

# Mutation analysis by direct and whole exome sequencing in familial and sporadic tooth agenesis

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**Abstract.** Dental agenesis is one of the most common congenital craniofacial abnormalities. Dental agenesis can be classified, relative to the number of missing teeth (excluding third molars), as hypodontia (1 to 5 missing teeth), oligodontia (6 or more missing teeth), or anodontia (lack of all teeth). Tooth agenesis may occur either in association with genetic syndromes, based on the presence of other inherited abnormalities, or as a non-syndromic trait, with both familial and sporadic cases reported. In this study, we enrolled 16 individuals affected by tooth agenesis, prevalently hypodontia, and we carried out direct Sanger sequencing of paired box 9 (PAX9) and Msh homeobox 1 (MSX1) genes in 9 subjects. Since no mutations were identified, we performed whole exome sequencing (WES) in the members of 5 families to identify causative gene mutations either novel or previously described. Three individuals carried a known homozygous disease mutation in the Wnt family member 10A (WNT10A) gene (rs121908120). Interestingly, two of these individuals were siblings and also carried a heterozygous functional variant in EDAR-associated death domain (EDARADD) (rs114632254), another disease causing gene, generating a combination of genetic variants never described until now. The analysis of exome sequencing data in the members of other 3 families highlighted new candidate genes potentially involved in tooth agenesis and considered suitable for future studies. Overall, our study confirmed the major role played by WNT10A in tooth agenesis and the genetic heterogeneity of this disease. Moreover, as more genes are shown to be involved in tooth

agenesis, WES analysis may be an effective approach to search for genetic variants in familiar or sporadic tooth agenesis, at least in more severe clinical manifestations.

## Introduction

Dental agenesis is one of the most common congenital anomaly of human dentition, with a prevalence (excluding third molars) ranging from 2.6 to 11.3% (1). Relative to the number of missing teeth, dental agenesis can be classified as hypodontia (when 1 to 5 teeth are missing), oligodontia (when 6 or more teeth are missing), or anodontia (when the complete failure of dentition development occurs). The most frequently missing teeth are the mandibular second premolars, followed by the maxillary lateral incisors and maxillary second premolars (2). It is generally accepted that both genetic and environmental factors play a significant role in the pathogenesis of this disease (3). Tooth agenesis may occur either in association with genetic syndromes characterized by other inherited anomalies as an isolated non-syndromic familial trait, or as sporadic cases. Familial tooth agenesis is characterized by moderate genetic heterogeneity and has been reported to have an autosomal-dominant, autosomal-recessive or X-linked mode of inheritance (4,5).

Tooth development requires a complex network of molecular interactions in which the activation of the nuclear factor- $\kappa$ B (NF- $\kappa$ B) and WNT signaling pathways indeed play a central role. The transcription factor, NF- $\kappa$ B, controls several cellular functions and it is activated by members of the tumor necrosis factor receptor (TNFR) superfamily, including EDA receptor (EDAR). Ectodysplasin (EDA) is a member of the tumor necrosis factor (TNF)-related ligand family and binds to its membrane receptor, EDAR, that in turn binds to its adaptor, EDAR-associated death domain (EDARADD). The EDA-EDAR-EDARADD complex activates the NEMO-IKK signaling cascade, allowing the nuclear translocation of NF- $\kappa$ B and thus the transcriptional control of genes that are crucial for tooth development (6).

WNT signaling molecules play important roles in the differentiation of tissues and organs during embryonic development. The secretion of WNTs, including WNT4, WNT6, and WNT10 from the dental epithelium is essential for tooth development. WNT10A is expressed in the dental epithelium

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and mesenchyme. Similar to other WNT proteins, it binds to the Frizzled (Fz) transmembrane receptors and to lipoprotein receptor-related protein 5/6 (LRP5/6) coreceptors, leading to the activation of the  $\beta$ -catenin pathway (7).

One of the direct downstream targets of WNT/ $\beta$ -catenin signaling during craniofacial development is Msh homeobox 1 (MSX1). MSX1 is a homeobox gene encoding a transcription factor that regulates the expression of bone morphogenetic protein 4 (BMP4), a member of the transforming growth factor- $\beta$  (TGF- $\beta$ ) superfamily, during the bud and cap stages of the tooth development. In these phases, MSX1 interacts with paired box 9 (PAX9) both at the gene and protein levels, with MSX1 and PAX9 being intimately involved in the genetic networks regulating tooth development (8). PAX9 is a transcriptional factor that plays a key role during embryogenesis by modifying the transcriptional activity of downstream genes. In particular, by physically associating with MSX1, it enhances both MSX1 and BMP4 gene expression during tooth development (9,10). On the other hand, Axin 2 (AXIN2), also known as conductin/axil, is an inhibitor of the WNT signaling pathway (11).

Previously, dominant loss-of-function mutations in MSX1, PAX9 and AXIN2 have been described in familial forms of non-syndromic tooth agenesis (12). Recently, the WNT10A variant has been described in up to 50% of analyzed patients with dental agenesis and also in various ectodermal dysplasia syndromes (13,14). Mutations in EDA, EDAR and EDARADD have also been shown to be associated with both isolated tooth agenesis and syndromic tooth agenesis, such as X-linked hypohidrotic ectodermal dysplasia (XLHED), a rare anomaly involving sparse hair, dental abnormalities, skin lesions and hypoplasia of sweat glands (15).

