Phenotype prediction of Mohr-Tranebjaerg syndrome (MTS) by genetic analysis and initial auditory neuropathy

Hongyang Wang†, Li Wang†, Ju Yang††, Linwei Yin‡, Lan Lan†, Jin Li†, Qiujing Zhang†, Dayong Wang†, Jing Guan†* and Qiuju Wang†*

Abstract

Background: Mohr-Tranebjaerg syndrome (MTS) is a rare X-linked recessive neurodegenerative disorder resulting in early-onset hearing impairment, gradual dystonia and optic atrophy. MTS is caused by variations in the nuclear TIMM8A gene, which is involved in mitochondrial transport of metabolites. This study aimed to identify the pathogenic gene variations in three Chinese families associated with predicted MTS with or without X-linked agammaglobulinaemia.

Methods: Otologic examinations, vestibular, neurological, optical and other clinical evaluations were conducted on the family members. Targeted genes capture combining next generation sequencing (NGS) was performed, and then Sanger sequencing was used to confirm the causative variation.

Results: A novel variation, c.232_233insCAAT, in TIMM8A was identified as the pathogenic variation in one Chinese family. This variation co-segregated with the most frequent phenotypic deafness and was absent in the 1000 Genomes Project, ExAC and 1751 ethnicity-matched controls. Clinically, otological examinations illustrated the typical postsynaptic auditory neuropathy for the proband without the symptoms of dystonia or optic atrophy. MRI demonstrated abnormal small cochlear symmetric nerves, while the vestibular function appeared to be less influenced. Furthermore, we found another two TIMM8A variations, the deletion c.133_135delGAG and a copy number variation (CNV) including the TIMM8A gene, in two independent case, when we performed NGS on an auditory neuropathy population.

Conclusion: We identified two novel variations in the TIMM8A gene (c.232_233insCAAT and c.133_135delGAG) and a CNV including the TIMM8A gene in three independent Chinese families with predicted MTS. To our knowledge, this is the first report of TIMM8A variations being identified in a Chinese population. Our results enrich the variation spectrum of TIMM8A and clinical heterogeneity of MTS. Genetic detection and diagnosis is a powerful tool for better understanding and managing syndromic hearing impairments, such as MTS, before they become full-blown.

Keywords: Auditory neuropathy, Mohr-Tranebjaerg syndrome (MTS), TIMM8A

Background

Mohr-Tranebjaerg syndrome, also known as deafness-dystonia-optic neuronopathy (MTS/DDON, MIM304700) syndrome, is a rare X-linked recessive neurodegenerative disease. It is characterized by early-onset progressive auditory neuropathy followed by dystonia and optic atrophy in adolescence or adulthood. Additional psychiatric disorders have been reported frequently, such as dementia, irritability and mental retardation. First described in 1960 [1], MTS was mapped to Xq22 by Tranebjaerg in 1995 [2, 3], and the disease-causing variations were found in a gene named translocase of mitochondrial inner membrane 8A/deafness-dystonia peptide-1 (TIMM8A/DDP1) [4]. The TIMM8A gene encodes a 97-amino-acid protein, a translocase involved in the import of metabolite transporters from the cytoplasm into the mitochondrial inner membrane [5, 6].

MTS is a progressive disease associated with multiple systems throughout the patient's life. However, it is not easy to discriminate it due to considerable variations...
both in the onset age and phenotypic expression [7]. To date, only 69 cases have been reported since the first MTS family was described in 1960. Most patients have suffered from deafness and subsequent dystonia for decades before they were diagnosed with MTS. Fortunately, molecular diagnosis through next-generation sequencing (NGS) has made it easier to diagnose deafness syndromes such as MTS, making diagnosis possible decades earlier, before all the phenotypes show. Targeted deafness genes combined with high-throughput sequencing can be regarded as a revolutionary technology with high speed and low cost to identify the pathogenic causes in a large number of syndromic hearing loss families [8, 9].

