

J Human Genet

TECTA mutations in Japanese with mid frequency hearing loss affected by zona
pellucida domain protein secretion

Running title: *TECTA* mutations in Japanese

¹Hideaki Moteki, ¹Shin-ya Nishio, ¹Shigenari Hashimoto, ¹Yutaka Takumi,

²Satoshi Iwasaki, ³Norihito Takeichi, ³Satoshi Fukuda, ¹Shin-ichi Usami

¹Department of Otorhinolaryngology, Shinshu University School of Medicine

²Department of Hearing Implant Science, Shinshu University School of Medicine

³Department of Otolaryngology Head and Neck Surgery, Hokkaido University School
of Medicine

Corresponding author:

Shin-ichi Usami, M.D., Ph.D.,

Department of Otorhinolaryngology,

Shinshu University School of Medicine,

3-1-1 Asahi, Matsumoto 390-8621, JAPAN.

Tel: +81-263-37-2666

Fax: +81-263-36-9164

E-mail: usami@shinshu-u.ac.jp

Abstract

TECTA gene encodes α -tectorin, the major component of noncollagenous glycoprotein of the tectorial membrane, and plays a role in intracochlear sound transmission. The *TECTA* mutations are one of the most frequent causes of autosomal dominant hearing loss and genotype-phenotype correlations are associated with mutations of *TECTA* in exons according to α -tectorin domains. In this study, we investigated the prevalence of hearing loss caused by *TECTA* mutations in Japanese autosomal dominant hearing loss families, and confirmed genotype-phenotype correlation, as well as the intracellular localization of missense mutations in the α -tectorin domain. *TECTA* mutations were detected in 2.9% (4/139) of our Japanese autosomal dominant hearing loss families, with the prevalence in moderate hearing loss being 7.7% (4/52), and all patients showed typical genotype-phenotype correlations as previously described. The present in vitro study showed differences of localization patterns between wild type and mutants, and suggested that each missense mutation may lead to a lack of assembly of secretion, and may reduce the incorporation of α -tectorin into the tectorial membrane.

Keywords: *TECTA*/ autosomal dominant hearing loss/ genotype-phenotype correlations / zona pellucida domain/ mid frequency hearing loss

Introduction

Hearing loss affects about 1 in 500 to 1000 newborns in developed countries and genetic causes account for at least 50% of all childhood nonsyndromic sensory neural hearing loss (SNHL)¹. Most of these cases are affected with severe and congenital pre-lingual deafness and autosomal recessive inheritance represented by *GJB2* gene mutations predominates (80%) over autosomal dominant (20%)². Mild to moderate SNHL and/or late onset SNHL, presenting with autosomal dominant inheritance pedigree pattern, is commonly supposed to be of genetic causes. Autosomal dominant nonsyndromic hearing loss (ADNSHL) is represented by heterogeneity of genetic and clinical features, as 60 loci have been mapped, 24 genes have been cloned, and correlation with audiological features have been reported [Van Camp and Smith, Hereditary Hearing Loss Homepage. WorldWide Web URL: <http://webhost.ua.ac.be/hhh/> 2011]. These types of SNHL can be characterized by age of onset, progression and pattern of audiogram.

As one cause of ADNSHL, *TECTA* mutations have been found in various types of hearing loss, age of onset, progression and frequency involvement in various populations³⁻¹⁰. This gene encodes α -tectorin, the major component of noncollagenous glycoprotein of the tectorial membrane that consists of an extracellular matrix overlying

the organ of corti, contacting the outer cochlear hair cells, and playing a role in intracochlear sound transmission¹¹. The α -tectorin is composed of three distinct modules: the entactin G1 domain, the zonadhesin (ZA) domain with von Willebrand factor type D repeats, and the zona pellucida (ZP) domain¹¹. No nonsense mutations of *TECTA* have been reported in autosomal dominant hearing loss. Missense mutations affecting the ZP domain are associated with mid-frequency hearing impairment, whereas mutations in the ZA domain are associated with hearing impairment primarily affecting the high frequencies¹². Phenotypes of hearing loss can range from mild to severe and have pre or postlingual onset⁸.

In this study, (1) we examined the prevalence of hearing loss caused by *TECTA* mutations in Japanese ADSHNL and confirmed genotype phenotype correlation, and (2) examined the impact of three missense mutations in the ZP domain on the cellular distribution of α -tectorin, known to cause mid-frequency hearing impairment. Many deafness-causing *TECTA* mutations have been reported, but the molecular mechanisms are unclear. To investigate the biological function of missense mutations in the ZP domain that were reported as causing ADSHNL GFP fusion proteins were generated and the effects of corresponding mutations on secretion patterns of the ZP domain of α -tectorin were examined.

