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ARTICLE

Identification of causative variants in *TXNL4A* in Burn-McKeown syndrome and isolated choanal atresia

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Burn-McKeown syndrome (BMKS) is a rare syndrome characterized by choanal atresia, prominent ears, abnormalities of the outer third of the lower eyelid, structural cardiac abnormalities, conductive and sensorineural hearing loss, and cleft lip. Recently, causative compound heterozygous variants were identified in TXNL4A. We analyzed an individual with clinical features of BMKS and her parents by whole-genome sequencing and identified compound heterozygous variants in TXNL4A (a novel splice site variant (c.258-2A > G, (p.?)) and a 34 bp promoter deletion (hg19 chr18:g.77748581_77748614del (type 1Δ) in the proband). Subsequently, we tested a cohort of 19 individuals with (mild) features of BMKS and 17 individuals with isolated choanal atresia for causative variants in TXNL4A by dideoxy-sequence analysis. In one individual with BMKS unrelated to the first family, we identified the identical compound heterozygous variants. In an individual with isolated choanal atresia, we found homozygosity for the same type 1Δ promoter deletion, whilst in two cousins from a family with choanal atresia and other minor anomalies we found homozygosity for a different deletion within the promoter (hg19 chr18: g.77748604_77748637del (type 2Δ)). Hence, we identified causative recessive variants in TXNL4A in two individuals with BMKS as well as in three individuals (from two families) with isolated choanal atresia.

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INTRODUCTION

Burn-McKeown syndrome (BMKS) was first described in five children by Burn *et al.*¹ It is a rare disorder characterized by choanal atresia, prominent ears, hypertelorism with short palpebral fissures and abnormalities of the outer third of the lower eyelids. Other features that can be observed are structural cardiac abnormalities, conductive and sensorineural hearing loss, and unilateral cleft lip.¹ There is clinical overlap with both Treacher Collins and CHARGE syndromes.

Burn *et al*¹ advised counseling a high recurrence risk in families with BMKS, but the inheritance pattern remained unclear until recently. As the syndrome was repeatedly described in siblings and mainly in males, an autosomal recessive¹ or X-linked inheritance pattern was suggested.² However, the identification of a chromosomal rearrangement, 46,XX,r(18)(p14q23), in an isolated female case with features of BMKS suggested terminal 18p or 18q as the locus for the disorder.¹ This was borne out by the detection of causative compound heterozygous variants in *TXNL4A* (located on chromosome 18q23), identified more than 20 years after the initial description of BMKS.³

TXNL4A is a member of the U5 spliceosomal complex that is critical for pre-mRNA splicing.⁴ It has been suggested that reduced

expression of *TXNL4A* influences the splicing of a specific subset of pre-mRNAs, resulting in the tissue-specific phenotypic spectrum of BMKS.^{3,5} In the present study, we describe new causative variants in *TXNL4A* and expand the associated phenotypic spectrum.

SUBJECTS AND METHODS

Subjects

Ethical approval from the board of the Medical Ethical Committee of the Erasmus MC, University Medical Center Rotterdam, the Netherlands, was given for whole-genome sequencing (WGS; MEC-2012-140) and for retrospective data collection (MEC-2013-547). UK samples were collected following approval from the Great Ormond Street Hospital for Children NHS Trust Ethics Committee (REC No. 08H0713/46). Informed consent was received from the individuals and parents.

WGS was performed on DNA of the members of family 1 (see Figure 1). To identify further mutation-positive individuals, dideoxy-sequence analysis of the promoter region, the exons and the exon/intron boundaries of *TXNL4A* was performed using DNA of 19 individuals who have clinical features that overlap with (mild) BMKS. In addition, 17 individuals with isolated choanal atresia were also tested for mutations in *TXNL4A*. In addition, 29 of the DNA samples were analyzed for copy number variations in *TXNL4A* using a