In this study, we performed variation screening of 127 known deafness-related genes in a young patient from Family 1 without any findings of common deafness-related gene screening. We identified a novel frameshift variation in the TIMM8A gene in this Chinese family with auditory neuropathy, who visited the outpatient department for deafness genetic consultation, as typical auditory neuropathy was the only phenotype in the proband at this stage. Furthermore, we performed targeted capture of 127 genes and NGS on another 167 patients diagnosed with auditory neuropathy and found another two TIMM8A variations, as well as agammaglobulinemia in one positive case. To our knowledge, this is the first report of MTS cases recognized efficiently with NGS before clinical diagnosis.

**Methods**

**Clinical evaluations**

Three probands, from Family 1, Family 2 and Family 3, who were diagnosed with auditory neuropathy and their parents were recruited (Fig. 1a, Fig. 2a and Fig. 3a), written informed consents were obtained from the next of kin on the behalf of the minors/children participants involved in this study. This study was approved by the Ethics Committee of Chinese PLA General Hospital.

Their medical histories were collected by a questionnaire. Otological examinations, including otoscopy, immittance testing, pure-tone audiometric examination (PTA, Madsen Astera², DENMARK), speech recognition score (SRS, Madsen Astera², DENMARK), distortion product otoacoustic emission (DPOAE, Madsen Capella, DENMARK), click-evoked auditory brainstem response (ABR), cochlear microphonics (CM) and electrocochleography (ECoG) tests (SmartEp, USA or Neuro-Audio, Russia), were performed to evaluate the auditory conditions. The diagnosis of auditory neuropathy was made according to the results of DPOAE/CM and ABR tests. The hearing level was assessed at 125, 250, 500, 1000, 2000, 4000 and 8000 Hz by PTA. The brain magnetic resonance imaging (MRI) examination for the temporal bone of both ears was also performed on the proband. Vestibular function evaluation included vestibular-evoked myogenic potentials (ocular VEMP, oVEMP and cervical VEMP, cVEMP), oculomotor function tests, positional nystagmus tests, positioning nystagmus tests and bithermal calorics tests. The ocular examinations included visual acuity, perimetry, flash and pattern visual-evoked potential testing, and detailed stereoscopic fundoscopy. The neurological examination included regular physical examinations and electromyography.

**Targeted gene capture and NGS**

Peripheral blood samples of these families were collected with consent. Genomic DNA was extracted from the whole blood of the probands and their parents (TIAN-GEN BIOTECH, BEIJING, CHINA). After the examination of DNA quality, Beijing Genomics Institute built the DNA libraries by following Illumina’s protocol, and then 127 known deafness-related genes, including exons, splicing sites and their flanking introns, were captured by using a custom probe and sequenced by the Illumina HiSeq2000 [10]. The paired-end reads generated by sequencing were aligned to the NCBI37/hg19 assembly by the BWA software, and variant calling was performed by GATK. The bioinformatics analysis method had been described in detail previously. To obtain rare variations, common variants and low-frequency variants in the 1000 Human Genomes Project database, HapMap Project database, ExAC database, EVS database and BGI in-house databases were excluded (0.5%). All remaining variants were considered to be rare variations and were annotated using Ensembl VEP, OMIM, MGI, Gene Ontology, HGNC gene annotation database and so on. Candidate variants were validated by Sanger sequencing.

**SNP array for CNV analysis**

Genome-wide chromosome copy number anomalies were detected using Illumina’s InfiniumOmniZhongHua-8 DNA Analysis Bead Chip (200925710145_R03C01), with a resolution of 20 kb.

