



HAL
open science

Whole exome sequencing identifies new causative mutations in Tunisian families with non-syndromic deafness.

Zied Riahi, Crystel Bonnet, Rim Zainine, Malek Louha, Yosra Bouyacoub, Nadia Laroussi, Mariem Chargui, Rym Kefi, Laurence Jonard, Imen Dorboz, et al.

► **To cite this version:**

Zied Riahi, Crystel Bonnet, Rim Zainine, Malek Louha, Yosra Bouyacoub, et al.. Whole exome sequencing identifies new causative mutations in Tunisian families with non-syndromic deafness.. PLoS ONE, 2014, 9 (6), pp.e99797. 10.1371/journal.pone.0099797 . pasteur-01060192

HAL Id: pasteur-01060192

<https://riip.hal.science/pasteur-01060192>

Submitted on 3 Sep 2014

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Whole Exome Sequencing Identifies New Causative Mutations in Tunisian Families with Non-Syndromic Deafness

Zied Riahi^{1,2,3}, Crystel Bonnet^{3,9}, Rim Zainine⁴, Malek Louha⁵, Yosra Bouyacoub¹, Nadia Laroussi^{1,2}, Mariem Chargui¹, Rym Kefi¹, Laurence Jonard⁵, Imen Dorboz⁶, Jean-Pierre Hardelin⁷, Sihem Belhaj Salah⁴, Jacqueline Levilliers⁷, Dominique Weil⁷, Kenneth McElreavey⁸, Odile Tanguy Boespflug⁶, Ghazi Besbes⁴, Sonia Abdelhak^{1,2*}, Christine Petit^{3,7}

1 Laboratoire de Génomique Biomédicale et Oncogénétique, Institut Pasteur de Tunis, Tunis, Tunisia, **2** Faculté des Sciences de Tunis, Université de Tunis El Manar, Tunis, Tunisia, **3** INSERM UMRS 1120, Institut de la Vision, Paris, France, **4** Service d'ORL et de Chirurgie Maxillo-faciale, CHU La Rabta, Tunis, Tunisia, **5** Centre de Référence des Surdités Génétiques, Hôpital Trousseau- APHP, Paris, France, **6** INSERM U931, Clermont Ferrand, France, **7** Unité de Génétique et Physiologie de l'Audition, Institut Pasteur de Paris, Paris, France, **8** Unité de Génétique du Développement Humain, Institut Pasteur de Paris, Paris, France

Abstract

Identification of the causative mutations in patients affected by autosomal recessive non syndromic deafness (DFNB forms), is demanding due to genetic heterogeneity. After the exclusion of *GJB2* mutations and other mutations previously reported in Tunisian deaf patients, we performed whole exome sequencing in patients affected with severe to profound deafness, from four unrelated consanguineous Tunisian families. Four biallelic non previously reported mutations were identified in three different genes: a nonsense mutation, c.208C>T (p.R70X), in *LRTOMT*, a missense mutation, c.5417T>C (p.L1806P), in *MYO15A* and two splice site mutations, c.7395+3G>A, and c.2260+2T>A, in *MYO15A* and *TMC1* respectively. We thereby provide evidence that whole exome sequencing is a powerful, cost-effective screening tool to identify mutations causing recessive deafness in consanguineous families.

Citation: Riahi Z, Bonnet C, Zainine R, Louha M, Bouyacoub Y, et al. (2014) Whole Exome Sequencing Identifies New Causative Mutations in Tunisian Families with Non-Syndromic Deafness. PLoS ONE 9(6): e99797. doi:10.1371/journal.pone.0099797

Editor: Osman El-Maarri, University of Bonn, Institut of experimental hematology and transfusion medicine, Germany

Received: February 3, 2014; **Accepted:** May 17, 2014; **Published:** June 13, 2014

Copyright: © 2014 Riahi et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported by the Tunisian Ministry of Public Health, the Ministry of Higher Education and Scientific Research (LR11IPT05) and by the E.C. Grant agreement N° 295097 for FP7 project GM-NCD-Inco and BNP Paribas foundation. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: sonia.abdelhak@pasteur.rns.tn

These authors contributed equally to this work.

