<td>GGGCTGAACATTATCTCTTTGTC</td>
<td></td>
</tr>
<tr>
<td>FAM83PF7</td>
<td>AGGCCATTCCTGGAGCCAAGAT</td>
<td>689</td>
</tr>
<tr>
<td>FAM83PR7</td>
<td>GACGGTGCAGAGATGAAGGT</td>
<td></td>
</tr>
</tbody>
</table>
Figure 1. Pedigree, clinical feature and molecular study of proband in family 1. A: The pedigree of family 1 showed two affected patients with ARHAI from a consanguineous family. A arrow indicates the proband. B: Panorex radiography of proband presented the tooth enamel loss. C: Clinical feature of proband demonstrating hypoplastic amelogenesis imperfecta. D: DNA sequencing revealed heterozygous genetic variation in the intron 5 of MMP20 gene (T18515C). E: Chromatogram from wild-type MMP20 gene (AY673603.1) (arrow indicates the position of nucleotide substitution).
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Results
The affected patients with ARHAI were studied from five Iranian consanguineous families (Fig. 1A, Fig. 2A). The proband from family 1 was a 17-year-old Iranian girl who had two healthy siblings, born to consanguineous parents. She and her siblings had been born with normal full-term pregnancy. Her first cousin was also affected with ARHAI. The clinical appearances of patients represented hypoplastic amelogenesis with yellow-brown discoloration. Their enamel showed also different range, from thin and smooth, to normal thickness with grooves, furrows and/or pits. All affected individuals were clinically and radiographically examined and showed non-syndromic enamel defects. Furthermore, the dental examination including an intra-oral evaluation of the parents and unaffected siblings showed no evidence of any enamel malformation or defect and no syndromic signs.

In family 2 the propositus was a 12-year-old boy affected with ARHAI. He and his healthy brother were born to first cousin parents. The proband's father, and his father's first cousin were also affected with ARHAI. The patients showed similar conditions to the index case of family 1.

PCR amplification of ENAM, KLK4, MMP20 and FAM83H genes were conducted and subsequently
subjected for direct PCR sequencing, and the sequencing data were analyzed. Two different MMP20 genetic variations were identified in two patients with amelogenesis imperfecta (Fig. 1D and Fig. 2D). Both mutations were heterozygous states, and they clustered in the intron4 of MMP20 gene.

**Discussion**

In this study we performed direct PCR sequencing for five families having at least one affected individual with ARHPIAI. The clinical appearances of hypoplastic amelogenesis imperfecta of our patients typically showed yellow-brown discoloration and verification of pathological enamel loss.

The amelogenesis imperfectas, are a clinically and genetically heterogeneous group of disorders characterized by faulty development of the tooth enamel due to hypoplasia or hypomineralization. Up to now, various enamel related genes such as ENAM, AMBN, AMELX, MMP20, KLK4, Amelotin and FAM83H genes have been suggested to be implicated in AIs. Our mutation screening in exon/intron boundaries of known candidate genes including, ENAM, KLK4, MMP20 and FAM83H could not detect any significant variations. However, we found two single nucleotide polymorphisms (SNP) including T18515C and G18504A in the forth intron of MMP20 gene.

Consistent with our results, different studies also, could not detect any significant mutations in the above genes in AIs [4, 25, 26]. In this regard, Santos et al (2007) evaluated genetic variation for the six major candidate genes including ENAM, AMBN, AMELX, MMP20, KLK4 and Amelotin genes in two Brazilian families with AI. This study indicated that the AI in these two families is not caused by any mutation in afore-mentioned genes for AI [26]. A study was carried out among seven Turkish families affected by AI. They evaluated the AMBN, AMELX, ENAM, FAM83H, KLK4, MMP20, and TUFT1 genes, but they also failed to detect any mutation in the studied genes [4]. More recent investigation has been performed in four Iranian families with AI[25]. In spite of the fact that the exons and intron/exon junctions of the candidate genes were sequenced, no gene mutations were identified in any individuals [25].

According to our results and previous studies, it is possible that the other genes and/or other mechanism such as epigenetic factors could act to the etiology of AI. Different polymorphisms in candidate genes responsible for amelogenesis imperfecta have been described in various studies [27-29]. Richard et al. studied polymorphism in AMELX exon 6 in an European population and they found low level of variations [28]. In the most recent study, a single nucleotide exchange was identified in the ENAM gene from a patient with the clinical features of AI [27]. Further studies are needed to find the exact nature of causation of amelogenesis imperfectas.

**References**

13. Simmer J.P.H., Lertlam R., Yamakoshi Y., and Hu J.C. Hypomutatuation enamel defects in Klk4 knockout/LacZ.