Materials and Methods

Subjects

139 Japanese autosomal dominant (AD; with two or more generations affected) sensorineural hearing loss families were screened for mutations in the *TECTA* gene. All probands were from independent families and none had any other associated neurological signs, visual dysfunction, or diabetes mellitus. Hearing level was classified by a pure-tone audiometry average over 500, 1000, 2000 and 4000 Hz in the better hearing ears as follows: normal hearing, <20 dB; mild hearing loss, 21-40dB; moderate hearing loss, 41-70 dB; severe hearing loss, 71-95 dB; and profound hearing loss, greater than 95 dB (GENDEAF, 2004). Of the 139 probands, four (3%) had normal hearing (only limited frequencies involved), 40 (29%) had mild hearing loss, 52 (37%) had moderate hearing loss, 23 (17%) had severe hearing loss, and 12 (9%) had profound hearing loss. Information on pure tone audiometry was not available for eight (6%) of these subjects. The mean age at the time of their participation (not onset of hearing loss) of the subjects were; normal hearing; 14.0 ± 10.6 years, mild hearing loss; 21.1 ± 15.5 years, moderate hearing loss; 25.3 ± 19.0 years, severe hearing loss; 32.1 ± 25.5 years and profound hearing loss; 27.5 ± 19.2 years.

All subjects gave prior informed written consent for participation in this study and the Ethical Committee of Shinshu University approved the study.

Mutation analysis

All 23 exons and flanking intronic sequences of the *TECTA* gene were amplified by polymerase chain reaction (PCR). Primers were designed to flank all of the exon-intron boundaries through use of the Primer3 web-based server (http://www-genome.wi.mi.edu/cgi-bin/primer/primer3_www.cgi). Each genomic DNA sample (40 ng) was amplified, using Ex-Taq polymerase (Takara), for 5 min at 95°C, followed by 37 three-step cycles of 95°C for 30s, 56-63°C for 30s, and 72°C for 1.5 min, with a final extension at 72°C for 10 min, ending with a holding period at 4°C in a Perkin-Elmer thermal cycler. The PCR products varied in size at about 200-700 bp, and they were treated with 0.1 µl exonuclease I (Amersham) and 1 µl shrimp alkaline phosphatase (Amersham) and by incubation at 37°C for 30 min, and inactivation at 80°C for 15 min. After the products were purified, we performed standard cycle sequencing reaction with ABI Big Dye terminators in an ABI 3100 autosequencer (Applied Biosystems, Foster City, CA, USA).

cDNA products ZP domain expression plasmids

Full length cDNAs of the ZP domains of α -tectorin genes were cloned by conventional PCR from the human fetal brain cDNA library (Invitogen, Carlsbad, CA, USA). Two pairs of primers for the entire coding regions of ZP domain including transmembrane domain (TMD) of α -tectorin were used. PCR steps were denaturing at 94°C for 2 min, followed by 30 cycles of 94°C for 30 s, 53°C for 30 s, and 72°C for 1 min, and then processing with a final extension at 72°C for 5 min. After amplification, expected sizes of PCR products were confirmed on 2% agarose gel, and the bands were visualized by ethidium bromide upon exposure to an ultraviolet transilluminator.

Produced cDNAs were digested with *Bam*HI/*Eco*RI and cloned into the *Bam*HI/*Eco*RI site of pEGFP C2 vector (Clontech, Palo Alto, CA, USA). Ligation reactants were transformed into *Escherichia coli* DH5a. A QIAprep spin miniprep kit (Qiagen, Valencia, CA, USA) was used for purification of plasmid DNA according to the manufacturer's protocol.

Site-directed Gene Mutagenesis

Gene mutagenesis of the ZP domain cDNA was performed by mega primer PCR¹³. The

mutagenesis primers for 5509TG, 5876A>G and 6063G>A which were previously reported from Spain⁴, Austria¹⁴ and Japan⁵ respectively, were designed. The following reverse primers were used to produce the mutations for initial PCR reaction: 5509TG CCCCTCGATGCCGGTGCCCTGTCTGTCA, 5876AG TCCAGAGTGTGTTTTTACACATGATATG and 6063GA ACCGAGCTGGAAGAACTTGCACTTAGAT. Initial PCR reactions (20µl) were prepared containing 0.1µg of template DNA, 0.4 µM of mutation primer, 0.4µM of ZP-Eco RI primer, 0.1 Unit of KOD pulse (TOYOBO, Japan), and KOD buffer, 2.0µM MgSO₄, and 0.8µM dNTP. These PCR reactions were denatured at 94°C for 2 min, followed by 30 cycles of 94°C for 30 s, 55°C for 30 s, and 72°C for 1 min, and then processed with a final extension at 72°C for 5 min. The subsequent mega primer reactions were prepared containing initial PCR products which were diluted 50-fold each, 0.4µM of ZP-BamHI R primer, 2.0 U of Takara Ex taq (Takara, Japan), and Ex-taq buffer (10x). These PCR products were inserted into a pEGFP-C2 vector with the same techniques as above. The sequences of all three cDNA constructs were confirmed by DNA sequencing using an ABI 3100 autosequencer.

Transfection and confocal microscopy

COS-7 cells grown in DMEM (Mediatech, Herndon, VA) supplemented with 10% fetal calf serum (Moregate, QLD, Australia) , were transiently transfected with the indicated plasmids, using Lipofectamine 2000 (Invitrogen) as described by the manufacturer.

Twenty-four hours after the transfection, cells were washed twice with PBS.

Cover slips were mounted onto glass slides and visualized under a Leica confocal microscope TCS SP2 AOBS (Leica Microsystems, Wetzlar, Germany).