Introduction

Profound congenital deafness affects about 1 out of 1000 newborns [1], and is mainly of genetic origin. Non-syndromic (isolated) deafness accounts for approximately 70% of inherited cases. To date, around 70 genes and more than 1000 mutations causing non-syndromic deafness have been reported (<http://deafnessvariationdatabase.org>). More than 45 genes and 69 loci are associated with autosomal recessive non-syndromic deafness (DFNB). Despite the broad genetic heterogeneity of DFNB, loss-of-function mutations in a single gene, *GJB2* (encoding connexin-26) account for more than 30% of the cases in most populations around the Mediterranean sea [2]. After the exclusion of *GJB2* in DFNB patients, finding the gene implicated is difficult due to the high degree of genetic heterogeneity [3]. In addition, many deafness genes consist of long and/or numerous exons, making conventional methods of mutation screening very expensive and time-consuming [4]. Recent advances in DNA enrichment and next generation sequencing techniques, however, allow rapid and cost-effective analysis to identify the causative mutations in deaf patients [5]. To date, ten syndromic or non-syndromic deafness genes have been identified using targeted genomic enrichment and

whole exome sequencing (WES): *TPRN*, *GPSM2*, *CEACAM16*, *SMPX*, *HSD17B4*, *HARS2*, *MASP1*, *OTOGL*, *DNMT1*, and *TSPEAR*. In addition several studies have shown the efficacy of WES to identify the causative mutations in recessive deafness forms [1,4,6,7]. After the exclusion of *GJB2* mutations and of most mutations previously identified in deaf individuals from Tunisia, we carried out WES in six deaf patients from four unrelated consanguineous Tunisian families.

Materials and Methods

Patients

Four unrelated Tunisian families including deaf individuals were included in this study based on parental consanguinity and the presence of at least two affected siblings (**Figure 1**). Written informed consent was obtained from all participants or their legal guardians. Audiological evaluation was carried out at the Otorhinolaryngology Department at La Rabta hospital in Tunis. All patients had bilateral profound sensorineural deafness. Patients had the following clinical investigations: computed tomography of the temporal bones, auditory brainstem response, magnetic

resonance imaging of the inner ear, tympanometry, fundus examination, cardiac and renal ultrasonography. Clinical examinations were unremarkable, and did not reveal symptoms or malformations that would suggest a syndromic form of deafness. Genomic DNA was extracted from peripheral blood samples using the standard salting-out method [8].

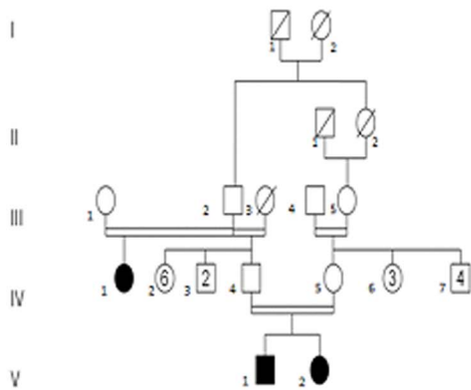
Ethics statement

This study has obtained the ethics approval (IPT/LR11-05/ Etude/05/2013) from the institutional review board of Pasteur Institute (Tunis- Tunisia- Registration number IRB00005445, FWA00010074). This study was conducted according to the principles of the declaration of Helsinki. Patients were anonymized and the corresponding code was conserved in a confidential file.

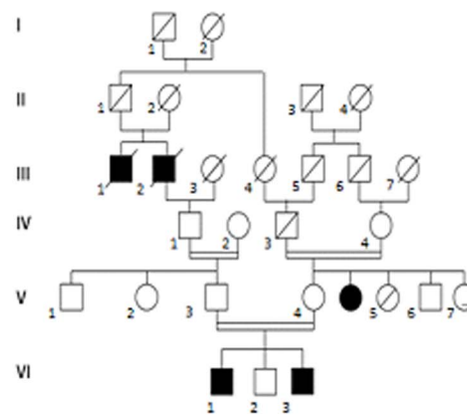
Whole exome sequencing and bioinformatics analysis

A DNA pooling strategy was taken for family DF7 (patients V.1 and V.2) and for family DF56 (patients VI.1 and VI.2). For families DF22 and DF137, DNA was only available from one affected sibling in each family, i.e. VI.1 and V.1, respectively.