**Variant filtration, confirmation and modelling**

Variants with allele frequencies higher than 5% in the 1000 Genomes Project and the local database were excluded. Splicing-site, frameshift and nonsense variants were taken into further consideration. Moreover, SIFT and PolyPhen2 software were used to evaluate the pathogenic possibility. Sanger sequencing was performed to establish the co-segregation of the candidate gene variations with the phenotype in the family members. A three-dimensional structure of TIMM8A was built by SWISS-MODEL (https://swissmodel.expasy.org/) and then visualized by Swiss-PdbViewer 4.1 (version 4.1, http://spdbv.vital-it.ch/).
Overview of MTS cases reported previously and related variations

The EMBASE and PUBMED databases were searched for literature review. The clinical phenotypes and associated genotypes of these MTS cases were summarized. Then, comparisons of the types of genotype and onset times of different clinical characteristics were analysed.
Results
Clinical multiple-discipline evaluations
The proband of Family 1 (Fig. 1a) was found to have hearing impairment from age three. Otological examinations, including DPOAE, ABR, CM, PTA and Phtmax, were performed on this 7-year-old boy, and the typical auditory neuropathy was illustrated. His parents had normal hearing, while the audiogram of the proband showed severe hearing impairment with profoundly damaged SRS (16 and 12% for the left and right ears, respectively). PTA audiogram on the left side was low-frequency ascending and on the right was mid-frequency U-shaped (Fig. 1b). As shown in Fig. 1c, DPOAE for the proband was normal, and no waves could be detected in ABR testing bilaterally, while CM waves from both ears existed (Fig. 1d). ABR testing on patient II:4 (mother of the proband) showed no abnormal latency or amplitude of ABR waves I, III and V (Additional file 1: Figure S1), as well as normal DPOAE, indicating the presence of outer hair cells in the cochlea (Additional file 2: Figure S2). In addition, waves of oVEMP...
and cVEMP testing showed normal amplitude and latencies, but decreased function of the horizontal semicircular canal was indicated by the bithermal caloric test (details recorded in Additional file 3: Figure S3 & Additional file 4: Table S1). MRI of the proband (Additional file 5: Figure S4) demonstrated an abnormally small cochlear nerve symmetrically. Regarding dystonia, no abnormal manifestation was observed by neurological examination, nor was any found on electromyography. Regarding optic atrophy, both the proband and his mother had normal visual acuity. Flash and pattern visual-evoked potential testing showed a normal latency and amplitude of P100 in both eyes, and detailed stereoscopic fundoscopy showed normal results (Additional file 6: Figure S5). Perimetry
was not available for this young child. The γ-globulin level of the proband was normal.

The probands from Family 2 (Fig. 2a) and Family 3 (Fig. 3a), who were diagnosed with auditory neuropathy (Fig. 2b, and c and Fig. 3b, c, d, e, and f), had similar audiological phenotypes as the proband from Family 1, as well as similar MRI findings and visual acuity. The major difference in the proband from Family 3 was the abnormal γ-globulin level, showing agammaglobulinemia when tested. The serum immunoglobulins of the proband from Family 2 were 0.30, 1.22 and 0.13 g/l of IgA, IgG and IgM, respectively.

**TIMM8A variation detection and analysis**

We identified a hemizygous variation, c.232_233insCAAT (p.Leu78SerfsX21), in the TIMM8A gene (NM_004085) in Family 1 (Fig. 4a and Table 1). This novel variation in the second exon of TIMM8A caused one amino acid substitution, leucine to serine at position 78, and a frame shift after that position. That position is highly conserved across species (Fig. 4b). Co-segregation of this variation with the disease in the family was confirmed using Sanger sequencing, as shown in the Fig. 4c. The c.232_233insCAAT variant was also detected in the proband’s mother with normal hearing, but it was absent in his father. This variation was not found in the 1000 Genomes Project, ExAC data and 1751 ethnicity-matched controls, which further supported its pathogenicity. By using SWISS-MODEL, a three-dimensional molecular structure was built to locate the mutated scope on the second exon of TIMM8A (Fig. 4d).

To further determine the frequency of TIMM8A variations in the auditory neuropathy population, we analysed the targeted gene capture and NGS from 167 cases in the auditory neuropathy population was 1.8% (3/168). Worldwide, the 14 previously reported TIMM8A abnormalities associated with MTS included missense/nonsense variations, deletions and splicing variants [7]. Therefore, the new insertion variation in this study presented a new genotype for MTS. Regarding phenotype, we observed that three variations in exon 2 did not result in visual loss, whereas other clinical manifestations showed poor correlations with genotypes.