Results

Mutation Screening of the *TECTA* Gene

Direct DNA sequencing identified four pathogenic mutation alleles from AD families including one family in which two mutations were found in one allele. Among those, the family with the c.6063G>A (p.R2021H) mutation was previously reported by Iwasaki et al⁵. Including those results, *TECTA* mutations were detected in 2.9% (4/139) of Japanese ADNSHL families, and the prevalence in moderate hearing loss was 7.7% (4/52).

The family F818 pedigree consisted of three generations and included nine affected members (four males and five females), four of whom participated in this study (Figure 1). This family had a p.R1773X (c.5318C>T) mutation affecting the ZA domain in exon 16, and had slowly progressive high frequency hearing loss. Segregation with hearing loss was confirmed in all cases for which DNA samples were available and none of the mutations were detected in controls.

The family F237 pedigree consisted of three generations and included five affected members (three males and two females), four of whom participated in this study (Figure 2). They demonstrated bilateral mild to moderate symmetric sensorineural hearing loss and showed a U-shaped audiogram, affected in the mid frequencies.

Vestibular disorder symptoms were not observed, and inner ear abnormalities were not found with CT scans. Two missense mutations in one allele, p. [H1400W; T1866M] (c. [4198C>T; 5597C>T]), were detected in α -tectorin in this family. The mutation H1400W in exon 12 was in the ZA domain of α -tectorin, whereas T1866M was in the ZP domain.

The family F652 pedigree consisted of four generations and included 16 affected members (Figure 3). I1997T (c.5990T>C) mutations were detected in α -tectorin in this family. The mutation in exon 19 located in the ZP domain of α -tectorin. This missense mutation appeared in heterozygosity and was shown to segregate almost completely with the affected status in this family. The audiograms were symmetric and often showed a U shape, which indicates that predominantly the mid frequencies are affected. But, one member, an 11-year-old girl (Figure 3. A; III-6), although bearing the responsible mutation, had a normal audiogram and no demonstrable hearing loss.

Localization of ZP domain mutants

The inherent fluorescence of GFP determined the intracellular localization of the

recombinant fusion proteins (Figure 4). Transfected GFP- ZP domains of α -tectorin wt (wild type) were found to be localized as labeled puncta, which may be secreted along the plasma membrane. In contrast, GFP- ZP domains of α -tectorin mutants, (GFP-ZP mut) C1837G, Y1870C and R2021H, were not recognized at the plasma membrane but were retained within the cytoplasm where they formed vesicles.

Discussion

We have identified four independent autosomal dominant families associated with four different *TECTA* mutations. Prior to this study, one Japanese family with 6063G>A (R2021H) mutation had been reported⁵. Including that family, prevalence of ADNSHL with *TECTA* mutation was 2.9% (4/139 families), which may be a relatively high incidence. Hildebrand et al. reported that its prevalence was about 4% in Spanish ADNSHL families (17/374 families)⁸. In our results, when limited to moderate hearing loss patients there was a higher rate of detection (7.7%; 4/52 families).

In this study, all patients showed typical genotype-phenotype correlations of SNHL with *TECTA* mutation as previously described^{8, 12, 15}. The family F818 in which the R1773X (c.5318C>T) mutation in the ZA domain was detected, showed high frequency hearing loss that was slowly progressive. The affected proband (Figure 1) noticed bilateral hearing impairment when she was around age 20, and her right hearing level was worse than the left because of cholesteatoma in her right ear. Hearing impairment was detected in her sons in school health checks but they had never suffered vertigo and no inner ear abnormality was seen in CT scans.

In the family F237, two missense mutations, H1400W (c.4198C>T) and T1866M (c.5597C>T), were detected in α -tectorin. The mutation H1400W in exon 12

was in the ZA domain of α -tectorin, whereas T1866M was in the ZP domain. Both amino acid residues were conserved among another species. It had been suggested in a previous report that *TECTA*-affected mid-frequency hearing impairment appeared to be related to the position of the mutations in the ZP domain of α -tectorin. Considering the phenotype and position of the mutation, T1866M was likely to be causative for hearing impairment in this family. The influence of the nucleotide change of c.4198C>T on apparent effect of splicing of the *TECTA* mRNA cannot be predicted. However, because this change was not present in the controls, it cannot be ruled out that it has an effect on the phenotype of these patients or it may even act synergistically with the T1866M (c.5597C>T) mutation. The similar results with two changes in one family were reported by Plantinga in 2006¹². The T1866M mutation that we detected in this study was previously reported in one family each in Korea, Spain and the USA^{7,8}. Hildebrand reported that the Spanish and American cases do not suggest a founder effect for this mutation⁸. Therefore, this T1866M mutation, now known to be existent in four independent families from four different countries, is suggested to be a possible mutational site hot spot.

In the family F652, we detected a novel mutation, I1997T (c.5990T>C), in exon 19 located in the ZP domain of α -tectorin. The audiograms of affected patients

indicated U-shaped mid-frequency hearing loss, associated with a ZP domain mutation previously reported. Regarding progression of hearing loss, these audiograms showed that the thresholds depend on age among these generations. Accordingly, this mutation would lead to slowly progressive mid-frequency hearing loss. Interestingly, an affected female (Figure 3. A; III-6) exhibited normal hearing at the age of twelve. The other affected male members had been diagnosed with hearing loss between the ages of ten to fifteen. Pfister had reported that there was gender difference in the severity of hearing loss in affected family members bearing the same *TECTA* mutations, with males being significantly more affected than females¹⁶. Therefore, there is a need for more detailed audiologic analysis and follow up in the other families to see whether they also show the same phenomenon in hearing impairment.