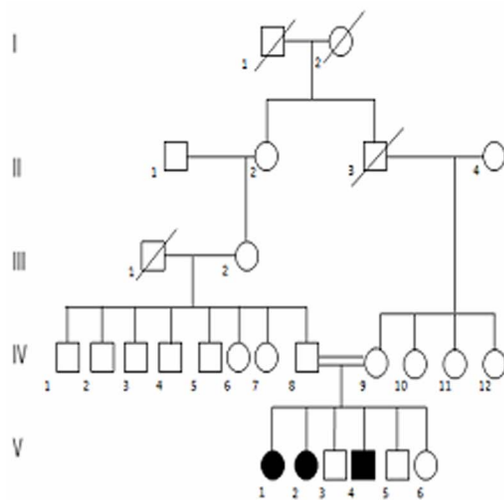
Targeted exome sequencing, library preparation, capture and sequencing, and sequence variant detection and annotation were performed by IntegraGen (Evry, France). Exons of genomic DNA samples were captured using the Agilent in-solution enrichment technique with a biotinylated oligonucleotide probe library, and paired-end 75-base massively parallel sequencing was carried out on an Illumina HiSeq2000. Sequence capture was performed according to the manufacturer's instructions (Human All Exon V5-50 Mb, Agilent). Briefly, 5 µg of each genomic DNA sample was fragmented by sonication and purified to yield fragments of 150–200 bp in length. Paired end adaptor oligonucleotides from Illumina were ligated on repaired A-tailed fragments that were purified and enriched by six polymerase chain reaction (PCR) cycles. Purified libraries (500 ng) were hybridized to the Sure Select oligonucleotide probe capture library for 24 h. After hybridization, washing and elution, the eluted fraction was PCR-amplified (10 to 12 PCR cycles), purified and quantified by quantitative PCR to obtain sufficient amounts of DNA template for downstream applications. Each eluted enriched DNA sample was then sequenced on an Illumina HiSeq 2000 as paired-end 75-



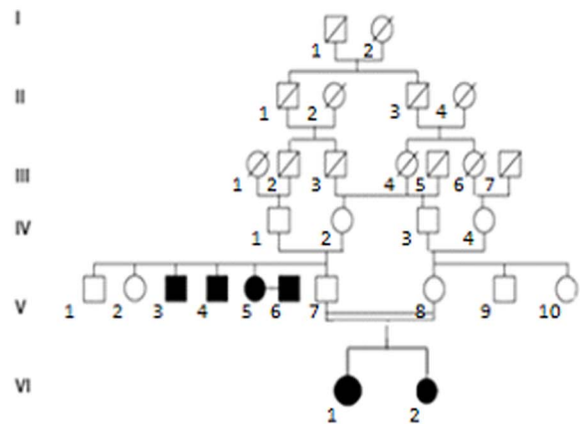
Family DF7



Family DF22



Family DF137



Family DF56

Figure 1. Pedigrees of the four Tunisian families analyzed using the whole exome sequencing strategy.
doi:10.1371/journal.pone.0099797.g001

base reads. Image analysis and base calling were performed using the Illumina Real-Time Analysis Pipeline version 1.14 with default parameters [9].

The exome design covers 51 Mb of the human genome corresponding to the exons and flanking intronic regions of 20 766 genes (220 000 exons) and also 700 miRNAs in the human reference sequence UCSC (hg19/GRCh37, February 2009 release) [7]. Bioinformatics analysis of sequencing data was based on the Illumina pipeline (CASAVA 1.8). CASAVA aligns reads to the human reference genome (hg19) with the alignment algorithm ELANDv2 (it performs multispeed and gapped alignments), calls SNPs on the basis of allele calls and read depth, and detects variants (SNPs and indels). Only positions included in the bait coordinates were conserved. Genetic variation was annotated with the Intergene in-house pipeline, consisting of gene annotation (using RefSeq), detection of known polymorphisms (using dbSNP132 and the 1000 Genomes Project database) and characterization of mutations as intronic or exonic, and silent, nonsense, missense and frame-shifting [9].

