A Novel Mutation Involving the Initiation Codon of FGF3 in a Family Described with Complete Inner Ear Agenesis, Microtia and Major Microdontia (LAMM Syndrome)

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Abstract

LAMM syndrome (OMIM #610706) is a rare autosomal recessive syndrome characterized by the association of Michel aplasia, microdontia and malformation of the external ear. Different mutations in FGF3 gene were reported in several families presenting with this syndrome.

Clinical features and genetic results observed in a family with LAMM syndrome are reported. The diagnosis of isolated Michel aplasia was initially made in this family composed of two affected children. Microtia and microdontia was recently evidenced in both patients suggesting the diagnosis of LAMM syndrome. New auditory and oro-dental iconography was performed permitting to describe the patients’ phenotype in depth and to report rare findings of LAMM syndrome.

The sequencing of FGF3 gene identified a novel missense mutation (c.2T>G), substituting the first initiator methionine in arginine, in the fibroblast growth factor 3 (FGF3) at the homozygous state in both patients. LAMM syndrome was confirmed and appropriate genetic counseling performed.

Keywords: LAMM syndrome; FGF3 gene; Autosomal recessive inheritance; Michel aplasia; Microtia; Genetic counseling

Introduction

Whereas developmental defects of ears and/or teeth are very common, the combination of both remains a rare syndromic event. Michel aplasia, first described in 1863 [1] is a very rare malformation with complete bony and membranous aplasia of the inner ear. Michel aplasia in association with microdontia and malformation of the external ear (microtia type I) defines the LAMM syndrome (Labyrinthine Aplasia, Microtia, and Microdontia; OMIM 610706), which is a rare recessive syndrome recently related to mutations in the FGF3 gene.

Only a dozen of FGF3 mutations have been reported to date in various studies. Initially, three different mutations were described in 3 unrelated Turkish families: p.Arg104*, p.Ser156Pro and p.Val206Serfs*13 [2]. Since then, 9 other mutations were reported in 6 different studies [3-8].

We report herein the identification of a novel mutation affecting the initiation codon of FGF3 gene at the homozygous state (p.[Met1Arg];[Met1Arg] / c.[2T>G];[2T>G]) in two adults’ siblings with LAMM syndrome for whom the inner ear aplasia was reported in 1997 [9]. In addition to the identification of the mutation, we provide here the clinical and radiological phenotype of these 2 patients with novel RMI imaging features as well as the in depth description of the oro-dental phenotype.

Methods

Patients and related phenotype

The two patients described are the two children of a non-consanguineous couple, native of France. Informed consent was obtained from all the family, in accordance with the tenets of the Declaration of Helsinki.

Case 1

The first child, a boy, was referred to the genetic clinic when he was 1 year old due to a profound sensorineural hearing loss diagnosed a few months after birth. The pregnancy had been uneventful with no history of antenatal exposure to medical treatment, toxics or infections. The tympanogram was flat and the brainstem auditory evoked potential were absent on both sides. Moreover, X-rays of temporal bones revealed asymmetric diameters of external auditory canals. The clinical examination evidenced no dysmorphia, small anteverted bones revealed asymmetric diameters of external auditory canals. The clinical examination evidenced no dysmorphia, small anteverted auricles (Figures 1A-1D) and microodontia of the primary dentition. A dextrorotation of the heart and ocular dryness were also diagnosed during the first year of life. The external ear was operated on for esthetic purposes (Figures 1E and 1F). The boy, although profoundly deaf,
learnt to use phonic vibrations and labial lecture enabling a perfectly integrated school and young adult life. At the age of 17, an MRI showed an internal ear aplasia [9], also called “Michel abnormality” in reference to the initial description reported in 1863 [1].

As a young adult (31 years old), the patient sought for genetic counseling in order to evaluate the risk of recurrence for his offspring. At this occasion an in-depth dental and otic evaluation was performed. The dental examination revealed a generalized microdontia with widely spaced teeth, associated with tooth agenesis (Figures 2A-2D). More precisely, at the maxillary level, the right first premolar was absent with persistence of the deciduous molar (white arrow, Figure 2C). On the left side, the first deciduous molar also persisted despite the fact that no permanent premolars were absent there (star, Figure 2C). At the mandibular level, permanent central incisors were absent, with persistence of the deciduous incisors (Figures 2C and 2D). The right second premolar and first molar were also missing due to previous avulsions (Figure 2C). In addition to microdontia and agenesis, the clinical examination also showed a dyschromia of the teeth (Figures 2A and 2B), as well as enamel defects corresponding to a thinning of enamel mantle and the presence of pits on the molar and premolar occlusal tables (Figure 2E). The radiographs and the CT scan, carried out for dental implant planning surgery, confirmed these clinical signs (Figures 2C, 2F and 2G) and highlighted a taurodontism (elongation of the pulp chamber) on the first permanent molars (Figures 2F and 2G). The radiographs also revealed root abnormalities, consisting in root elongation (Figures 2C, 2D and 2F) and C-shaped morphology (Figures 2H and 2I).