The initial aim of this study was to evaluate the MSX1 and PAX9 mutation rate in a cohort of patients and subsequently to identify the causative gene mutations associated with autosomal tooth agenesis in some families by employing whole exome sequencing (WES).

## Subjects and methods

**Study subjects.** The probands (n=9; Fig. 1) were patients who were 10 to 20 years of age enrolled at the Dental Clinic of the University of Brescia, Italy, from 2008 to 2014, with a confirmed diagnosis of dental agenesis, but without systemic or syndromic diseases. When available, parents and siblings either affected or not, were also enrolled in the study. All individuals enrolled were informed of the purpose of this study and signed an informed consent. All clinical and genetic studies were approved by the Ethics Committee of the Spedali Civili of Brescia (AOBS-GENI-2011; NP 1119) and were conducted according to the principles expressed in the Declaration of Helsinki. In addition to panoramic radiography, a clinical examination was conducted on a dental chair, under artificial light, with a probe and a dental mirror by two trained dentists.

**Mutation analysis by Sanger sequencing.** Genomic DNA was isolated from the peripheral blood of affected individuals and available affected/non-affected relatives using the Purgene Blood Core kit (Qiagen, Valencia, CA, USA) following the instructions of the manufacturer. DNA from peripheral blood of a control (CTRL) unaffected subject was also isolated. The

PAX9 and MSX1 coding sequences were amplified by PCR using the primers and the thermal cycler protocols previously described (16-18). The amplified products were sequenced using the BigDye Terminator v3.1 Cycle Sequencing kit and analyzed using the ABI PRISM 310xl Genetic Analyzer (both from Applied Biosystems, Foster City, CA, USA). The results were compared with the reference sequences for each (MSX1, NM\_002448.3; PAX9, NM\_006194.3).

The exons and approximately 30 bp of the flanking introns of the WNT10A and EDARADD genes were amplified with the primers previously described (19). The results were compared with the reference sequences for each gene (WNT10A, NM\_025216.2; EDARADD, NM\_080738.3).

**Whole exome sequencing (WES).** WES was carried out in selected affected individuals and their relatives. In total, 12 cases were examined by WES and the details of the subjects analyzed are reported in Fig. 1. The exomes of the probands of families 1, 5, 7 and 8 were sequenced at Personal Genomics (University of Verona, Italy) using the TruSeq Exome Enrichment kit (Illumina, San Diego, CA, USA) for whole exome capture. The members of family 5 (I-1, I-2, II-2), family 8 (I-1, I-2) and family 9 (I-2, II-2, II-3) had their exomes sequenced at Erasmus MC, University Medical Center Rotterdam using the Nimblegen SeqCap EZ Exome version 2.0 exome capture kit. In both protocols, paired-end libraries were sequenced on the Illumina HiSeq instrument. Sequencing reads were aligned to the human genome reference sequence (hg19) using BWA-mem (20) and then processed according to GATK 3.4-46 best practices (21) for variant discovery. Briefly, aligned reads were processed for duplicate removal using Picard and the base quality score recalibration was performed using GATK. The processed BAM files of single probands or the proband and relatives were then analyzed using GATK HaplotypeCaller for the identification of both single nucleotide polymorphisms (SNPs) and indel variants. Reported variants in VCF file format were then filtered removing those with a read depth <6 and a genotype quality (GQ) <20. Subsequently, the filtered variants were annotated using ANNOVAR (22) and selected according to the following criteria: i) exonic, non-synonymous variants; ii) reported allele frequency in 1000G phase 3 data and ExAc 0.3<1%; and iii) not located in segmental duplicated region. These variants were searched for mutations in known genes related to dental agenesis. Where no mutations emerged, the data of the family pedigree were analyzed to select variants segregating according to the disease hereditary model. These variants were further annotated with known phenotypes from the ClinVar database (23) and the RVIS score (24) that provides an estimate of the tolerance to functional variations for each gene. Candidate genes variants were then prioritized based on PolyPhen2 (25) and FATHMM (26) deleteriousness predictions and the RVIS score of the associated gene.

## Results

**Study subjects.** A total of 9 females and 7 males ranging in age from 8 to 48 years affected by tooth agenesis from 9 different families were enrolled in this study (Fig. 1). In particular, 5 individuals were affected by oligodontia (number of missing teeth  $\geq 6$ ) and 11 individuals by hypodontia (number