Confirmation of the mutations by Sanger sequencing

Sanger sequencing was carried out to validate the mutations identified by WES as previously described [7]. Specific PCR and sequencing primers were designed using Primer3 (**Table 1**) (<http://primer3.ut.ee/>).

Results

Mutations in *GJB2*, the gene most frequently involved in autosomal recessive deafness in Tunisia [10–12], and mutations in other DFNB genes that had previously been reported in Tunisian deaf patients (**Table S1**) were first excluded by PCR and Sanger sequencing of these genes in the patients.

For the WES data analysis, based on familial history and pedigree we hypothesized an autosomal recessive mode of disease transmission and the presence of the causative mutations in the homozygous state in the patients.

To identify pathogenic variants, we filtered out polymorphisms using the Single Nucleotide Polymorphism Database dbSNP132. We excluded all the variants reported in 1000 genomes, Hapmap, and Exome variant server databases. In the second step, we focused on variants which are present in the coding exons and flanking splice sites. From the SNP and indels files, we selected nonsense, frame-shifting (indels), missense, and splice-site mutations, as they were more likely to be pathogenic. Only the variants with a read depth greater than 5 were retained.

After application of these 5 filtering steps (also listed in table 2), we found 3 SNPs and 1 indel inpatients DF7-V.1 and V.2, 11 SNPs and 1 indel inpatient DF22-VI.1, 3 SNPs and 1 indel inpatients DF56-VI.1 and VI.2, and 6 SNPs and 0 indel inpatient DF137-V.1 (**Table 2 and Table S2**).

The biallelic sequence variants predicted to be the causative mutations in the patients are all located in genes already known to be involved in deafness (*MYO15A*, *TMC1* and *LRTOMT*), but these particular mutations have not been previously reported (**Table 3**). Their presence in the homozygous state in all affected siblings, and in the heterozygous state in the clinically unaffected parents, was shown, in each family, by Sanger sequencing of the corresponding DNA fragment (except for the father in family DF56, whose DNA was not available). Finally, none of these mutations was present in the Exome Variant Server database or in 150 ethnically matched normally hearing individuals.

Discussion

We have identified previously unreported biallelic mutations in three different DFNB genes (*MYO15A*, *LRTOMT*, *TMC1*) in patients affected by congenital profound deafness, who belong to four unrelated Tunisian families, using a WES strategy.

In the DF7 patients, a missense mutation, c.5417T>C (p.L1806P), was identified in exon 22 of *MYO15A* (NM_016239). This mutation is predicted to be deleterious according to SIFT (<http://sift.jcvi.org/>) and Mutation Taster (<http://www.mutationtaster.org/>). In family DF137, a different mutation, c.7395+3G>A, was present in the same gene. This mutation is predicted to abolish the splice donor site of intron 37 and to create a cryptic splice donor site 3 bp upstream from the original site, according to the Alamut 2.3 software (<http://www.interactive-biosoftware.com>), which could lead to intron inclusion or exon skipping in the mature transcript. Mutations in *MYO15A* are responsible for the DFNB3 form of deafness [13]. The gene encodes myosin XVa, a 3530 amino acid motor protein involved in the differential elongation of the inner ear hair cells stereocilia [14]. Two mutations in *MYO15A* had previously been identified in Tunisian patients whose hearing impairment loss ranged from severe to profound: c.7395+3G>C and c.4998C>A (p.C1666X) [15]. Mutations in *MYO15A* have also been identified in families originating from Pakistan, India [16] and Japan [17], and were associated with profound deafness in the patients. In family DF22, a nonsense mutation c.208C>T (p.R70X) was identified in exon 7 of *LRTOMT* (NM_001145309). The premature stop codon is predicted to result in a truncated protein with impaired function or no protein at all, due to nonsense mediated mRNA decay, according to Alamut 2.3 software. *LRTOMT* is responsible for the

Table 1. Sequences of the primers used to validate the mutations by Sanger sequencing.

