Simultaneous Occurrence of an Autosomal Dominant Inherited MSX1 Mutation and an X-linked Recessive Inherited EDA Mutation in One Chinese Family with Non-syndromic Oligodontia

Xiao Xia ZHANG¹, Sing Wai WONG¹,², Dong HAN¹, Hai Lan FENG¹

Objective: To describe the simultaneous occurrence of an autosomal dominant inherited MSX1 mutation and an X-linked recessive inherited EDA mutation in one Chinese family with non-syndromic oligodontia.

Methods: Clinical data of characteristics of tooth agenesis were collected. MSX1 and EDA gene mutations were detected in a Chinese family of non-syndromic oligodontia.

Results: Mild hypodontia in the parents and severe oligodontia in the son was recorded. A novel missense heterozygous mutation c.517C>A (p.Arg173Ser) was detected in the MSX1 gene in the boy and the father. A homozygous missense mutation c.1001G>A (p.Arg334His) was detected in the EDA gene in the boy and the same mutant occurred heterozygously in the mother.

Conclusion: Simultaneous occurrence of two different gene mutations with different inheritance patterns, which both caused oligodontia, which occurred in one subject and in one family, was reported.

Key words: EDA, inheritance, MSX1, mutation, oligodontia

Hypodontia (the congenital missing of teeth, which is also known as tooth agenesis) is the most common developmental anomaly in men¹. Oligodontia, usually referring to a severe hypodontia, is defined as the congenital absence of six or more teeth, excluding the third molars, and may present as part of the syndrome, while the non-syndromic oligodontia is more commonly found². Although many oligodontia cases presented sporadically without an inheritance trait, researchers have found more evidence that genetic defects play a major role in the aetiology³,⁴. In the last two decades, mutations were identified to be associated with non-syndromic oligodontia in many genes, including muscle segment homeobox 1 (MSX1), paired box 9 (PAX9), axin inhibition protein 2 (AXIN2), wingless-type mouse mammary tumour virus integration site family member 10A (WNT10A), nuclear factor-kappa-B essential modulator (NEMO), keratin 17 (KRT17) and ectodysplasin A (EDA), that formerly was found to cause the most typically syndromic oligodontia of hypohidrotic ectodermal dysplasia (HED)/anhidrotic ectodermal dysplasia (EDA)⁵-¹⁷.

Non-syndromic oligodontia has wide phenotypic heterogeneity and can be inherited in an autosomal dominant, autosomal recessive, or X-linked manner¹⁸, which were mostly autosomal dominant in the MSX1 mutation or an X-linked recessive trait in the EDA mutation.
The MSX1 gene located in 4p16.2 encodes a member of the muscle segment homeobox gene family. The encoded protein functions as a transcriptional repressor during embryogenesis through interactions with components of the core transcription complex and other homeoproteins. It may also have roles in limb-pattern formation, craniofacial development, particularly odontogenesis, and tumour growth inhibition. MSX1 was the first gene identified causing family non-syndromic tooth agenesis in humans, which was also reported to be associated with orofacial clefting (lip and palate cleft), tooth agenesis and Witkop syndrome, known as tooth and nail syndrome (TNS), a rare autosomal dominant disorder, with the characteristics of affected individuals having nail dysplasia and several congenitally missing teeth. Later, more mutations in MSX1 were identified causing non-syndromic oligodontia, mostly missense mutations or nonsense mutations in autosomal dominant inheritance.

Almost at the same time when the MSX1 mutation was first identified to be a cause of oligodontia, mutation in a novel transmembrane protein of EDA was found to cause X-linked anhidrotic ectodermal dysplasia (EDA)/hypohidrotic ectodermal dysplasia (HED). Then, different EDA mutations were continuously identified to be associated with EDA/HED families or patients. Until 2006, it was reported that a novel missense mutation in the EDA gene in all affected males and carrier females of a Mongolian family, with only a congenital absence of teeth, inherited the mutation in an X-linked fashion. The affected members of the family did not show other HED characteristics, except hypodontia. Ever since, many literatures reporting EDA mutations in non-syndromic oligodontia appeared, and the EDA gene became one of the most likely genes to cause non-syndromic oligodontia.

In this study, two mutations in MSX1 and EDA were identified, both in the boy and one in the father and the mother, which was inherited as an autosomal dominant trait and X-linked recessive trait respectively, in a Chinese family with non-syndromic oligodontia. It is very interesting that a rare co-occurrence arose in such a way. To our knowledge, no literature has been found reporting similar conditions.

Materials and methods

Subjects

A 16-year-old male patient requested treatment advice regarding some of his congenitally missing permanent teeth, and underwent the oral examination at the Department of Prosthodontics, Peking University, School and Hospital of Stomatology. It was confirmed that no teeth extraction occurred or any other developmental abnormalities, except the missing teeth, according to their medical history and the description given by his parents. Oral examination, cast analysis and panoramic

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<th>Subjects</th>
<th>Congenitally missing teeth</th>
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<tr>
<td>Son</td>
<td>8 7 6 5 4 3 2 1 1 2 3 4 5 6 7 8</td>
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<tr>
<td></td>
<td>Max.</td>
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<td>Mother</td>
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Note: Max – maxillary; Man – mandibular; 1 – incisor; 2 – lateral incisor; 3 – canine; 4 – first premolar; 5 – second premolar; 6 – first molar; 7 – second molar; 8 – third molar; * – congenitally missing permanent teeth; # – extracted teeth; & – conical-shaped teeth; § – ectopic eruption.
radiographs revealed the diagnosis of non-syndromic oligodontia. His father and mother were recruited later and clinical data were collected as described above. They were both finally diagnosed with mild hypodontia. Peripheral blood was taken from the family for the subsequent detection of genomic mutations of candidate genes that associated with oligodontia, including MSX1 and EDA. One hundred normal volunteers were selected as control individuals for the mutation detection experiments. The study was approved by the Institutional Review Board of Peking University School and Hospital of Stomatology. Informed consent was obtained from the family, as well as the normal controls.