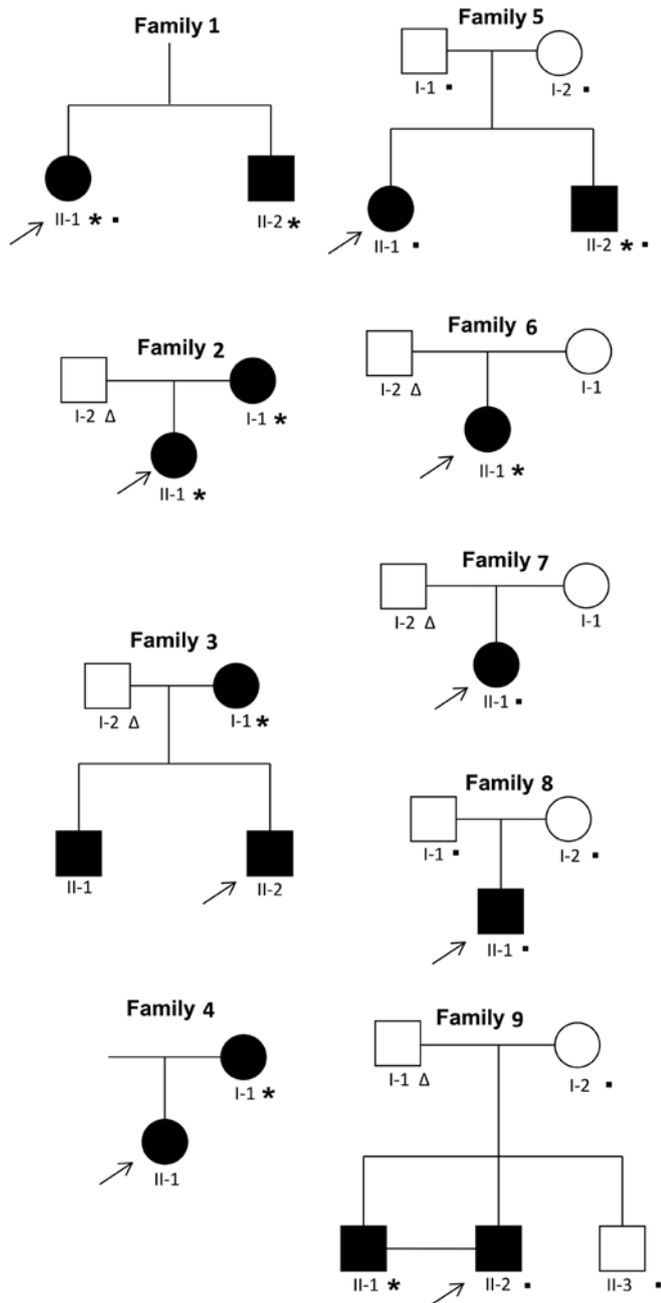


Figure 1. Pedigrees of the 9 families in this study. Squares indicate males, and circles indicate females. Filled symbols represent individuals diagnosed with tooth agenesis; empty symbols represent unaffected subjects. The asterisks indicate individuals for which mutational analysis for the PAX9 and MSX1 genes was performed; filled squares indicate subjects who underwent whole exome sequencing (WES) analysis; empty triangles indicate that no DNA was available. Arrows indicate the probands of each family.

of missing teeth  $\leq 5$ ). The most common missing teeth were the mandibular second premolars (35 in 10/16 affected; 45 in 9/16 affected), maxillary lateral incisors (12 in 5/16 affected; 22 in 5/16 affected) and maxillary second premolars (15 in 4/16 affected; 25 in 4/16 affected) (Tables I and II).

**Mutation analysis of PAX9 and MSX1 genes.** To identify causative mutations in subjects with tooth agenesis, we performed direct sequencing of the PAX9 and MSX1 genes in 9 affected subjects (Fig. 1, individuals marked with an asterisk).

Table I. Characteristics of the probands and the affected relatives.

Characteristic	N	%
Males	7	44
Females	9	56
Type of tooth agenesis (total number of cases)	16	
Hypodontia (1-5 permanent teeth missing)	11	69
Oligodontia (>6 permanent teeth missing)	5	31
Type of permanent teeth missing (total teeth missing)	81	
Central incisor (11, 21, 31, 41)	8	9.9
Lateral incisor (12, 22, 32, 42)	16	19.7
Canine (13, 23, 33, 43)	4	4.9
First premolar (14, 24, 34, 44)	9	11.1
Second premolar (15, 25, 35, 45)	27	33.3
First molar (16, 26, 36, 46)	1	1.2
Second molar (17, 27, 37, 47)	6	7.4
Third molar (18, 28, 38, 48)	10	12.3
No. of missing incisors	35	43.2
No. of missing canines	4	4.9
No. of missing premolars	25	30.8
No. of missing molars	17	21

Ultimately, the mutation analysis of the PAX9 and MSX1 genes did not highlight the presence of any causative mutation in the 9 affected individuals.

**WES.** Since the genetic heterogeneity of tooth agenesis has become apparent and has expanded since our first screening, to include 17 different genes, we decided to apply WES to identify novel and/or previously described causative mutations, as well as new candidate genes. Overall, we performed WES on 12 subjects from 5 distinct families (Fig. 1, individuals marked with a filled square; Table II, individuals highlighted in grey) with a mean coverage across target region of 75X-171X and 22,482-37,874 variants identified per subject in exonic regions (Table III).