To better characterize the otic abnormalities found in this patient, a novel generation CT scan was performed. Compared to Marsot-Dupuch et al. [9] this analysis confirmed the Michel aplasia (characterized by the bilateral aplasia of the inner ear structures associated with bilateral aplasia of the petrous apex and absence of development of the internal auditory canal) as well as the hypoplasia of the middle ear ossicles associated with a normal external auditory canal (Figures 3A-3C). However, abnormalities of the middle ear ossicles were better described with this new imagery. Indeed, the stapes, initially reported as absent, was in fact very hypoplastic. Moreover, the incus and the malleus, initially considered as normal, were deformed with a shortening of the long process of the incus. A new MRI was also realized, confirming the initial findings (described but not shown), notably the bilateral absence of the cochlea vestibular nerve and absence of development of the internal auditory canal in high resolution T2 weighted sequences focusing on the cerebellopontine angles (Figure 3D). The resolution of the new radiographic images permitted to define the abnormal course of the facial nerve. Indeed, after a normal origin, the course of the facial nerve passed in the lower part of the cerebellopontine angle and then followed a usual posterior path in the petrous apex. The CT scan confirmed its usual emergence in the stylo-mastoid foramen behind the styloid processes and sideward of jugular foramen.

On the basis of these clinical and radiological data, the diagnosis of isolated Michel aplasia initially established [9] moved to LAMM syndrome.
sequences. Informed consent was obtained from the 4 members of the family for DNA analysis. Genomic DNA was extracted from blood samples according to the manufacturer's protocol (Flexigene DNA kit, Qiagen). The three exons and exon-intron boundaries of the gene were PCR amplified with 50 ng of genomic DNA template. The primers were designed with Primer 3 (http://frodo.wi.mit.edu/primer3); detailed information on specific SNPs was accessible on supplementary table.

**Results**

Sequencing of FGF3 in both patients identified a novel homozygous missense mutation (p.[Met1Arg]; c.[2T>G]) in the first exon. The parents were heterozygous for the mutation (Figure 4A). This mutation substitutes the first amino acid, methionine, into an arginine. Because of the homozygous status of the mutations, we suspected a distant consanguinity, and performed a genome scan for case II.1 using an Affymetrix setting. Haplotype sharing of SNPs identified a homozygote region of 10 Mb on chromosome 11 encompassing FGF3 gene, between rs 1944130 and rs 568421 corresponding to physical position of 69,291,704 - 78,714,313 bp (Figure 4B). This homozygous block, associated with several others disturbed along the patient’s genome, suggested a distant consanguinity.

**Discussion**

The first case of LAMM syndrome was reported in 1991 by Hersch et al., who described a patient presenting with Michel/labyrinthine aplasia, microtia and microodontia [11]. The acronym “LAMM syndrome” was used first in 2007 by Tekin et al. with the identification of a homozygous mutation in FGF3 in three unrelated Turkish families including nine affected individuals [2]. The authors described three major phenotypic manifestations common to all individuals, which were then confirmed by further studies [3-8]: i) profound congenital neurosensorial deafness associated with the complete absence of inner ear structures bilaterally, including cochlea, vestibule and semicircular canals (Michel aplasia), ii) type I microtia with shortening of auricles, and iii) microodontia with widely spaced teeth. The individuals appeared to have normal middle ears structures and normal cognitive abilities, even if a delay in gross motor skills was noted for all of them, probably in relation with absence of inner ear and impaired balance. Type I microtia was associated with anteverted auricles in seven of nine individuals. The dental anomalies reported were variable with supernumerary teeth, absence of teeth, peg-shaped lateral incisors and loss of tooth heights due to abrasion. A mild micrognathia was also reported in some patients [2].

Herein, we report the identification of a novel FGF3 mutation in two siblings presenting with LAMM syndrome with the characteristic association of Michel/labyrinthine aplasia, microtia and microodontia. Compared to the literature, our patients presented the classical clinical and radiological symptoms of LAMM syndrome associated to mutations in FGF3, i.e., inner ear agenesis, microtia and microdontia associated with the anteverted auricles. Radiological investigations demonstrated anomalies classically described of the inner ear, but also defects at the level of the middle ear. This anomaly was reported only once before by Sensi et al., in two sibs presenting a bilateral involvement of middle ear structures in addition to the inner ear aplasia [8] (Figure 3). Our observation therefore confirms that middle ear defects can be found in LAMM syndrome. We also described bilateral absence of the cochlea vestibular nerve with otherwise normal brain and cerebellum.
The orodental examination showed abnormalities very similar to those reported in the literature: the CT-scan carried out for dental implant planning surgery, however also highlighted new features such as the presence of a taurodontism, as well as root abnormalities consisting in root elongation and C-shaped morphology. To our knowledge, this is the first time that such defects are reported with the orodental phenotype of LAMM syndrome.

Interestingly, the abnormal external ears and orodental phenotype were initially overlooked as the diagnosis of isolated Michel aplasia was suggested at first [9]. Years later a reappraisal of the orodental syndrome was completed. This finding was previously reported once in a family with LAMM syndrome [2].

In conclusion, we describe a novel homozygous mutation implying the first affected patient compared to 5 controls. Grey box: homozygous SNPs (AA or BB). White box: heterozygous SNPs (AB).

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Indeed, a very low risk genetic counseling could be advocated to the siblings for their offspring concerning the recurrence of this disabling disease that necessitate a specialized education for severe deafness (no cochlear implantation for obvious anatomical reasons) as well as dental implants in adulthood.

Clinically, our report confirmed the possibility of middle ear abnormalities in LAMM syndrome, described only once to date. Moreover, we described in detail the orodental phenotype observed in LAMM syndrome. This report highlights the importance of a global clinical examination of the patient behind a symptom seemingly isolated and especially the importance of dental examination as a highly valuable diagnostic guidance.

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References


Figure 5: Mutations reported in FGF3 to date. Each mutation was reported in only one family except for the mutations p.Arg95Trp and p.Arg104* described respectively in 2 and 3 independent families (indication in exponent in the figure). The mutation identified in this study is underlined.