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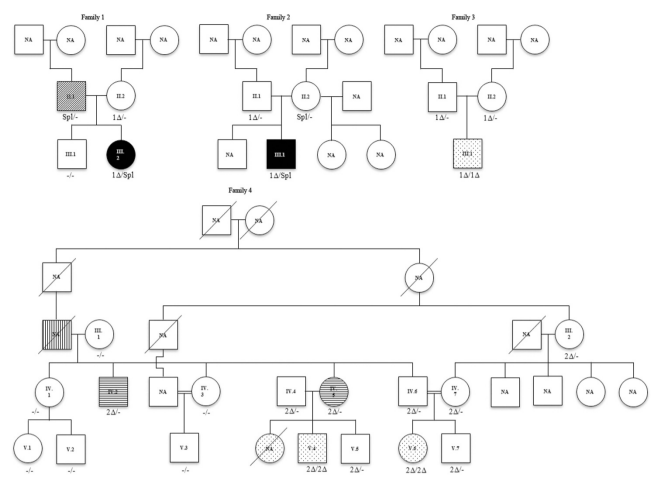


Figure 1 Pedigrees of families. Symbol definitions: \square onta affected, $\blacksquare \bullet$ BMKS, $\square \otimes$ lagophtalmos, $\square \otimes$ choanal atresia, $\blacksquare \oplus$ prognathism and $\square \otimes \oplus$ maxillary hypoplasia. 1Δ , promoter deletion type 1Δ (hg19 chr18:g.77748581_77748614del), 2Δ , promoter deletion type 2Δ (hg19 chr18:g.77748604_77748637del); NA, not available; SpI, splice site variant, -/-, no causative variant in TXNL4A.

multiplex ligation-dependent probe amplification (MLPA) assay as described by Wieczorek $\it et al.^3$

The variants identified were described according to HGVS nomenclature, ⁶ using the reference sequences NM_006701.2, ENST00000269601 and ENSG00000141759, on GRCh37/hg19 and were submitted to the Leiden Open Variation Database (http://www.lovd.nl/TXNL4A).

Whole-genome sequencing

Before publication of the paper of Wieczorek *et al*,³ WGS was performed on DNA of the proband of family 1 (III.2) and her parents (see Figure 1) by Complete Genomics, a BGI company (Mountain View, CA, USA) as described by Drmanac.⁷ Variants were annotated using NCBI build 36.3/hg18 and dbSNP build 130. Data were analyzed using cga tools version 1.6.0.43 and TIBCO Spotfire 7.0.0 (TIBCO Software Inc., Boston, MA, USA). The annotations were converted to GRCh37/hg19 by using Human BLAT search on the UCSC website (Kent Informatics, Inc., Santa Cruz, CA, USA) as described previously.⁸

An autosomal dominant disease model was tested in family 1. The analysis was restricted to novel non-synonymous variants, variants disrupting a splice site (\pm two bp), and insertions or deletions in the coding sequence (\pm 50 bp).

The remaining variants were analyzed with Annovar, 9 to get an indication of the pathogenicity and the ESP frequency as given in Exome Variant Server (EVS, NHLBI GO Exome Sequencing Project (ESP), Seattle, WA (URL: http://evs.gs.washington.edu/EVS/) [April 2014 accessed]). We focused on variants that were located in genes without loss-of-function mutations in EVS.

Confirmation by dideoxy-sequence analysis

The variants identified by WGS were validated by dideoxy-sequence analysis (all primer sequences are provided in Supplementary Table 1). Amplification reactions were performed according to standard procedures. PCR products were purified using Agencourt AMPure (Agencourt, Beckman Coulter Inc., Brea, CA, USA). Direct sequencing of both strands was performed using Big Dye terminator version 3.1 (Applied Biosystems, Foster City, CA, USA) as recommended by the manufacturer. Dye terminators were removed using Agencourt CleanSeQ (Agencourt) and loaded on an ABI 3130XL Genetic Analyzer (Applied Biosystems). The sequences were analyzed using SeqPilot version 4.1.2 build 507 (JSI Medical Systems Gmbh, Kippenheim, Germany).

Confirmation of effect of the variants on RNA expression

To analyze the effect of the variants on RNA expression, cDNA was analyzed by restriction enzyme digestion and deep sequencing. First, RNA was extracted according to the manufacturer's protocol from venous blood collected into PAXgene Blood RNA tubes (Qiagen N.V., Venlo, The Netherlands) from individuals II.1, II.2 and III.2 (Figure 1, family 1). cDNA was synthesized using the RevertAid First Strand cDNA kit (Thermo Scientific Inc., Waltham, MA, USA), with random hexamer primers according to the manufacturer's instructions. Primers were designed for all exons and intron two to screen for truncations and intron retention (Supplementary Table 1). cDNA was amplified and products were electrophoresed on agarose gels and DNA was visualized by staining with ethidium bromide (EtBr).