**Discussion**

In the present study, we identified three novel TIMM8A variations in three Chinese families by applying targeted gene capture to 127 deafness-related genes combined with NGS, as well as a SNP array. The variations of TIMM8A were co-segregated in the families and absent in the 1000 Genomes Project, ExAC data and 1751 ethnicity-matched normal controls. According to previous studies [11], the Tim10/DDP family zinc finger domain includes amino acids 22 to 85 of TIMM8A. Both the frameshift variation c.232_233insCAAT and the deletion variation c.133_135delGAG were located in this special domain and resulted in amino acid sequence alterations starting at positions 78 and 45, respectively, impairing the formation of TIMM8A protein. TIMM8A belongs to a family of evolutionary conserved proteins assembled into a hetero-hexameric complex with human Tim13 [12]. Thus, these variations in the TIMM8A gene might interfere with formation of the heterodimer complex between Tim8 and Tim13.

Auditory neuropathy is a unique hearing dysfunction characterized by absent or abnormal ABR and the presence of OAE and/or CM [13]. Disruption of auditory nerve activity may involve the auditory nerve (postsynaptic auditory neuropathy), inner hair cells and/or the synapses with auditory nerve terminals (presynaptic auditory neuropathy) [14]. The auditory neuropathy in our study belonged to the postsynaptic kind since previous studies on temporal bone histopathology from MTS patients showed a 90–95% loss of cochlear function [15]. Meanwhile, MRI examination of the probands further confirmed the abnormally small cochlear nerve compared with nearby constructions. Somewhat differently, electrophysiologic evaluations of the probands demonstrated the absence of vestibular and optical deficits and dystonia, which may be ascribed to the early stage in the course of MTS. Since few studies have indicated that the carrier female also developed deafness or dystonia [3, 16], the mothers of probands in our study also underwent overall examinations and were assessed as unaffected. Considering MTS is a progressive degeneration [17], long-term follow-up would benefit the patients. Recently, internal globus pallidus (GPI) deep brain stimulation (DBS) was utilized to treat successfully at severe dystonia [18], while cochlear implantation produced little effect [19].

Clinical phenotypes and related known genotypes are overviewed in Table 1, which highlights the high

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**Overview of known MTS cases on genotype and phenotype**

Genotype and phenotype correlation analysis for MTS cases is summarized in Table 1. Worldwide, the 14 previously reported TIMM8A abnormalities associated with MTS included missense/nonsense variations, deletions and splicing variants [7]. Therefore, the new insertion...
clinical heterogeneity of MTS. All 69 patients reported have hearing impairments within the first decade, but the onset time of dystonia varies, and most occur in the second decade of life. Remarkably, all three probands in this study appeared to be without visual loss at the tested age. The possible explanation for the incomplete optical penetrance in MTS could be related to the damage extent of the structure or function of TIMM8A. To our knowledge, this study is the first report of TIMM8A-related hearing loss in China, and it is not rare in the auditory neuropathy population, making the case for better understanding the underlying mechanism. Only one patient with CNV in this study showed agammaglobulinaemia. The 16,334-bp deletion including TIMM8A and BTK identified in Family 3 may be related to contiguous deletion syndrome with the co-existence of X-linked agammaglobulinemia (XLA) and MTS.