The present study further investigated whether the molecular mechanisms of hearing loss associated with *TECTA* mutations could be explained by protein expression. In contrast to COS-7 cells transfected with GFP-ZP wt, which were found to be localized in punctate spots along the plasma membrane (Figure 4B), the localization of GFP-ZP mutation proteins were not seen on the cellular membrane but mainly aggregated in the cytoplasm (Figure 4B). These mutations were located in the ZP domain of α -tectorin, this domain is responsible for secretion and polymerization of

extracellular proteins into supramolecular structure¹⁷⁻¹⁹. The results of these findings suggest that each missense mutation may lead to the lack of assembly of secretion, and may reduce the incorporation of α -tectorin into the tectorial membrane.

In this study, we have reported the prevalence of *TECTA* mutations in Japanese ADNSHL patients detected by genetic screening, and confirmed the genotype phenotype correlations. We also elucidated how mutation in the ZP domain of α -tectorin causes hearing loss through protein expression study of ZP domain proteins. *TECTA* mutation screening should be considered for patients with mild to moderate inherited autosomal dominant hearing loss because of its higher incidence. Further investigation of this gene is necessary to identify the function in the cochlea responsible for the distinct phenotype.

Acknowledgements

We sincerely thank the families for their participation in this study. And we also thank A. C. Apple-Mathews for help in preparing the manuscript. This work was supported by the Ministry of Health and Welfare, Japan (S.U.), and a Grant-in-Aid for Scientific Research from the Ministry of Education, Science and Culture of Japan (S.U.).

Figure legends

Figure 1

A: Pedigree of the family F818 and audiograms of four different patients. Black and white symbols indicate the affected and the unaffected subjects, respectively. B: Electropherograms for unaffected (wild type) and affected family members showing the heterozygous c.5318C>T mutation of *TECTA* co-segregating with hearing loss in this family. C: Audiograms of four different affected patients show high frequency hearing loss. Patient II-1 suffered decreasing hearing level in the right ear with cholesteatoma and postoperative change.

Figure 2

A: Pedigree of the family F237 and audiograms of four different patients. The marks and symbols are as described in Figure 1. B: Electropherograms for unaffected (wild type) and affected family members showing the heterozygous c.4198C>T and c.5597C>T mutations of *TECTA*. C: Multiple amino acid alignment of proteins homologous to the alpha-tectorin zona pellucida domain containing these mutated

positions. Amino acid residues that are identical among all of the homologs are enclosed.

D: Audiograms of four different affected patients showing deterioration in mid frequency as a U-shaped audiogram.

Figure 3

A: Pedigree of the family F652, B: Electropherograms for unaffected (wild type) and affected family members showing the heterozygous c.5990T>C mutation of *TECTA*. C: c.5990T>C is predicted to substitute isoleucine for threonine acid at amino acid position 1997. Multiple amino acid alignment of protein homologs was conserved. D: Audiograms of affected patients were flat to U-shaped, and there was a tendency to decreased hearing level associated with age.

Figure 4

A: Domain structure of the human alpha-tectorin protein. Three mutants in ZP domain protein including TM (Transmembrane domain) were generated by Site-directed mutagenesis. Resulting cDNAs digested with *EcoRI*/*Bam*HI and cloned into the site of

pEGFP-C2 plasmid are shown below. B: Protein expression in COS-7 cells transfected with GFP-ZP wild type (wt) showing a characteristic puncta along the plasma membrane. In contrast, GFP-ZP mutants (mut) C1837G, R2021H and Y1870C, which were associated with mid-frequency hearing loss phenotypes, were not recognized at the plasma membrane but were retained within the cytoplasm as indicated by the white arrows.

References

1. Smith R.J., Bale J.F., Jr., White K.R. Sensorineural hearing loss in children. *Lancet*. **365**, 879-890 (2005).
2. Petersen M.B., Willems P.J. Non-syndromic, autosomal-recessive deafness. *Clin Genet*. **69**, 371-392 (2006).
3. Verhoeven K., Van Laer L., Kirschhofer K., Legan P.K., Hughes D.C., Schatteman I., et al. Mutations in the human alpha-tectorin gene cause autosomal dominant non-syndromic hearing impairment. *Nat Genet*. **19**, 60-62 (1998).
4. Moreno-Pelayo M.A., del Castillo I., Villamar M., Romero L., Hernandez-Calvin F.J., Herraiz C., et al. A cysteine substitution in the zona pellucida domain of alpha-tectorin results in autosomal dominant, postlingual, progressive, mid frequency hearing loss in a Spanish family. *J Med Genet*. **38**, E13 (2001).
5. Iwasaki S., Harada D., Usami S., Nagura M., Takeshita T., Hoshino T. Association of clinical features with mutation of TECTA in a family with autosomal dominant hearing loss. *Arch Otolaryngol Head Neck Surg*. **128**, 913-917 (2002).
6. Plantinga R.F., de Brouwer A.P., Huygen P.L., Kunst H.P., Kremer H., Cremers C.W. A novel TECTA mutation in a Dutch DFNA8/12 family confirms genotype-phenotype correlation. *J Assoc Res Otolaryngol*. **7**, 173-181 (2006).
7. Sagong B., Park R., Kim Y.H., Lee K.Y., Baek J.I., Cho H.J., et al. Two novel missense mutations in the TECTA gene in Korean families with autosomal dominant nonsyndromic hearing loss. *Ann Clin Lab Sci*. **40**, 380-385 (2010).
8. Hildebrand M.S., Morin M., Meyer N.C., Mayo F., Modamio-Hoybjor S., Mencia A., et al. DFNA8/12 caused by TECTA mutations is the most identified subtype of nonsyndromic autosomal dominant hearing loss. *Hum Mutat*. **32**, 825-834 (2011).
9. de Heer A.R., Pauw R.J., Huygen P.L., Collin R.W., Kremer H., Cremers C.W. Flat threshold and mid-frequency hearing impairment in a Dutch DFNA8/12 family with a novel mutation in TECTA. Some evidence for protection of the inner ear. *Audiol Neurootol*. **14**, 153-162 (2009).
10. Balciuniene J., Dahl N., Jalonen P., Verhoeven K., Van Camp G., Borg E., et al. Alpha-tectorin involvement in hearing disabilities: one gene--two phenotypes.