Digestion with PshAI and AhdI and electrophoresis of digests

To assess the effect of the paternal variant on RNA expression in family 1, cDNA amplification products were digested with PshAI or AhdI (New England Biolabs Inc., Ipswich, MA, USA) and analyzed by agarose gel electrophoresis. The forward primer was mutated to allow digestion with AhdI (underlined and bold 'a' in Supplementary Table 1).

Deep sequencing by Ion PGM Sequencing

To distinguish between the wild-type and mutant allele, two heterozygous SNPs in the 3'-UTR of TXNL4A that were identified in the WGS data from the parents of the family 1 proband were used. To quantify the effect of the maternal variant, deep sequencing covering the SNP was performed on cDNA of the proband. Primers are given in Supplementary Table 1. The cDNA amplification products were diluted 1/100 and 2 µl was used in a second round PCR, with a reverse primer including the Ion PGM P1 adapter, and a forward primer with the Ion PGM A adapter sequence and a barcode sequence (reaction conditions available on request). Amplification products (roughly equal amounts as judged by EtBr staining by agarose gel electrophoresis) were combined and then purified with AMPure beads (Beckman Coulter). Emulsion PCR and enrichment was performed with the Ion PGM Template OT2 200 Kit (Life Technologies) according to the manufacturer's instructions. Sequencing of enriched templates was performed on the Ion Torrent PGM (Life Technologies) for 125 cycles using the Ion PGM Sequencing 200 kit v2 and Ion 316 chips. Data were processed with Ion Torrent platform-specific pipeline software v4.2.1.

RESULTS

Whole-genome sequencing of family 1

The proband of family 1 and her parents were analyzed by WGS. Variants remaining after the various filtering steps are available on request. By testing the expected autosomal dominant disease model, we identified a novel splice site variant upstream of the last exon in TXNL4A (c.258-2A>G, (p.?)), that was inherited from the father. According to both cga tools and ANNOVAR, the splice site variant was predicted to be deleterious. As this variant was present in the 3' splice site consensus sequence, it was highly conserved with a PhyloP score of 1.824×10^{-5} . The variant was not present in ESP, 1000 Genomes (The 1000 Genomes Project Consortium, 2012, URL: http://www.1000genomes.org/10), Ensembl,11 nor in our in-house Huvariome database.12 However, it did have an allele frequency of 7.31×10^{-6} in gnomAD.13 During the analysis, the phenotype was recognized as suspected BMKS. Therefore, recessive variants on

chromosome 18 were implicated and the DNA sequence of *TXNL4A* was scrutinized manually. Based on the publication of Wieczorek *et al*,³ the likely significance of a 34 bp deletion within the promoter became apparent (hg19 chr18:g.77748581_77748614del, referred to as type 1Δ and inherited from the mother).³ The presence of both variants was confirmed by dideoxy sequencing (Figure 2).

cDNA analysis of TXNL4A

RT-PCR analysis of the proband of family 1 and her parents, followed by agarose gel electrophoresis, did not show any aberrantly spliced products in any of the samples, despite the presence of the predicted splice site variant in both the proband and her father. WGS data indicated that both parents each carried a different sequence change within the 3' UTR of *TXNL4A* that was not inherited by their daughter (thus located on the reference alleles): the father was heterozygous for hg19 chr18:g.77733297dupC (rs77355432, dbSNP build 146) and the mother for hg19 chr18:g.77733273C>T (rs4798931). Primers were designed to amplify both SNPs in one amplicon.

The effect of the 3' acceptor variant c.258-2A > G on splicing, was assessed using the restriction enzymes AhdI and PshAI. AhdI was specific for the 3' UTR dupC (reference allele, not passed to the proband), while PshAI was specific for the mutant allele in cDNA from the father (passed to the proband). The primer was designed to amplify the alleles equally, had they both been present in the cDNA. Using AhdI, the cDNA of the father was digested almost to completion (Figure 3), and using PshAI the cDNA remained undigested (data not shown), indicating that there was no or extremely reduced mature transcript from the mutant allele.

Deep sequencing of maternal cDNA demonstrated reduced expression from the allele with the type 1Δ : in a total of 36 924 reads, 8 636 reads (23%) were of the reference C (the allele with the promoter variant), and 28 283 reads (77%) contained the variant T (wildtype promoter allele).