Our original aim in studying these three families was to ask for genetic consultation, as the probands were solely characterized by early onset auditory neuropathy without any signs of dystonia or visual impairment. Genetic diagnosis of MTS for the young patient was confirmed ahead of clinical diagnosis. In fact, it was NGS technology that brought them a clear answer and perspective on the disease. Therefore, targeted capture of deafness genes combined with NGS has become a powerful tool to provide with precise molecular diagnoses and optimal rehabilitation for syndromic hearing-impaired patients as well as accurate genetic counselling for their whole family.
| Year          | Source                          | Affected members, No. of Generations | Phenotypes                        | Onset of Deafness | Onset of Dystonia | Onset of Impaired vision | Psychiatric disorders | TIMM8A Variations | Type       |
|---------------|---------------------------------|--------------------------------------|-----------------------------------|-------------------|-------------------|--------------------------|-----------------------|-------------------|------------|
| 1976          | America [20]                    | 3,2                                  |                                   | 2–6               | 7                 | None                     |                       |                   | NR         |
| 1960,1995,1996| Norway                          | 16,5                                 |                                   | 2–7               | 7–50              | mid-thirties             | Peripheral neuropathy, mental deterioration, et al | c.151delT       | Frameshift  |
| 1996          | America [3]                     | 5,2                                  |                                   | NR                | NR                | none                     | Mental impairment     | c.183delI0       | Frameshift  |
| 1998          | Australia [21]                  | 3,2                                  |                                   | 2                 | 6–12              | 12–16                    | Possible              | c.52C > T        | NR         |
| 2000          | Dutch [22]                      | 1,1                                  |                                   | 2.5               | 10                | None                     | Hyper-reflexia, dyspraxia | c.233C > G      | missense    |
| 2001          | Denmark [23]                    | 3,5                                  | Early infancy-2                   | 21                | 14–18             | Mildly demented           |                       | c.105G > T       | nonsense   |
| 2001          | America [16]                    | 8,4                                  |                                   | congenital        | Teens-adult       | early fifties            | Dementia              | c.108delIG      | Frameshift  |
| 2001          | Japan [24]                      | 5,4                                  |                                   | 0.5–9             | 16–30             | none                     | Mild mental impairment | c.238C > T       | Nonsense    |
| 2003          | Germany [25]                    | 1,1                                  |                                   | 1                 | 1                 | 37                       | None                  | c.38G > C        | Missense    |
| 2004          | Italy [26]                      | 1,1                                  |                                   | 2                 | 19                | 15                       | Cognitive decline      | complete deletion  |             |
| 2005          | Spain [27]                      | 2,2                                  |                                   | 4–11              | 11–20             | 24                       | Cognitive decline      | IVS1-23A > C     | Splice      |
| 2006          | Spain [28]                      | 2,3                                  |                                   | 3–7               | 8                 | none                     | Attention deficit and hyperactivity disorder | c.127delT       | Frameshift  |
| 2007          | UK [29]                         | 1,1                                  |                                   | < 1               | 25                | 30                       | None                  | IVS1+ 1G > A     | Splice      |
| 2007          | Spain [30]                      | 1,1                                  |                                   | 3                 | 30                | 15                       | Distractibility, irritability, and childish manners | c.112C > T       | Nonsense    |
| 2007          | Czech Republic [17]             | 2,1                                  |                                   | 2.5               | 33                | NR                       | Aggressive behaviour  | complete deletion  |             |
|               | Czech Republic                  | 1,1                                  |                                   | 4                 | 5                 | NR                       | NR                   | complete deletion  |             |
|               | Estonia                         | 2,1                                  |                                   | 2.5,4             | NR                | NR                       | Progressing psychomotor retardation | complete deletion  |             |
|               | African-America                | 1,1                                  |                                   | 3                 | 3                 | NR                       | NR                   | complete deletion  |             |
| 2008          | Spain [31]                      | 1,1                                  |                                   | 4                 | 23                | 14                       | Mild mental retardation | IVS1+ 1G > A     | Splice      |
| 2008          | America [19]                    | 1,1                                  |                                   | 2                 | NR                | NR                       | exon 1 deletion        |                   |             |
| 2011          | Japan [32]                      | 1,1                                  |                                   | 1                 | NR                | NR                       | complete deletion      |                   |             |
| 2013          | Japan                            | 1,1                                  |                                   | 1.5               | NR                | NR                       | complete deletion      |                   |             |
| 2013          | France [18]                     | 1,1                                  |                                   | 2.5               | 8                 | None                     | Congenital mental retardation | IVS1+ 1G > A     | Splice      |
| 2016          | Poland [33]                     | 1,1                                  |                                   | 3                 | 12                | None                     | NR                   | NR               |             |
| Year | Source          | Affected members, No. of Generations | Phenotypes | Genotypes |
|------|----------------|--------------------------------------|------------|-----------|
|      |                |                                      | Onset of Deafness | Onset of Dystonia | Onset of Impaired vision | Psychiatric disorders | TIMM8A Variations | Type         |
| 2016 | French-Canadian [34] | 4,3                                  | 2          | NR         | NR         | progressive psychomotor deterioration | complete deletion |
| 2018 | China          | 1,2                                  | 3          | Not yet    | Not yet    | Not yet                               | c.232_233insCAAT  | p.L78fs21     | Frameshift   |
| 2018 | China          | 1,2                                  | 13         | Not yet    | Not yet    | Not yet                               | c.133_135delGAG   | p.Glu45del    | Indel        |
| 2018 | China          | 1,2                                  | 13         | Not yet    | Not yet    | Not yet                               | arr[hg19] Xq22.1  | /            | CNV          |
Conclusion
In conclusion, three novel hemizygous variations of TIMM8A were the pathogenic variations in three Chinese families with predicted MTS. Our data extend the variation spectrum of the TIMM8A gene and give deeper insight into phenotypes of MTS. Target deafness gene capture and