TMC1-EX22_23F	TTTAAGAAGTATCTTGGGGAAGCTG
TMC1-EX22_23R	ATGCCACTCACCATCCAATG
LRTOMT-7F	AGGATAATAATTGCTACTGGCAAAA
LRTOMT-7R	ATCCCAAATATTCCTTCACTGTCTT
MYO15A-EX21F	CCTTTTGCACATGGCTGAG
MYO15A-EX21R	GCCTGGGTTGTGTATTCTG
MYO15A-EX36F	GGTGTGCTAGAATGGAGGCA
MYO15A-EX36R	GAGAGGTGGCAGTGGTGAC

doi:10.1371/journal.pone.0099797.t001

Table 2. Evolution of the number of variants during whole exome.

Patients	DF7-V.1 + V.2		DF22-Vi.1		DF56-Vi.1+Vi.2		DF137-V.1	
	SNP	Indel	SNP	Indel	SNP	Indel	SNP	Indel
Total of variants	42959	6432	70417	5395	77906	6402	72500	5517
After exclusion of variants on chromosomes X and Y	42108	6234	67819	5275	73529	6210	70365	5357
After exclusion of heterozygous variants	12714	1635	28098	2162	21905	1733	26570	1901
After dbSNP132 filtering	247	374	266	601	114	450	179	470
After additional database filtering (Hapmap,1000G, Exome variant server)	32	17	92	30	26	47	68	22
after exclusion of intronic, 5'UTR, 3'UTR, and synonymous variants	3	1	11	1	3	1	6	0

doi:10.1371/journal.pone.0099797.t002

Table 3. Biallelic mutations identified in DFNB genes using a whole exome sequencing strategy.

Family	Genomic position(Hg19)	Gene	Refseq	Exon/intron	cDNA change	Amino acid change	Mutation type
DF7	chr.17 (18044343)	MYO15A	NM_01623	Exon 22	c.5417T>C	p.L1806P	missense
DF137	chr.17 (18054082)	MYO15A	NM_016239.3	Intron 37	c.7395+3G>A		splice site
DF22	Chr.11 (71817106)	LRTOMT	NM_001145309	Exon 7	c.208C>T	p.R70X	nonsense
DF56	chr.9 (75445600)	TMC1	NM_138691	Intron 23	c.2260+2T>A		splice site

doi:10.1371/journal.pone.0099797.t003

DFNB63 deafness form. It produces five different alternatively spliced transcripts, which encode two different leucine rich transmembrane and O- methyltransferase domain containing proteins, *LRTOMT1* and *LRTOMT2*, both expressed in the inner ear sensory cells [18]. Mice carrying a mutation of the orthologous gene (*COMT2*) suffer from vestibular dysfunction, profound deafness and progressive degeneration of the organ of Corti [19]. Two mutations in *LRTOMT* had previously been reported in Tunisian patients, c.242G>A (p.R81Q) and c.313T>C (p.W105R). Two other mutations, c.358+4G>A and c.328G>A, have been identified in deaf patients from Pakistan and Turkey, respectively. In all cases, the hearing impairment was profound [18]. The p.R81Q mutation has also been found in deaf patients from Morocco at a frequency of 8.75%, which makes this gene the second most frequently involved deafness gene in Morocco, after *GJB2* [20]. Finally, the DF56 patients carried a c.2260+2T>A mutation in intron 23 of *TMC1* (NM_138691). According to Alamut 2.3, a cryptic splice donor site will be created 2 bp upstream from the original site, which is expected to result in the skipping of exon 23. *TMC1* has been implicated both in autosomal recessive (DFNB7/11) and autosomal dominant (DFNA36) deafness forms. In Tunisia, one particular mutation in exon 7 of *TMC1*, c.100C>T (p.R34X) has been identified in Tunisian patients affected by autosomal recessive non syndromic profound deafness at a frequency of 5.55% [21]. Mutations in this gene are also described in Turkey [22] and Pakistan [23] with prelingual severe to profound deafness. *TMC1* gene encodes a transmembrane channel-like protein and is expressed in mouse vestibular and cochlear hair cells. The precise role of the encoded protein remains to be established, but it has been to be required for mechanotransduction and proposed to be a component of the mechanotransduction channel. Its absence in *TMC1* deficient mouse lead to hair cells degeneration [24].