For 2 families, it was possible to identify known causative mutations, thus providing a molecular diagnosis. The already described p.F228I mutation (rs121908120) in the homozygous state in the WNT10A gene was detected in the proband of family 7 and confirmed by Sanger sequencing; her unaffected mother displayed the same variant in the heterozygous state (Fig. 2). The same mutation was also detected in both the affected siblings from family 1 (Fig. 3). Moreover, the proband (II-1, oligodontia) and her brother (II-2, more severe phenotype) also carried a p.S103F mutation (rs114632254) in the EDARADD gene in the heterozygous state (Fig. 3). Both mutations were confirmed by Sanger sequencing. In 3 other families, our analysis led to the identification of novel candidate variants in genes with a potential role in tooth agenesis.

In family 5, an autosomal recessive form of inheritance seemed to be most probable (Fig. 1). In both the affected



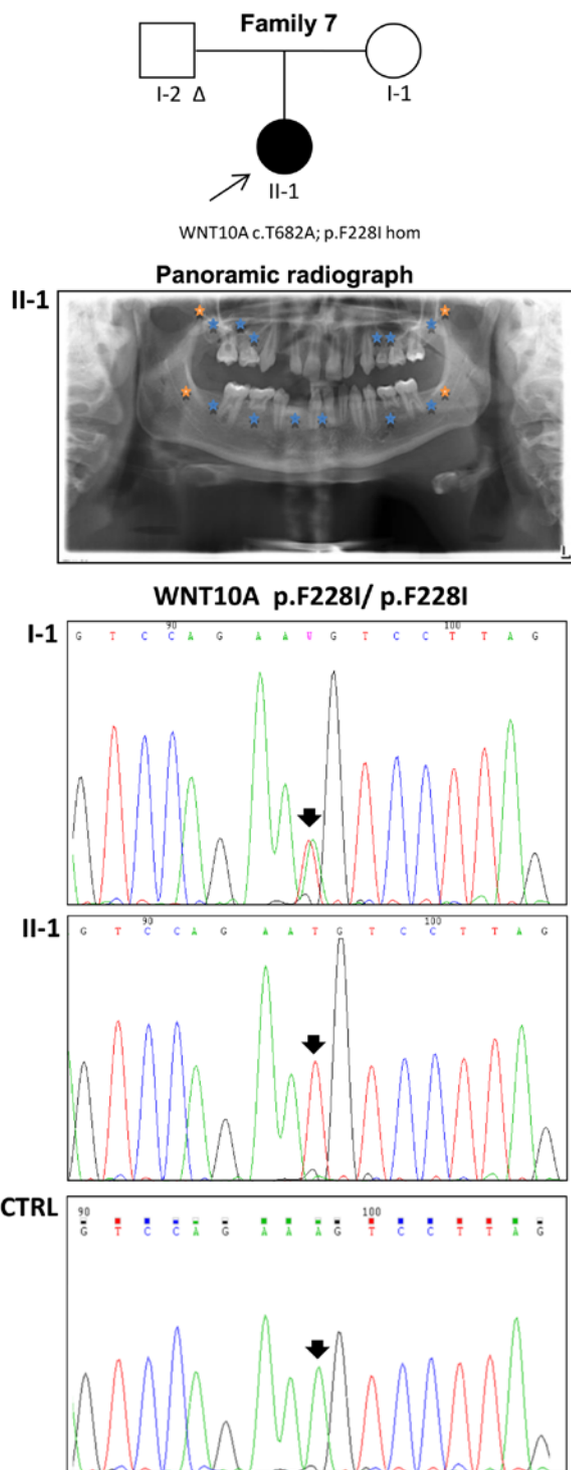


Figure 2. Panoramic radiograph of dentition of the proband (II-1) of family 7. The missing teeth are indicated by asterisks. Partial DNA sequences of exon 3 of WNT10A from the unaffected mother (subject I-1), the proband (II-1) and from a control subject (CTRL). The nucleotide substitution is indicated by an arrow.

siblings, we found compound heterozygous mutations in the Mucin 16 (MUC16) and Titin (TTN) genes and a homozygous genetic variation c.-387delC/G in the 5'UTR portion of the paired-like homeodomain 2 (PITX2) gene. Furthermore, in the proband (II-1) and in her brother, we detected an SNP (c. T455C:p.V152A; rs17563) in the homozygous state in the BMP4 gene previously associated with tooth agenesis (Table IV).

Table III. Summary of whole exome sequencing results.

Sample	Family	Mean coverage in target region	% bases covered at least 20X	No. of exonic variants
II-1	5	100X	93.6	23,297
II-1	1	171X	95.5	22,801
II-1	7	77X	91.8	22,482
II-1	8	82X	92.2	23,564
I-1	5	75X	83.9	22,603
II-2	5	82X	85.3	22,822
I-2	5	82X	85.3	22,706
I-1	8	83X	85.8	22,825
I-2	8	80X	84.9	23,129
I-2	9	82X	91.6	37,874
II-2	9	100X	92.7	37,822
II-3	9	88X	91.1	37,556

All results are reported considering the human CDS sequences as target region. Family ID are the same as those reported in Fig. 1.