**DNA extraction and detection of mutations**

Genomic DNA was extracted from peripheral blood lymphocytes using the Biotec DNA minikit (Biotek, Beijing, China) according to the manufacturer’s instruction. Also genomic DNA of normal volunteers were extracted from the buccal epithelial cells using swab DNA minikit (Tiangen, Beijing, China), according to the manufacturer’s instructions.

The entire coding region and the intron-exon junctions of MSX1 and EDA genes were amplified by polymerase chain reaction (PCR) with AmpliTaq Gold 360 Master Mix (Applied Biosystems, California, USA). The PCR products were sent to the Sangon Biotech Company (Beijing, China) for direct sequencing, using a Big Dye terminator v3.1 (Applied Biosystems) and a 3730 DNA sequencer (Applied Biosystems). Primer sequences and PCR conditions were available upon request. SeaMan Pro genetic analysis software (DNASTAR, Wisconsin, USA) was used for sequencing analysis.

**Results**

**Clinical data**

The whole family including the proband boy and his parents, all of whom were in good physical health and well-being. Hypodontia only affected permanent dentitions. The original status of congenitally missing teeth and oral abnormalities at the first visit was recorded according to the examination and the medical history as in Table 1. The father had a full dentition of 28 teeth excluding the third molars, but there were diastemas in the maxillary anterior region due to the maxillary conical lateral incisors and the obvious smaller width of all present teeth (Table 1; Figs 1a and 1b). The father’s panoramic radiograph in Figure 1 was taken when some of the molars were extracted for periodontal disease after several years had passed. The mother’s two mandibu-
molars (17, 27 and 37) and one premolar (15), with retained deciduous teeth i.e. 55, 53, 63 and 71. Coinciding with the congenitally missing teeth, ectopic eruption of both maxillary canines and the torquing left maxillary incisor were observed (Table 1; Figs 1e, 1f and 1g). The boy was the only child of the family, and there was no exact or reliable medical records obtained from the relatives of the parents about the tooth agenesis and other ectodermal dysplasia-related conditions.

**Mutations detected**

**MSX1**: A missense heterozygous mutation c.517C>A (p.Arg173Ser) was detected in exon 2 of MSX1 in the boy and the father, however it was not detected in the mother (Fig 2)

**EDA**: A homozygous missense mutation of c.1001G>A (p.Arg334His) was detected in exon 8 of the EDA in the boy; the same mutant occurred heterozygously in the mother, while it was not detected in the father (Fig 3).

None of the mutations were detected in normal controls in this study.

**Discussion**

In a systematic review and meta-analysis on the genetic background of non-syndromic oligodontia, which includes articles published up to March 2012, it was reported that PAX9, EDA and MSX1 are the first three genes currently known to have a potential for causing non-syndromic oligodontia. Numerous literatures reported that different mutations in different genes may cause family non-syndromic oligodontia, but to our knowledge, it is unreported that two different mutations in the most common oligodontia-associated genes occurred in one family with different inheritance traits, which is a very uncommon occurrence.

The MSX1 gene mutation identified in this study is a novel mutation which occurred in exon 2 of the gene. Like most of the mutations previously identified in MSX1, the mutation found in this study was also a missense mutation c.517C>A that changed one amino acid (p.Arg173Ser) in the highly conserved homeodomain. However, the mutation of c.1001G>A (p.Arg334His) in exon 8 of the EDA gene in this family is the same as the known mutation reported, and the previous structural analysis indicated that it would produce conformational changes, potentially altering the stability of the EDA homotrimers.

The most interesting thing in this study is the relationship between the genotypes and phenotypes, both
in individual appearance and in the family inheritance. It has been reported that the size of remaining teeth in oligodontia patients is significantly smaller compared to normal controls\(^2,3\). This is the same with our clinical observation and previous studies\(^29\). The summary of non-syndromic oligodontia phenotypes associated with mutations, showed that MSX1 mutations are more likely linked to the agenesis of the second premolar and the four third molars\(^5,30\). While the hypodontia phenotype in EDA-associated non-syndromic tooth agenesis seems to favour the lack of incisors, other teeth are involved as well, and carrier females generally show a much milder phenotype with only one or two teeth missing or no symptoms at all\(^27\).

In the present family, although the father had a full dentition of 28 teeth, he had no developed third molars and all remaining teeth showed a reduction in size, causing diastemas. The EDA mutation carrier mother just had mild symptoms i.e. absence of two mandibular incisors and peg-shaped maxillary incisors. However, the son inherited two sites of mutations through different traits, had a more severe phenotype of tooth agenesis than his parents, with congenitally missing teeth, mainly in the incisors and molars. Since c.518G>C (p.Arg173Pro) has been recorded as one of the MSX1 SNPs\(^31\), which is similar with the mutation c.518G>C (p.Arg173Pro) has been recorded as one of missing teeth, mainly in the incisors and molars. Since of tooth agenesis than his parents, with congenitally

Conflict of interest
The authors reported no conflicts of interest related to this study.

Author contribution
Dr Xiaoxia Zhang for the clinical data and blood sample collection, for designing the study and writing the paper; Dr Singwai Wong and Dr Dong Han for the mutation detection experiments; and Dr Hailan Feng for the overall design of the research.

References