Expansion of the mutation spectrum

Dideoxy sequencing of *TXNL4A* was performed on DNA obtained from 19 index cases with (mild) features of BMKS. The proband of family 2 (Figure 1, III.1 of family 2) was found to carry the identical variants to those of the family 1 proband. Analysis of parental samples showed that they were present in compound heterozygous state with

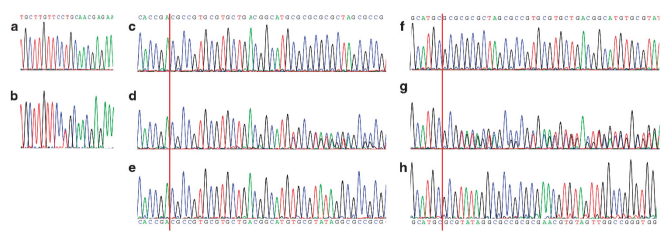


Figure 2 Electropherograms of dideoxy-sequence analyses. Red line indicates the start of the deletion. (a) control DNA. (b) splice site variant in DNA of III.2 of family 1. (c) region of type 1Δ deletion in control DNA. (d) heterozygous type 1Δ deletion in DNA of III.2 of family 1. (e) homozygous type 1Δ deletion as seen in DNA of III.1 of family 3. (f) region of type 2Δ deletion in control DNA. (g) heterozygous type 2Δ deletion as seen in DNA of IV.6 of family 4. (h) homozygous type 2Δ deletion as seen in DNA of V.4 of family 4.

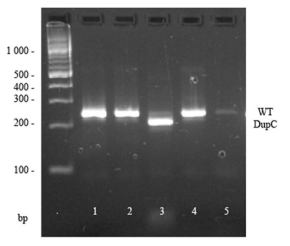


Figure 3 Restriction enzyme digest analysis of rs77355432 SNP in cDNA from family 1; expected fragment sizes reference allele 249 bp (WT), alternative allele 215+34 bp (dupC). Lane 1, uncut DNA. Lane 2, cDNA from the proband is not digested with Ahdl. Lane 3, cDNA of the father is cut almost to completion with Ahdl. Lane 4, cDNA of the mother is uncut with Ahdl. Lane 5, negative control.

the type 1Δ deletion inherited from the father and the splice site variant inherited from the mother. In addition, 17 index cases with choanal atresia were tested for mutations in *TXNL4A*. In two probands, homozygosity for a promoter deletion was identified. The proband of family 3 (Figure 1, III.1 of family 3) had a uniparental disomy of chromosome 18p11.32q23, leading to homozygosity for the type 1Δ deletion. The family 4 proband (Figure 1, V.4 of family 4) was homozygous for a slightly different promoter deletion: hg19 chr18: g.77748604_77748637del corresponding to type 2Δ in Wieczorek *et al.*³ The same genotype was also present in his first cousin (Figure 1, V.6 of family 4).

MLPA analysis was performed on the 29 samples with sufficient DNA available. This included samples of both families 3 and 4 (Table 2). Large rearrangements were not identified in any of these samples.

Subjects

Nineteen individuals with clinical features overlapping with BMKS syndrome and 17 individuals with isolated choanal atresia were tested for mutations in *TXNL4A*. Causative variants were identified in five individuals. An overview of the clinical phenotype of these individuals is given in Table 1. The proband of family 4 and his first cousin (Figure 1, V.4 and V.6 of family 4) were previously described as patients 1 and 3 in the paper of Ramos-Arroyo *et al.*¹⁴ All individuals had choanal atresia and normal development. In three out of five, their pregnancy was complicated by polyhydramnios. The same three had a prominent nasal bridge. Maxillary hypoplasia was also seen in the same three individuals, and micrognathia was seen in two out of them. Two out of five individuals had defects of the lower eyelids. The proband of family 4 and his first cousin had oligodontia, however, due to the high prevalence of this condition, ¹⁵ we cannot rule out that this is a separate condition that is segregating within this family. ¹⁴