Family 8 was also likely to present a recessive form of inheritance (Fig. 1). The proband displayed homozygous non-synonymous single nucleotide variants (SNVs) in the aryl-sulfatase family member H (ARSH), proline rich 32 (PRR32) and apurinic/apyrimidinic endodeoxyribonuclease 2 (APEX2) genes, a nucleotide change in homozygosis in transglutaminase 4 (TGM4) resulting in a premature stop codon and a 40 bp frameshift insertion in homozygosis in the placenta specific 4 (PLAC4) gene. Putative *de novo* heterozygous genetic variants were identified in 14 genes, 4 of whom displayed compound heterozygous gene alterations. Furthermore, the homozygous genetic variant in the 5'UTR portion of the PITX2 gene identified in family 6 was also present in the proband of family 8 (Table V).

Finally, an autosomal recessive form of inheritance was likely also for family 9. The father could not be included in the study and the analysis was conducted on the twin proband (II-2) and on his unaffected mother and brother (Fig. 1). The proband evidenced a non-synonymous SNP in the heterozygous state in the ankyrin repeat and EF-hand domain containing 1 (ANKEF1), COMM domain-containing protein 7 (COMMD7), DDHD domain containing 1 (DDHD1), early B-cell factor 2 (EBF2), envoplakin (EVPL), SPT6 homolog, histone chaperone (SUPT6H) genes and a compound heterozygous mutation in the succinate-CoA ligase GDP-forming beta subunit (SUCLG2) gene (Table VI).

## Discussion

The main objective of this study was to identify the causative genetic defects in subjects affected by non-syndromic tooth agenesis. We began with Sanger sequencing of the PAX9 and MSX1 genes, since they were considered in the literature the most probable candidate genes (27). This analysis performed on 9 individuals affected by agenesis did not evidence any muta-

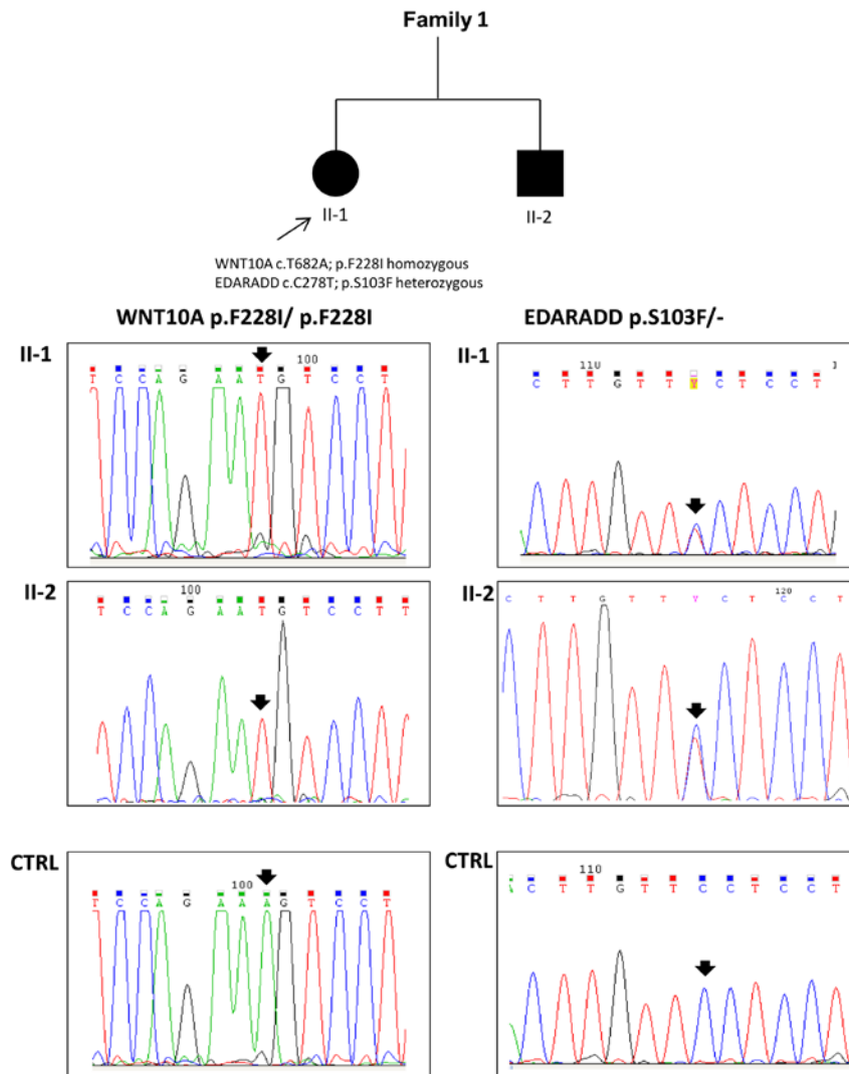


Figure 3. Direct sequencing analysis of mutation found by whole exome sequencing (WES) in family 1. Partial DNA sequences of exon 3 of WNT10A (left panels) and exon 7 of EDARADD (right panels) from the proband (II-1), her brother (II-2) and from a control subject. Nucleotide substitution is indicated by an arrow.