Fig. 5 Copy number variation including TIMM8A of Family 3. a NGS result showing the deletion in the TIMM8A gene. b Molecular cell karyotype of Xq22.1(100,593,213-100,609,547)x0 of the proband from Family 3
high-throughput NGS increased the diagnosis yield of syndromic genetic hearing loss such as MTS before all symptoms and signs were fully developed.

Additional files

Additional file 1: Figure S1. ABR and CM waves of the proband’s mother (II 8) in Family 1. Normal latency and amplitude of ABR waves I, III and V are shown. (TIF 1369 kb)

Additional file 2: Figure S2. DPOAE results of the proband’s mother in Family 1. (TIF 2415 kb)

Additional file 3: Figure S3. VEMP waves of the proband in Family 1 showing normal latency and amplitude. (TIF 4272 kb)

Additional file 4: Table S1. Vestibular Function Evaluation and results of proband of Family 1. (DOCX 17 kb)

Additional file 5: Figure S4. Brain MRI examination of the proband from Family 1. A. Axial view of the cerebellopontine angle and the temporal bone. The IAC is visible, and the facial nerve is normal. B. 3D-fast-spin echo sequence image on oblique plane sagittal from normal age-matched control. Left side demonstrates a normal cochlear nerve (Cn, red arrow), normal-sized IAC, facial (Fm), superior (Vsn) and inferior vestibular nerves (Vin) (yellow arrows). C&D image from proband: abnormally small cochlear nerve (red arrows) in both sides. (TIF 4158 kb)

Additional file 6: Figure S5. Visual-evoked potential testing and stereoscopic fundoscopy of the proband from Family 1. (TIF 3098 kb)

Abbreviations

ABR: Auditory brainstem response; CM: Cochlear microphonics; CNV: Copy number variation; cVEMP: Cervical vestibular-evoked myogenic potentials; DPOAE: Distortion product otoacoustic emission; ECochG: Electrocochleography; MRI: Magnetic resonance imaging; MTS: Mohr-Tranebjaerg syndrome; NGS: Next generation sequencing; oVEMP: Ocular vestibular-evoked myogenic potentials; PTA: Pure-tone audiometric examination; SRS: Speech recognition score; XLA: X-linked agammaglobulinaemia

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Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

Conceived and designed the experiments: QW HW LW JY. Performed the experiments: HW LW JY. Analyzed the data: HW LW LY JG LL JL. Contributed reagents/materials/analysis tools: JY QZ DW. Wrote the paper: HW LW. Critical reading and discussion of manuscript: DW QW. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The study was approved by the Committee of Medical Ethics of Chinese PLA General Hospital. Written informed consent from all the participants in the family was obtained.

Consent for publication

Written informed consents for publication from all the participants in the family were obtained from the next of kin on behalf of the minors/children participants involved in this study.

Competing interests

The authors declare that they have no competing interests.

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