DISCUSSION

In two out of 20 unrelated individuals with clinical features of BMKS, we identified compound heterozygosity in *TXNL4A* for a novel splice site variant, c.258-2A>G, (p.?) and the type 1Δ promoter variant, hg19 chr18:g.77748581_77748614del. Our findings support the idea

that causative TXNL4A variants do not completely ablate function in BMKS 3

In nine out of 11 affected families, Wieczorek *et al*³ found a rare loss-of-function variant (nonsense, frameshift or microdeletion) on one allele in conjunction with a promoter deletion on the other allele. In our deep sequencing analysis of RNA isolated from blood, expression of the type 1Δ mutant allele was reduced by ~69%. This is in keeping with a luciferase reporter gene assay, where the promoter activity of constructs containing type 1Δ deletion was reduced by 59% compared to a wild-type construct.³ The presence of the splice-site variant on the other *TXNL4A* allele of the family 1 proband most likely leads to loss-of-function, since analysis of paternal cDNA based on the closely adjacent SNP rs77355432, indicated no evidence of transcription of this allele (Figure 3). Wieczorek *et al*³ have shown that deletion of exon three combined with type 1Δ causes BMKS. Hence, we can conclude that the variants identified in the probands of families 1 and 2 are the underlying causes of the clinical phenotype.

In the paper of Wieczorek et al^3 the allele frequency of the promoter deletions in German control samples was estimated at 0.76%. A single homozygous type 1∆ deletion was identified in 3343 population-based German and South Asian samples. Unfortunately, data about the craniofacial phenotype in that individual were not available. In this study, we sequenced a cohort of 17 individuals with choanal atresia and also identified a homozygous type 1Δ deletion (family 3). Although the predicted frequency of homozygous type 1∆ deletions was estimated at 1:17 300, which seemed to exclude this genotype as a cause for BMKS, Wieczorek et al3 stated that this genotype might lead to a mild phenotype. It seems likely that the isolated bilateral choanal atresia identified in the proband of family 3 might be part of this mild phenotype, especially since choanal atresia is one of the major anomalies of BMKS. As isolated choanal atresia is a feature of the Tbx22 knock-out mouse, 16 the individuals with isolated choanal atresia were screened for mutations in TBX22, but causative variants were not found.

In family 4, homozygosity for the pathogenic promoter variant type 2Δ was found to segregate in the affected individuals. Wieczorek et al^3 also found a homozygous type 2Δ case in an individual with true BMKS and showed that promoter activity of a construct containing type 2Δ was reduced by 72% compared to wild-type, indicating a stronger reduction of TXNL4A expression than type 1Δ . However, individuals with the type 2Δ (family 4) seem to be less severely affected than the compound heterozygotes; only choanal atresia, hypodontia and some minor anomalies were seen (Table 2). Further screening of family 4 identified eight members who were heterozygous for type 2Δ , and interestingly minor anomalies were present in two of them (Figure 1 and Table 2). Our data support the proposed dosage-specific effect.^{3,5}

Wieczorek et al³ identified causative variants in nine out of 11 families, compared to two out of 18 in this study. This may be attributed to the fact that in this work individuals were included with only mild features of BMKS and may overlap with other conditions such as bilateral craniofacial microsomia, CHARGE phenotype, and asymmetrical Treacher-Collins like phenotype. Recently, it has been shown that ribosomopathies, such as Treacher Collins syndrome, and spliceosomopathies can have similar craniofacial phenotypes.⁵ In contrast to CHARGE and Treacher Collins syndrome, individuals with BMKS tend to have a normal intellectual development, however, there is otherwise significant clinical overlap. Therefore, we suggest it will be important to test for mutations in TXNL4A if the clinical phenotype is indicative of BMKS, but also if the clinical phenotype is

Table 1 Clinical phenotype of individuals with causative variants using the features mentioned by Wieczorek et al³ that are associated with BMKS