tion, thus suggesting that other genes may be involved in tooth agenesis, in agreement with other published data (28). Mutations in the WNT10A gene emerged as frequently involved (13,14). We decided to use the WES technology to identify causative mutations. We identified pathogenic mutations in the proband of family 1 and the same alterations in her brother. The subjects displayed a known homozygous disease mutation in the WNT10A gene (p.F228I, rs121908120) and a heterozygous mutation in the EDARADD gene (p.S103F, rs114632254). The frequent p.F228I aminoacid substitution is associated with the autosomal dominant or autosomal recessive form of isolated hypodontia and ectodermal dysplasia (29). The EDARADD mutation p.S103F was previously predicted to be harmful for protein function by bioinformatics analysis (19). This particular combination of genetic variants involving two genes related to teeth development has never been previously described, at least to the best of our knowledge. Both the affected individuals had oligodontia, but individual II-1 had 17 missing teeth and subject II-2 had 6 missing teeth. By WES, we identified the pathogenic mutation in the proband of family 8 that also displays the p.F228I mutation in WNT10A in the homozygous state.

To extend the WES analysis, we focused on families 5, 8 and 9, as genomic DNA was available from all the members with the exception of the unaffected father (I-1) of family 9. For family 5, an autosomal recessive form of inheritance seemed most probable. Our analysis identified the PITX2, TTN and MUC16 genes as affected by candidate mutations. The TTN and MUC16 genes were not considered good candidate genes for agenesis prediction, given their high RVIS score, confirming the propensity of these genes to accumulate rare functional mutations with neutral effect (30). Of note, we found a homozygous c.-387delC/G variation in the 5'UTR of the PITX2 gene in the siblings that could be associated with tooth agenesis as we mentioned above. PITX2 is a transcription factor that initiates tooth development, activating amelogenin expression, whose protein product is necessary for enamel formation. Since the mutation is localized in the promoter region of the gene, it may affect its transcriptional activity (31). A more in depth analysis of genetic variants in this family also identified an interesting variant in the BMP4 gene, a non-synonymous SNP in the homozygous state (rs17563; c.T455C:p.V152A) in the affected siblings II-1 and II-2. In a Brazilian study conducted

Table IV. Selected candidate genes and variants in family 5.

Gene.refGene	ExonicFunc. refGene	Chr	Ref	Alt	Func.refGene	AAChange.refGene
Compound						
MUC16	Non-synonymous SNV	chr19	G	T	Exonic	MUC16:NM_024690:exon3:c.C12953A:p.A4318D
MUC16	Non-synonymous SNV	chr19	T	C	Exonic	MUC16:NM_024690:exon3:c.A23897G:p.E7966G
MUC16	Non-synonymous SNV	chr19	C	T	Exonic	MUC16:NM_024690:exon12:c.G36581A:p.R12194Q
TTN	Non-synonymous SNV	chr2	C	G	Exonic	TTN:NM_003319:exon186:c.G75952C:p.E25318Q, TTN:NM_133432:exon187:c.G76327C:p.E25443Q, TTN:NM_133437:exon187:c.G76528C:p.E25510Q, TTN:NM_133378:exon307:c.G95443C:p.E31815Q, TTN:NM_001256850:exon308:c.G98224C:p.E32742Q, TTN:NM_001267550:exon358:c.G103147C:p.E34383Q
TTN	Non-synonymous SNV	chr2	C	T	Exonic	TTN:NM_003319:exon73:c.G18133A:p.D6045N, TTN:NM_133432:exon74:c.G18508A:p.D6170N, TTN:NM_133437:exon74:c.G18709A:p.D6237N, TTN:NM_133378:exon194:c.G37624A:p.D12542N, TTN:NM_001256850:exon195:c.G40405A:p.D13469N, TTN:NM_001267550:exon245:c.G45328A:p.D15110N
TTN	Non-synonymous SNV	chr2	T	C	Exonic	TTN:NM_133378:exon90:c.A23131G:p.I7711V, TTN:NM_001256850:exon91:c.A25912G:p.I8638V, TTN:NM_001267550:exon93:c.A26863G:p.I8955V
TTN	Non-synonymous SNV	chr2	G	A	Exonic	TTN:NM_133378:exon77:c.C19445T:p.S6482L, TTN:NM_001256850:exon78:c.C22226T:p.S7409L, TTN:NM_001267550:exon80:c.C23177T:p.S7726L
Known						
PITX2	UTR5	chr4	C	-	NM_000325: c.-387delG	
BMP4	Exonic	chr14	T	C	NM_001202: c.T455C: p.V152A	

on 46 individuals with tooth agenesis and 88 control cases, the CC genotype of BMP4 was more frequent in individuals with 3 or more missing teeth than in the control group ( $p < 0.0001$ ), leading the authors to the conclusion that this variant was associated with tooth agenesis (32). The same SNP was previously found in two Mexican families with oligodontia (33). Capasso *et al* (34) observed that the c455T>C substitution altered the BMP4 mRNA secondary structure and that the BMP4 mRNA and protein expression levels were higher for the T allele in a population in southern Italy with cutaneous melanoma. PAX9 and MSX1 synergistically activate BMP4, and BMP4 is a gene crucial to tooth development as it regulates the passage from bud to cap stages. Since the unaffected parents displayed the same genetic variations of PITX2 and BMP4 in the homozygous state, it was deemed that these variations alone were unlikely to be the direct cause of the observed tooth

hypodontia; however, they may still play a role as risk factors or modulators of the phenotype.