- Hum Genet.* **105**, 211-216 (1999).
11. Legan P.K., Lukashkina V.A., Goodyear R.J., Kossi M., Russell I.J., Richardson G.P. A targeted deletion in alpha-tectorin reveals that the tectorial membrane is required for the gain and timing of cochlear feedback. *Neuron.* **28**, 273-285 (2000).
 12. Plantinga R.F., Cremers C.W., Huygen P.L., Kunst H.P., Bosman A.J. Audiological evaluation of affected members from a Dutch DFNA8/12 (TECTA) family. *J Assoc Res Otolaryngol.* **8**, 1-7 (2007).
 13. Sarkar G., Sommer S.S. The "megaprimer" method of site-directed mutagenesis. *Biotechniques.* **8**, 404-407 (1990).
 14. Govaerts P.J., De Ceulaer G., Daemers K., Verhoeven K., Van Camp G., Schattelman I., et al. A new autosomal-dominant locus (DFNA12) is responsible for a nonsyndromic, midfrequency, prelingual and nonprogressive sensorineural hearing loss. *Am J Otol.* **19**, 718-723 (1998).
 15. Moreno-Pelayo M.A., Goodyear R.J., Mencia A., Modamio-Hoybjor S., Legan P.K., Olavarrieta L., et al. Characterization of a spontaneous, recessive, missense mutation arising in the Tecta gene. *J Assoc Res Otolaryngol.* **9**, 202-214 (2008).
 16. Pfister M., Thiele H., Van Camp G., Franssen E., Apaydin F., Aydin O., et al. A genotype-phenotype correlation with gender-effect for hearing impairment caused by TECTA mutations. *Cell Physiol Biochem.* **14**, 369-376 (2004).
 17. Jovine L., Qi H., Williams Z., Litscher E., Wassarman P.M. The ZP domain is a conserved module for polymerization of extracellular proteins. *Nat Cell Biol.* **4**, 457-461 (2002).
 18. Qi H., Williams Z., Wassarman P.M. Secretion and assembly of zona pellucida glycoproteins by growing mouse oocytes microinjected with epitope-tagged cDNAs for mZP2 and mZP3. *Mol Biol Cell.* **13**, 530-541 (2002).
 19. Jovine L., Darie C.C., Litscher E.S., Wassarman P.M. Zona pellucida domain proteins. *Annu Rev Biochem.* **74**, 83-114 (2005).

Figure legends

Figure 1

A: Pedigree of the family F818 and audiograms of four different patients. Black and white symbols indicate the affected and the unaffected subjects, respectively. B: Electropherograms for unaffected (wild type) and affected family members showing the heterozygous c.5318C>T mutation of *TECTA* co-segregating with hearing loss in this family. C: Audiograms of four different affected patients show involving in high frequency hearing loss. The affected 1 patient suffered decreasing at right hearing with cholesteatoma and post operative change.

Figure 2

A: Pedigree of the family F237 and audiograms of four different patients. The marks and symbols are as described in Figure 1. B: Electropherograms for unaffected (wild type) and affected family members showing the heterozygous c.4198C>T and c.5597C>T mutation of *TECTA*. C: Multiple amino acid alignment of proteins homologous to the alfa-tectorin zona pellucida domain containing these mutated

position. Amino acid residues that are identical among all of the homologs are indicated in square. D: Audiograms of four different affected patients showing deteriorate in mid frequency as a “U-shape” audiogram.

Figure 3

A: Pedigree of the family F652, B: Electropherograms for unaffected (wild type) and affected family members showing the heterozygous c.59990T>C mutation of *TECTA*.

C: c.5990T>C is predicted to substitute isoleucine for threonine acid at amino acid position 1997. Multiple amino acid alignment of proteins homologous was conserved.

D: Audiograms of affected patients show flat to U-shape audiogram, and a tendency to decrease hearing level associate with age.