Clinical phenotype of indiv	Clinical phenotype of individuals with causative variants				
ID in pedigree	III.2 family 1	III.1 family 2	III.1 family 3	V.4 family 4	V.6 family 4
Gender	Female	Male	Male	Male 7 years	Female
Genetic testing	JV years TCOF1	Affymetrix 750 K microarray	Affymetrix 750 K microarray -> 18p11.32q23	Karyo	TXNL4A
	СНD7	TXNL4A	(136,300-77,997,392) IIIIZ uniparental disorny for chromosome 18	TXNL4A	MLPA
	SNP array	MLPA	TXNL4A	MLPA	
	WGS		MLPA		
Variant carried by father	MLPA c.258-2A>G, (p.?)	hg19 chr18:	hg19 chr18:g.77748581_77748614del (type 1Δ)	hg19 chr18: g.77748604_77748637del	hg19 chr18: g.77748604_77748637del
Variant carried by mother	hg19 chr18:	g.77748581_77748614del (type 1Δ) c.258-2A > G, (p.?)	hg19 chr18:g.77748581_77748614del (type 1Δ)	(type 2Δ) hg19 chr18: g.77748604_77748637del	(type 2A) hg19 chr18: g.77748604_77748637del
Positive family history Normal pregnancy	g.////48581_///48514del (type 1Δ) Y father scleral show Y	N Polyhydramnios	N IVF ICSI	(type Z∆) Y Polyhydramnios	(type ZA <i>)</i> Y Polyhydramnios
			Failure to progress		
Asymmetry of the face Hypertelorism	>- Z			ZZ	zz
Short palpebral fissures Defect of lower eyelids	>- >-	Y (B)		zz	zz
Aplasia of puncta	Y (L)				
Prominent nasal bridge	> :	≻		≻ :	> :
Short philtrum Thin lips	Z >			>- >-	> Z
Cleft lip or palate	. Z 2	Y (L)		· z ;	: z ;
Bifid uvula Bilateral choanal atresia/	N Bilateral (bony)	Bilateral	Bilateral (bony)	lateral (bony)	N Bilateral
stenosis	:			:	
Prominent ears Preauricular tags	Y Y (R)	Y (R)		>-	>-
Maxillary hypoplasia	Y (R)	:		>- 3	>
Micrognathia Cardiac defect	(X) ~ Z	Y Asymptomatic ASD and VSD		Z	
Hearing loss	Z >	>		Z>	z,
development	-	_		-	-
Short stature Other	Y Hypoplasia infra-orbital rim (R)	Dimple on the cheek		N Downslanting palpebral fissures	N Cowlick
	Upslanting palpebral fissures	Hypermetropia		Prognathism	Downslanting palpebral fissures
	Eyelashes were longer laterally than			High arched narrow palate	Prognathism
				Absence of the upper and lower deciduous	Dental malocclusion
	Choroid coloboma (L)			premolars	Narrow palate
	Microstrabismus			Hypodontia of four permanent premolars	Inilateral absence of the parmenent unar
	Amblyopia				lateral incisor

Abbreviations: ASD, atrial septal defect; B, bilateral; Hom, homozygous; ICSI, intracytoplasmic sperm injection; IVF, in vitro fertilization; Karyo, karyotype; L, left; M, maternal; N, no; P, paternal; R, right; VSD, ventricular septal defect; WGS, whole-genome sequencing; Y, yes. Blank entries indicate that information was not available.