Family 8 (a trio with unaffected parents) displayed a probable autosomic recessive form of inheritance. Unfortunately, we were unable to identify a strong candidate gene with a clear role in the physio-pathology of tooth formation and development. Based on RVIS score prioritization criteria, 6 genes emerged affected by functional mutation in this family and not expected to accumulate functional mutation by chance, namely PRR32, PLAC4, nuclear factor 1A (NFIA), tripartite motif containing 14 (TRIM14), carbohydrate sulfotransferase 8 (CHST8) and synaptic Ras GTPase activating protein 1 (SYNGAP1). Of these, 3 genes were unlikely to be involved in tooth agenesis: PLAC4 is mainly involved in placenta tissue formation (35); CHST8 has been associated with autosomal recessive peeling skin syndrome (36); SYNGAP1 is

Table V. Selected candidate genes and variants in family 8.

Chr	Ref	Alt	Func.refGene	Gene.refGene	GeneDetail.refGene	ExonicFunc.refGene
<i>Recessive</i>						
chrX	G	A	Exonic	ARSH	NA	Non-synonymous SNV
chrX	G	A	Exonic	PRR32	NA	Non-synonymous SNV
chr3	C	T	Exonic	TGM4	NA	Stopgain
chr21	-	40 bp	Exonic	PLAC4	NA	Frameshift insertion
chrX	G	A	Exonic	APEX2	NA	Non-synonymous SNV
<i>De novo</i>						
chr17	21 bp	-	Splicing	MYO19	NA	NA
chr16	T	G	Exonic	IFT140	NA	Non-synonymous SNV
chr1	A	C	Splicing	NFIA	NM_001145511:exon9:c.12312A>C, NM_001145512:exon10:c.1390-2A>C, NM_005595:exon9:c.12552A>C, NM_001134673:exon9:c.1255-2A>C	NA
chr11	-	T	Exonic	MUC6	NA	Frameshift insertion
chr6	G	A	Exonic	GPANK1	NA	Non-synonymous SNV
chr1	G	C	Splicing	NFIA	NM_001145511:exon9:c.1231-1G>C, NM_001145512:exon10:c.1390-1G>C, NM_005595:exon9:c.12551G>C, NM_001134673:exon9:c.1255-1G>C	NA
chr6	A	G	Exonic	HSPA1L	NA	Non-synonymous SNV
chr11	G	A	Exonic	MUC6	NA	Non-synonymous SNV
chr19	-	18 bp	Exonic	RSPH6A	NA	Non-frameshift insertion
chr9	C	G	Exonic	TRIM14	NA	Non-synonymous SNV
chr19	-	AGC	Exonic	CHST8	NA	Non-frameshift insertion
chr6	CACC ACCA CCAT	-	Exonic	SYNGAP1	NA	Non-frameshift insertion
chrX	C	T	Exonic	RBMX	NA	Non-synonymous SNV
chr11	G	T	Exonic	MUC6	NA	Non-synonymous SNV
chr11	A	G	Exonic	MUC6	NA	Non-synonymous SNV
chr11	G	-	Exonic	MUC6	NA	Frameshift deletion
chr1	G	-	Splicing	SEC22B	NM_004892:exon6:c.1066-1G>-	NA
chr3	C	G	Exonic	IL17RE	NA	Non-synonymous SNV
<i>Compound</i>						
chr2	-	CTGC	Exonic	DNAH7	NA	Frameshift insertion
chr2	A	G	Exonic	DNAH7	NA	Non-synonymous SNV
chr2	C	T	Exonic	DNAH7	NA	Non-synonymous SNV
chr11	-	T	Exonic	MUC6	NA	Frameshift insertion
chr11	G	A	Exonic	MUC6	NA	Non-synonymous SNV
chr11	-	TA	Exonic	MUC6	NA	Frameshift insertion
chr11	G	T	Exonic	MUC6	NA	Non-synonymous SNV
chr11	A	G	Exonic	MUC6	NA	Non-synonymous SNV
chr11	G	-	Exonic	MUC6	NA	Frameshift deletion
chr1	A	C	Splicing	NFIA	NM_001145511:exon9:c.1231-2A>C, NM_001145512:exon10:c.1390-2A>C, NM_005595:exon9:c.1255-2A>C, NM_001134673:exon9:c.1255-2A>C	NA

Table V. Continued.

Chr	Ref	Alt	Func.refGene	Gene.refGene	GeneDetail.refGene	ExonicFunc.refGene
chr1	G	C	Splicing	NFIA	NM_001145511:exon9:c.1231-1G>C, NM_001145512:exon10:c.1390-1G>C, NM_005595:exon9:c.1255-1G>C, NM_001134673:exon9:c.1255-1G>C	NA
chr10	A	T	Splicing	NRAP	NM_006175:exon38:c.4431+2T>A, NM_198060:exon39:c.4536+2T>A, NM_001261463:exon39:c.4536+2T>A	NA
chr10	T	G	Exonic	NRAP	NA	Nonsynonymous SNV
Known PITX2	UTR5	chr4	C	-	NM_000325:c.-387delG	

Table VI. Selected candidate genes and variants in family 9.