Figure 4

A: Domain structure of the human alfa-tectorin protein. Three mutants in ZP domain

protein including TM (Transmembrane domain) were generated by Site-directed mutagenesis. Resulting cDNAs digested with *XhoI*/*Bam*HI and cloned into the site of pEGFP-C2 plasmid is shown below. B: Protein expression in transfected COS-7 cells. COS-7 cells transfected with GFP-ZP wt showed a characteristic puncta along the plasma membrane. In contrast, GFP-ZP mut C1837G, R2021H and Y1870C, which were associated with mid-frequency hearing loss phenotypes, were not recognized at the plasma membrane but was retained within the cytoplasm.

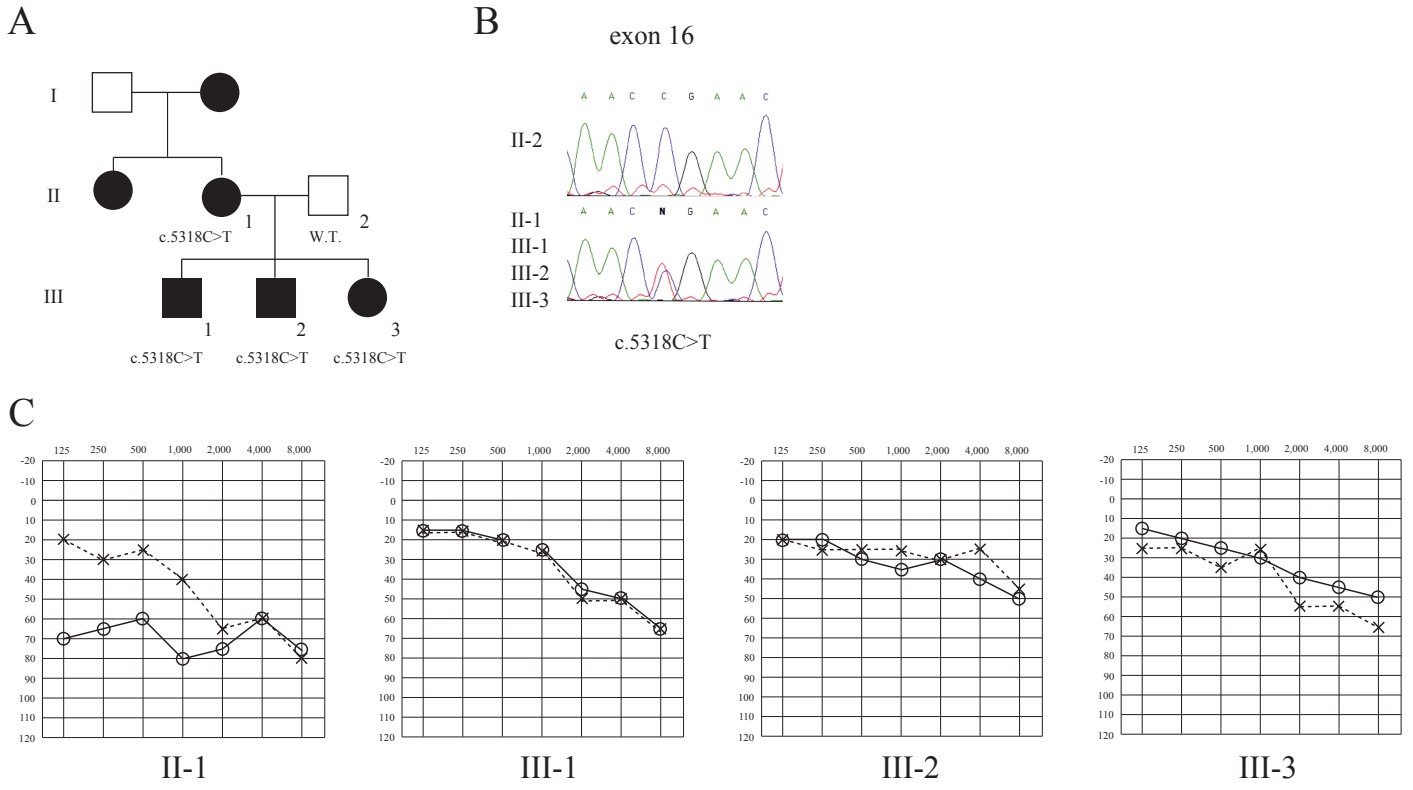


Figure 1

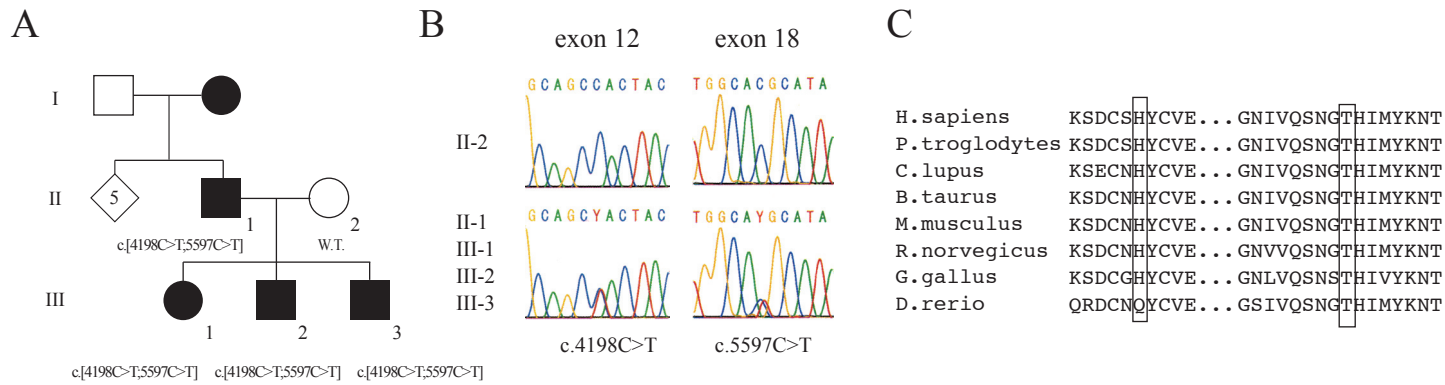
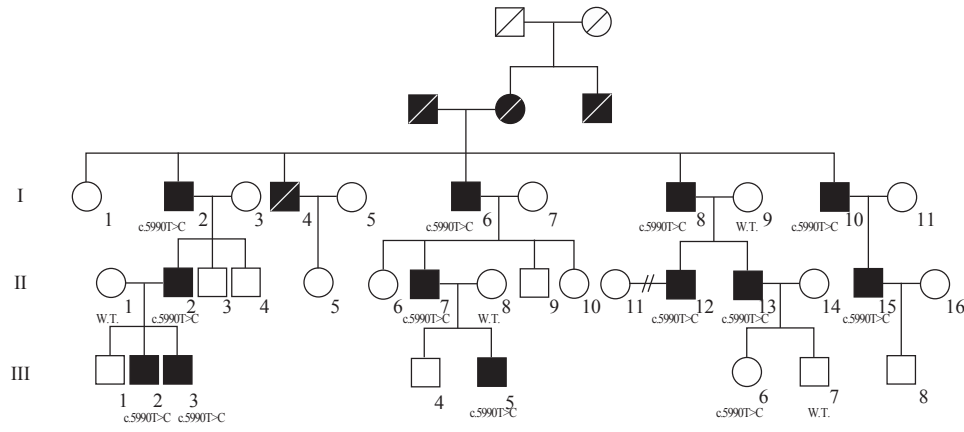
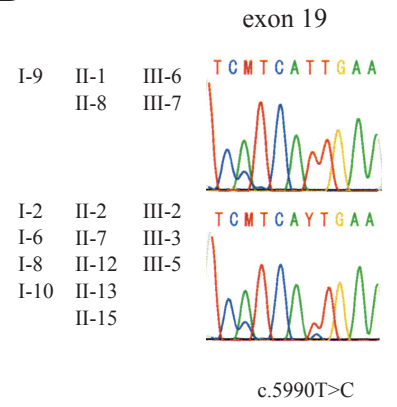


Figure 2

A



B



C

H. sapiens	DKLRYFIIIEGGQNL
P. troglodytes	DKLRYFIIIEGGQNL
C. lupus	DKLRYFIIIEGGQNI
B. taurus	DKLRYFIIIEGGQNI
M. musculus	DKLRYFIIIEGGQNI
R. norvegicus	DKLRYFIIIEGGQNI
G. gallus	DKLRYFIIIEGGQNL
D. rerio	DRLRYIIIEGCPNI