Table 2 Overview of testing done for 41 affected and 19 unaffected individuals

							BMK S/isolated
Family	Sample	Gender	Genetic testing	Variants identified	MLPA	Clinical features	choanal atresia
Family 1	III.2	ட	<i>TCOF1, CHD7</i> , SNP array, WGS	hg19 chr18.g.77748581_77748614del (1∆) (M), c.258-2A>G, (p.?) (P)	>-	Normal pregnancy, facial asymmetry, short palpebral fissures, defect of lower eyelids, aplasia of puncta lacrimalis L, hypoplasia infra-orbital rim R, upslanting palpebral fissures, longer eyelashes laterally than medially, chronid coldoma L, microstrabismus, ambiopa bilateral choanal atresia, preauricular tag R, maxillary hypoplasia R, micrograpthia R and normal psychomotor development	BMKS
	11.1	Σι⊥	WGS WGS	c.258-2A>G, (p.?) hg19 chr18:g.77748581_77748614del (1A)	> >	Scleral show	
	Uncle Husband	≥≥	TXNL4A TXNL4A		> >		
Family 2	III.1	Σ	Affymetrix 750 K microarray, TXNL4A	hg19 chr18;g.77748581_77748614del (1∆) (P), c.258-2A>G, (p.?) (M)	· > -	Polyhydramnios, bilateral defect of lower eyelids, hypermetropia, prominent nasal bridge, cleft lip and palate L, bilateral choanal atresia, preauricular tag R, micrograthia, normal psychomotor development, dimple on the cheek, asymptomatic ASD and VSD	BMKS
	11.2	цΣ	TXNL4A TXNL4A	c.258-2A>G, (p.?) hg19 chr18:g.77748581_77748614del (1A) (het)	zz		
Family 3	II:1	Σ	Affymetrix 750 K microarray -> isodisomy of chr18 TXNL4A	hg19 chr18:g.77748581_77748614del (1A) (hom)	>-	IVF ICSI, failure to progress and bilateral choanal atresia	Choanal atresia
	11.2	Ŀ	TXNL4A	hg19 chr18:g.77748581_77748614del (1∆) (het)	z		
	1.1	Σ	TXNL4A	hg19 chr18:g.77748581_77748614del (1A) (het)	z		
Family 4	4.7	Σ	TXNL4A, TBX22	hg19 chr18: g.77748604_77748637del (2.Δ) (hom)	>-	Polyhydramnios, downslanting palpebral fissures, prominent nasal bridge, bilateral choanal atresia, maxillary hypoplasia, prognathism, high arched narrow palate, absence of the upper and lower deciduous premolars, hypodontia of four permanent bremolars and normal psychomotor development.	Choanal atresia
	9.	ш	Karyo, <i>TXNL4A</i> , <i>TBX22</i>	c245212del34, (p.?) (2A) (hom)	>-	Polyhydramnios, prominent nasal bridge, bilateral choanal atresia, maxillary hypoplasia, cowlick, downslanting palpebral fissures, prognathism, dental malocclusion, narow palate and unilateral absence of the permanent upper lateral incisor	Choanal atresia
	11.1	Ŀ	TXNL4A		z		
	III.2	L L	TXNL4A	hg19 chr18: g.77748604_77748637del (2 Δ) (het)	zz		
	IV.1	⊾≥	TXNL4A	hg19 chr18: g.77748604_77748637del	zz	Prognathism	
	o ≥	Ц	TV NI A	(2∆) (het)	Z	,	
	V. V.	. ≥	TXNL4A	hg19 chr18: g.77748604_77748637del	zz		
	17.5	Σ	TXNL4A	(2A) (het) hg19 chr18: g.77748604_77748637del	z	Prognathism	
	<u>ل</u> ا	Σ	TXN/ 74	(2A) (het)	z		
	0.	Ξ	V+74/V	(2 Δ) (het)	_		
	17.7	ட	TXNL4A	hg19 chr18: g.77748604_77748637del (2∆) (het)	z		
	V.1	ь :	TXNL4A		z		
	7.7	≥ ≥	1XNL4A TXN1.48		z z		
	7.5	Σ	TXNL4A	hg19 chr18: g.77748604_77748637del	zz		
	٧.7	Σ	TXNL4A	(2.Δ.) (het) hg19 chr18: g.77748604_77748637del (2.Δ.) (het)	z		

Table 2 (Continued)