In the present study, no genetic linkage analysis was performed prior to exome sequencing in order to identify candidate

chromosomal regions and to reduce the analysis to those regions. Patients were selected based on familial history of deafness and the presence of parental consanguinity. Compared to the classical genetic method (homozygosity mapping and Sanger sequencing) WES is a cost effective strategy to identify mutations causing autosomal recessive deafness in consanguineous families.

Finally, since the Tunisian population shares a common genetic background with other populations of the Mediterranean basin and Middle- East [25], direct screening of the identified mutations in profoundly deaf patients from those regions would be useful to get an estimate of their prevalence.

Supporting Information

Table S1 List of the mutations excluded by Sanger sequencing in ascertained families before whole exome sequencing. (DOCX)

Table S2 List of the mutations retained after the 5 filtering steps. (DOCX)

Acknowledgments

We would like to thank the patients and their families for their collaboration. We are grateful to the technical staff at Armand Trousseau Hospital in Paris for their assistance.

Author Contributions

Conceived and designed the experiments: SA CP. Performed the experiments: ZR CB YB NL. Analyzed the data: ZR CB. Contributed reagents/materials/analysis tools: ML LJ RK JL DW. Wrote the paper: ZR. Contributed in the critical revision of the manuscript for important intellectual content: JPH CB SA CP. Clinical evaluations of patients: RZ GB. Evaluation of the outcome of cochlear implant: SBS. Participated in the study of the control population: MC ID KM OTB.