D

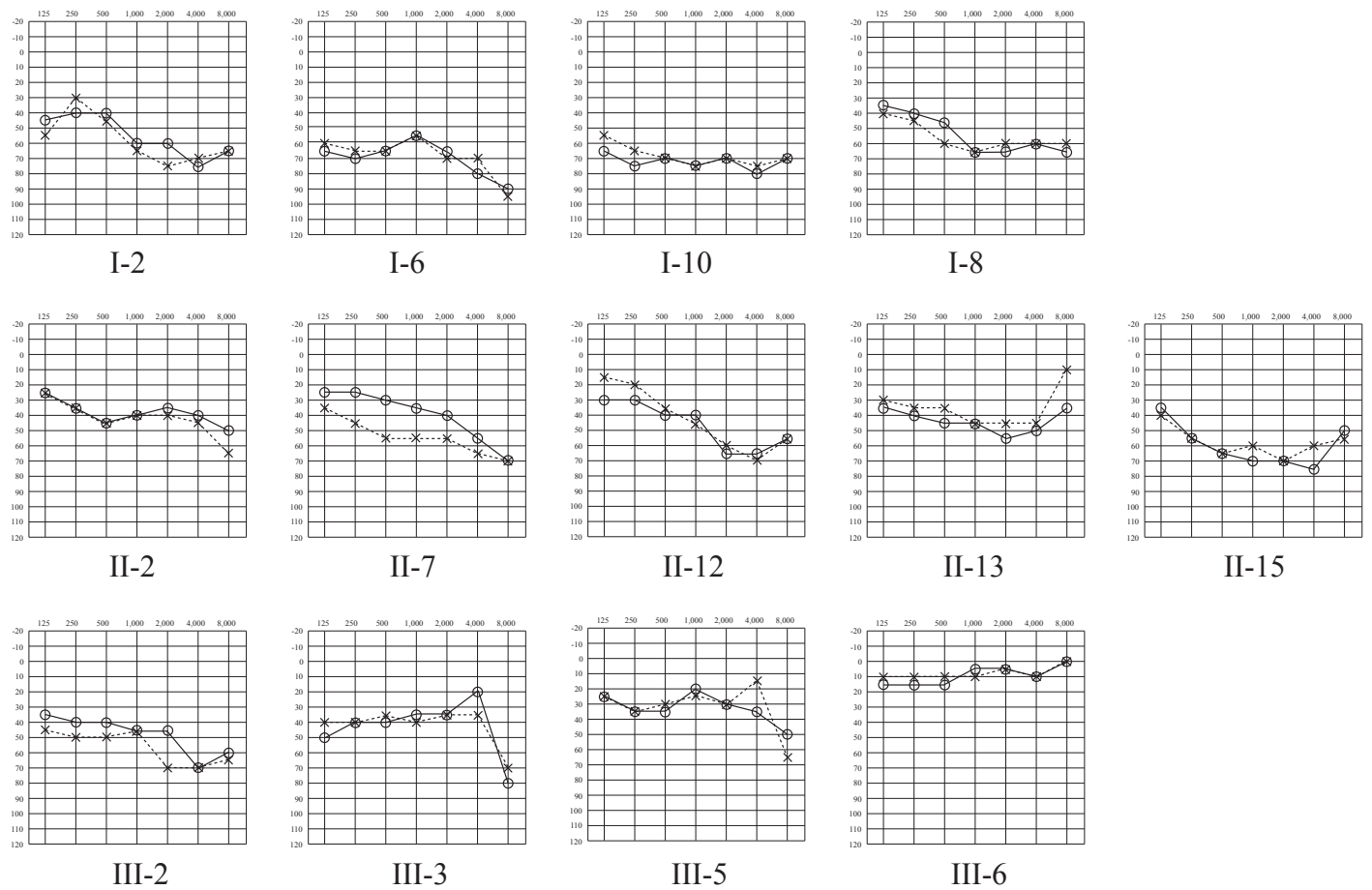
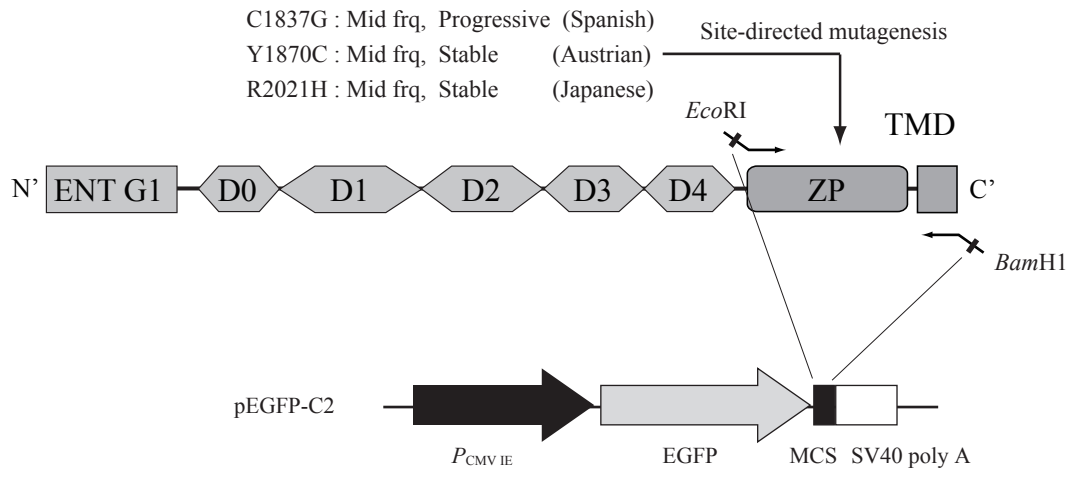


Figure 3

A



B

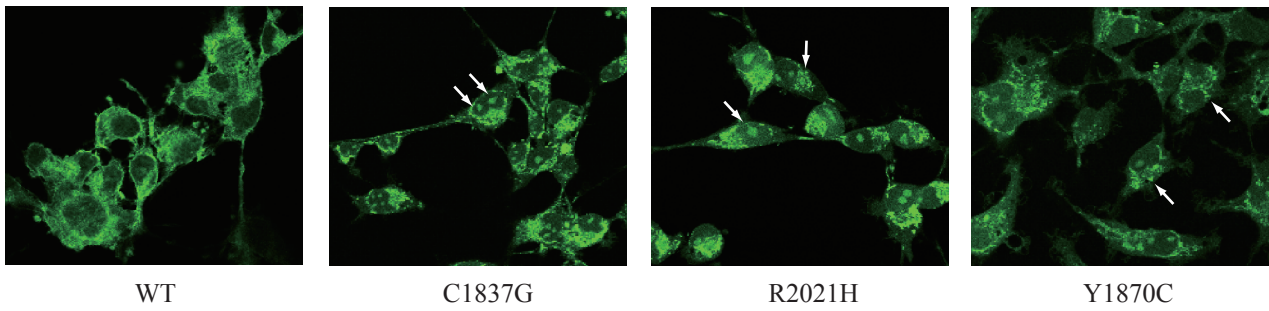


Figure 4