Chr	Ref	Alt	Func. refGene	Gene. refGene	GeneDetail. refGene	ExonicFunc. refGene	AAChange.refGene
<i>Compound</i>							
chr20	A	G	Exonic	ANKEF1	NA	Non-synonymous SNV	ANKEF1:NM_198798:exon2:c.A332G:p.D111G, ANKEF1:NM_022096:exon3:c.A332G:p.D111G
chr20	C	T	Exonic	COMMD7	NA	Non-synonymous SNV	COMMD7:NM_001099339:exon1:c.G5A:p.G2D, COMMD7:NM_053041:exon1:c.G5A:p.G2D
chr14	T	C	Exonic	DDHD1	NA	Non-synonymous SNV	DDHD1:NM_001160148:exon6:c.A1411G:p.I471V, DDHD1:NM_030637:exon6:c.A1411G:p.I471V, DDHD1:NM_001160147:exon7:c.A1432G:p.I478V
chr8	A	G	Exonic	EBF2	NA	Non-synonymous SNV	EBF2:NM_022659:exon7:c.T560C:p.L187S
chr17	G	T	Exonic	EVPL	NA	Non-synonymous SNV	EVPL:NM_001988:exon11:c.C1213A:p.L405M
chr3	C	T	Exonic	SUCLG2	NA	Non-synonymous SNV	SUCLG2:NM_001177599:exon10:c.G1124A:p.G375E, SUCLG2:NM_003848:exon10:c.G1124A:p.G375E
chr3	C	T	Exonic	SUCLG2	NA	Non-synonymous SNV	SUCLG2:NM_001177599:exon10:c.G1123A:p.G375R, SUCLG2:NM_003848:exon10:c.G1123A:p.G375R
chr17	T	C	Exonic	SUPT6H	NA	Non-synonymous SNV	SUPT6H:NM_003170:exon32:c.T4393C:p.C1465R
<i>De novo</i>							
chr3	C	T	Exonic	SUCLG2	NA	Non-synonymous SNV	SUCLG2:NM_001177599:exon10:c.G1124A:p.G375E, SUCLG2:NM_003848:exon10:c.G1124A:p.G375E
chr3	C	T	Exonic	SUCLG2	NA	Non-synonymous SNV	SUCLG2:NM_001177599:exon10:c.G1123A:p.G375R, SUCLG2:NM_003848:exon10:c.G1123A:p.G375R

responsible for non-syndromic mental retardation autosomal dominant form 5 (OMIM 612621) (37), a phenotype not reported in this family. The PRR32 gene is poorly characterized and difficult to evaluate. NFIA and TRIM14 are two transcription factor genes and they may represent good candidates for further studies aimed at assessing their involvement in the regulatory network driving teeth morphogenesis. NFIA

may be of particular interest, since mutations in this gene cause craniofacial abnormalities in mice and craniosynostosis in humans (38). However, homozygous null mutations in the gene have been associated with kidney, nervous and fertility phenotypes that were not observed in this family. Of note, this family was also a carrier of the genetic variant c.387delC/G in the 5'UTR of PITX2 (found in the homozygous state in the

proband and in his unaffected brother and in the heterozygous state in their unaffected mother in family 5; discussed above), suggesting that a complex interplay of genetic risk factors may be involved in this disease.

Finally, family 9 included a pair of monozygotic twins with hypodontia and the recessive inheritance model was likely. Unfortunately, DNA for the father was not available for the study. Among the genetic variants found, only the one in *COMMD7* may have a link with the disease. *COMMD7* is in fact a NEMO interacting protein involved in the termination of NF- $\kappa$ B signaling that is activated by the EDA-EDAR-EDARADD complex (39). This mutation is predicted to be damaging with a score of 1 by PolyPhen-2. Further studies are necessary to establish whether this heterozygous mutation in the proband twin may be involved in the pathological phenotype.

In conclusion, in this study, using WES analysis, we identified the causative mutations in cases with a more severe phenotype (families 1 and 7, where 17 and 16 teeth were missing, respectively). We did not immediately identify variants that may be disease-causing for families with a light phenotype, such as families 5 and 9 (maximum 4 missing teeth) and with a mild oligodontia, such as family 8 (6 missing teeth). However, our analysis of these families highlighted some interesting candidate genes that may be considered targets of future functional studies. However, we cannot exclude that the manifestation of the phenotypic traits may also depend or be modulated by other mechanisms, including long and small non-coding RNAs (i.e., miRNAs), epigenetic modifications, such as DNA methylation and histone modifications or regulatory DNA variants. Finally, since a complex regulatory network contributes to tooth formation with multitudes of genes potentially involved, we consider that WES may be an effective strategy with which to detect the genetic defects related to tooth agenesis, particularly in severe cases of oligodontia in sporadic and familial cases. The determination of the genetic causes of tooth agenesis is important for genetic counseling and for anticipating the problems related to the clinical management of dental anomalies, in order to ensure a correct occlusion, particularly during developing dentition in children.

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