References

- Yan D, Tekin M, Blanton SH, Liu XZ (2013) Next-generation sequencing in genetic hearing loss. *Genet Test Mol Biomarkers* 17: 581–587.
- Denoyelle F, Marlin S, Weil D, Moatti L, Chauvin P, et al. (1999) Clinical features of the prevalent form of childhood deafness, DFNB1, due to a connexin-26 gene defect: implications for genetic counselling. *Lancet* 353: 1298–1303.
- Sirmaci A, Edwards YJ, Akay H, Tekin M (2012) Challenges in whole exome sequencing: an example from hereditary deafness. *PLoS One* 7: e32000.
- Diaz-Horta O, Duman D, Foster J, 2nd, Sirmaci A, Gonzalez M, et al. (2012) Whole-exome sequencing efficiently detects rare mutations in autosomal recessive nonsyndromic hearing loss. *PLoS One* 7: e50628.
- Woo HM, Park HJ, Baek JI, Park MH, Kim UK, et al. (2013) Whole-exome sequencing identifies MYO15A mutations as a cause of autosomal recessive nonsyndromic hearing loss in Korean families. *BMC Med Genet* 14: 72.
- Bonnet C, Louha M, Loundon N, Michalski N, Verpy E, et al. (2013) Biallelic nonsense mutations in the otogelin-like gene (OTOGL) in a child affected by mild to moderate hearing impairment. *Gene* 527: 537–540.
- Delmaghani S, Aghaie A, Michalski N, Bonnet C, Weil D, et al. (2012) Defect in the gene encoding the EAR/EPTP domain-containing protein TSPEAR causes DFNB98 profound deafness. *Hum Mol Genet* 21: 3835–3844.
- Miller SA, Dykes DD, Polesky HF (1988) A simple salting out procedure for extracting DNA from human nucleated cells. *Nucleic Acids Res* 16: 1215.
- Ishida S, Picard F, Rudolf G, Noe E, Achaz G, et al. (2013) Mutations of DEPDC5 cause autosomal dominant focal epilepsies. *Nat Genet* 45: 552–555.
- Riahi Z, Chahed H, Jaafoura H, Zainine R, Messaoud O, et al. (2013) A novel frameshift mutation (c.405delC) in the GJB2 gene associated with autosomal recessive hearing loss in two Tunisian families. *Int J Pediatr Otorhinolaryngol* 77: 1485–1488.
- Riahi Z, Hammami H, Ouragini H, Messai H, Zainine R, et al. (2013) Update of the spectrum of GJB2 gene mutations in Tunisian families with autosomal recessive nonsyndromic hearing loss. *Gene* 525: 1–4.
- Riahi Z, Zainine R, Mellouli Y, Hammachi R, Bouyacoub Y, et al. (2013) Compound heterozygosity for dominant and recessive GJB2 mutations in a Tunisian family and association with successful cochlear implant outcome. *Int J Pediatr Otorhinolaryngol* 77: 1481–1484.
- Wang A, Liang Y, Fridell RA, Probst FJ, Wilcox ER, et al. (1998) Association of unconventional myosin MYO15 mutations with human nonsyndromic deafness DFNB3. *Science* 280: 1447–1451.
- Belyantseva IA, Boger ET, Naz S, Frolenkov GI, Sellers JR, et al. (2005) Myosin-XVa is required for tip localization of whirlin and differential elongation of hair-cell stereocilia. *Nat Cell Biol* 7: 148–156.
- Belguth H, Aifa-Hmani M, Dhoubi H, Said MB, Mosrati MA, et al. (2009) Screening of the DFNB3 locus: identification of three novel mutations of MYO15A associated with hearing loss and further suggestion for two distinctive genes on this locus. *Genet Test Mol Biomarkers* 13: 147–151.
- Liburd N, Ghosh M, Riazuddin S, Naz S, Khan S, et al. (2001) Novel mutations of MYO15A associated with profound deafness in consanguineous families and moderately severe hearing loss in a patient with Smith-Magenis syndrome. *Hum Genet* 109: 535–541.
- Yano T, Ichinose A, Nishio S-y, Kobayashi Y, Sato H, et al. (2013) A Novel Mutation of MYO15A Associated with Hearing Loss in a Japanese Family. *Journal of Clinical Case Reports*.
- Ahmed ZM, Masmoudi S, Kalay E, Belyantseva IA, Mosrati MA, et al. (2008) Mutations of LRTOMT, a fusion gene with alternative reading frames, cause nonsyndromic deafness in humans. *Nat Genet* 40: 1335–1340.
- Du X, Schwander M, Moresco EM, Viviani P, Haller C, et al. (2008) A catechol-O-methyltransferase that is essential for auditory function in mice and humans. *Proc Natl Acad Sci U S A* 105: 14609–14614.
- Charif M, Bounaceur S, Abidi O, Nahili H, Rouba H, et al. (2012) The c.242G>A mutation in LRTOMT gene is responsible for a high prevalence of deafness in the Moroccan population. *Mol Biol Rep* 39: 11011–11016.
- Ben Said M, Hmani-Aifa M, Amar I, Baig SM, Mustapha M, et al. (2010) High frequency of the p.R34X mutation in the TMC1 gene associated with nonsyndromic hearing loss is due to founder effects. *Genet Test Mol Biomarkers* 14: 307–311.
- Sirmaci A, Duman D, Ozturkmen-Akay H, Erbek S, Incesulu A, et al. (2009) Mutations in TMC1 contribute significantly to nonsyndromic autosomal recessive sensorineural hearing loss: a report of five novel mutations. *Int J Pediatr Otorhinolaryngol* 73: 699–705.

23. Kitajiri SI, McNamara R, Makishima T, Husnain T, Zafar AU, et al. (2007) Identities, frequencies and origins of TMC1 mutations causing DFNB7/B11 deafness in Pakistan. *Clin Genet* 72: 546–550.
24. Kawashima Y, Geleoc GS, Kurima K, Labay V, Lelli A, et al. (2011) Mechanotransduction in mouse inner ear hair cells requires transmembrane channel-like genes. *J Clin Invest* 121: 4796–4809.
25. Romdhane L, Kefi R, Azaiez H, Ben Halim N, Dellagi K, et al. (2012) Founder mutations in Tunisia: implications for diagnosis in North Africa and Middle East. *Orphanet J Rare Dis* 7: 52.