choanal atresia	BMKS		BMKS		BMKS			Choanal atresia	BMKS				BMKS	3	DIVINO	BMKS			BMKS		BMKS		BMKS		BMKS	BMKS	Choanal atresia	-		Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	Choanal atresia	BMKS	BMKS	BMKS	BMKS	BMKS
Clinical features	Ptosis, shallow orbits, prominent beaked nose and micrognathia		Polyhydramnios, premature, respiratory distress, macroglossia, micrognathia, retrognathia, cleff palate, bilateral microtia, preauricular fistula, short down-	slanting palpebral fissures, shallow orbits and hearing loss	Small for gestational age, conductive hearing loss, bilateral ear tags, down-	slanting palpebral fissures, hypertelorism, micrognathia, perianal tag and patent	foramen ovale	Bilateral choanal atresia	Bilateral Tessier 7 cleft, dental crowding, bilateral ear tags, hearing loss based on	absent cochlear nerves, coloboma of the left papill, retina and pupil, amblyopia,	esophageal atresia with tracheoesophageal fistula, hemivertebrae, hypoplastic	thumbs L > K, bilateral hemitacial macrosomia, hypotonia, USA and mild PMR	Micrognathia, retrognathia, narrow maxilla, bilateral ear tags, hearing loss,	epibulibar dermold (K), dental crowding and USA	Asymment of the face, fleaming loss, normal stature, dysplastic ears, amblyopia	and abeliant racial iterve. Asymmetry of the face hypertelorism short palpebral fissures prominent pasal	hidge cleft lip and palate, microtia, maxillary hypoplasia, micrographia, hearing	loss, normal psychomotor development and short stature	Unilateral cleft lip and palate, maxillary hypoplasia, micrognathia and normal	psychomotor development	Renal cysts and microtia (unilateral)		Bilateral preauricular tags, bilateral Tessier 7 cleft, microtia R and anal atresia		Maxillary hypoplasia, micrognathia, microtia R and hypoplasia zygomata (mild)	Maxillary hypoplasia, micrognathia and unilateral hearing loss	Hypertelorism, hypoplastic uvula, right sided choanal atresia, Mum Carbimazole	treatment, blue eyes and ear pit (cocnlear implant)	Choanal atresia R	Choanal atresia B	Choanal atresia R	Birid uvula, choanal atresia K, ASD	Choanal atresia R	Choanal atresia R		CHARGE					Choanal atresia L, ID and syngnathia					
MLPA	>-	>-	>		>			>-	>-			:	>-	>	-	>	-		>-		>		>-	:	>- :	>-	>-	>	>- :	>- 1	>- :	> - ;	> - i	Z	Z	Z	Z	Z	Z	Z	Z	Z	Z	Z	z	Z
Variants identified																deletion of POI R1D																				R157X in CHD7										
Genetic testing	WGS, TXNL4A	WGS, TXNL4A	WGS, TXNL4A, TCOFI, POLRID. POLRIC.	POLRIA, array	TXNL4A, array, TCOF1,	SALLI, POLRIC, POLRID		TXNL4A, CHD7	TXNL4A			;	Karyo, <i>TXNL4A</i>	CO C	All ICS gelles, WES, BOR	genes TCOF1 POLR1C POLR1D	WFS. TXNI 4A		all TCS genes, TXNL4A		TCOF1, array, BOR syn-	drome genes, TXNL4A	TCOF1, SNP array,	FULKIC, FULKIU, IXINL4A	TCOF1, TXNL4A	TCOF1, TXNL4A	TXNL4A, TBX22	00% CH		TXNL4A, TBX22	TXNL4A, TBX22	IXNL4A, IBXZZ	IXNL4A, IBXZZ	IXNL4A, IBX22	TXNL4A, TBX22	CHD7, TXNL4A, TBX22		TXNL4A, TBX22	TXNL4A, TBX22	TXNL4A, TBX22	TXNL4A, TBX22	TXNL4A	TXNL4A	TXNL4A	TXNL4A	TXNL4A
Gender	≥	L	≥		Σ			ட	ட				Σ	L	L	ш	-		Σ		∑		L	:	∑ :	Σ																	L	∑	Σ	Σ
Sample	11.1	I.2 (mother)	9		7			œ	6				10	:	11	12	1		13		14		15	;	16	17	18		61	20	21	77	23	24	25	26	27	28	59	30	31	32	33	34	35	36
Family	Family 5																																													

Abbeviations: ASD, arial septal defect; B. bilateral; f, female; Hom, homozygous; ICSI, intracytoplasmic sperm injection; ID, intellectual disability; IVF, in vitro fertilization; karyogram; L, left; m, male; M, maternal; N, no; OSA, obstructive sleep apnea; P, paternal; PMR, psychomotor retardation; R, right; VSD, ventricular septal defect; WES, whole-exome sequencing; WGS, whole-genome sequencing; Y, yes.

indicative of these other conditions should their appropriate gene tests report negative.

CONCLUSION

We describe the finding of the splice site variant c.258-2A>G, combined with promoter deletion type 1Δ , in *TXNL4A* as the genetic cause of BMKS in two unrelated individuals. Homozygosity for the type 1Δ deletion was identified in a third individual with isolated choanal atresia. Homozygosity for the type 2Δ deletion was identified in a family with choanal atresia and other minor anomalies. These results confirm that variants affecting function of *TXNL4A* are the cause of BMKS and possibly a cause of isolated choanal atresia, underlining the tissue-specific nature of craniofacial disorders caused by spliceosomal defects.⁵

CONFLICT OF INTEREST

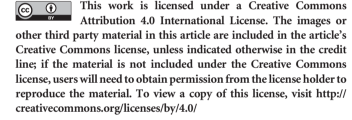
The authors declare no conflict of interest.

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