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Connexin 26 mutations in cases of sensorineural deafness in eastern Austria

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Mutations in the connexin 26 (Cx26) gene (GJB2) are associated with autosomal nonsyndromic sensorineural hearing loss. This study describes mutations in the Cx26 gene in cases of familial and sporadic hearing loss (HL) by gene sequencing and identifies the allelic frequency of the most common mutation leading to HL (35delG) in the population of eastern Austria. For this purpose we have developed and applied a molecular beacon based real-time mutation detection assay. Mutation frequencies in the Cx26 gene of individuals from affected families (14 out of 46) and sporadic cases (11 out of 40) were 30.4% and 27.5%, respectively. In addition to known disease related alterations, a novel mutation 262 G→T (A88S) was also identified. 35delG accounted for almost 77% of all Cx26 mutations detected and displayed an allelic frequency in the normal hearing population of 1.7% (2 out of 120). The high prevalence of the 35delG mutation in eastern Austria would therefore allow screening of individuals and family members with Cx26 dependent deafness by a highly specific and semi-automated method. European Journal of Human Genetics (2002) 10, 427 – 432. doi:10.1038/sj.ejhg.5200826

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Introduction

Congenital hearing loss affects 1 in 1000 children^{1,2} and about 60% of all cases are estimated to be of genetic origin.³ In approximately 80% of cases, the hearing impairment is inherited in an autosomal recessive fashion.⁴ More than 100 genes are estimated to be involved in hearing impairment and to date 60 loci (eg⁵⁻⁷) have been described in previous studies and 21 related genes have been cloned

including Connexin 26 $(Cx26)^8$, Connexin 31^9 and alpha tectorin. 10

Connexins are a large family of gap junction forming proteins involved in direct cell-to-cell transfer of small molecules and ions. Cx26 is thought to be important for the recycling of endolymphatic potassium ions. $^{11-13}$

Mutations in the gap junction protein Cx26 play an important role in autosomal nonsyndromic sensorineural HL. Several mutations have been identified which display a variable population distribution^{14–19} and can lead to autosomal dominant^{20,21} and recessive forms of HL. The 35delG recessive frame shift mutation is the most frequent alteration in European populations with carrier frequencies of 1 out of 35 in southern and 1 out of 79 in northern

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Europe.²² 35delG is one of the most frequent pathogenic mutations in humans and displays a carrier frequency similar to the deltaF508 mutation of the cystic fibrosis transmembrane conductance regulator gene (CFTR).²³

In this study, chromosomal sequencing of familial and sporadic cases of nonsyndromic hearing impairment was performed to identify mutations in the Cx26 gene. The 35delG mutation was the most common alteration found in the Cx26 gene. Five further sporadic mutations were additionally found, one of which has not yet been described. To estimate the 35delG carrier frequency in the eastern Austrian population, a simple and rapid screening procedure was developed using real time PCR with molecular beacons which represents a highly sensitive, semiautomated method for the detection of 35delG which may be used in routine, clinical diagnosis.

Methods

Individuals suffering from sensorineural hearing impairwere recruited from the Department Otorhinolaryngology, University hospital of Vienna and informed consent was obtained from all individuals or the parents of minors. The complete medical histories of all individuals were examined and included otolaryngological status and pure tone audiometrical analysis. Cases of acquired HL and syndromic forms were excluded from the study. In several cases, otoacoustic emissions (OAE), brainstem evoked response audiometry (BERA), ophthalmic evaluations, computer tomography and magnetic resonance imaging of temporal bone and blood analyses were performed to ascertain disease status.

Individuals (n=57) from families (n=23) with at least two affected individuals and sporadic cases (n=20) with no further case of familial deafness were recruited into the study. Inheritance of bilateral deafness was autosomal recessive in 13 familial cases and dominant in seven families and unclear in three cases. HL was classified as moderate in seven families, severe in three and profound in 13 families, congenital in 18 familial cases and postlingual in five families. All 20 sporadic cases (3-42 years; mean 19.8) included in this study demonstrated bilateral moderate to profound congenital HL with no signs of progression. The allelic frequency of the 35delG mutation was determined in 120 unrelated, unaffected Austrian individuals. The DNA samples were randomised and showed a normal sex distribution.

Screening of the Cx26 gene by direct PCR-sequencing

Genomic DNA was extracted from peripheral blood with a E.Z.N.A. Blood DNA Kit II kit (peqlab, Erlangen, Germany). The Cx26 gene was amplified in a Perkin-Elmer thermal cycler using the primer sets Cx26f, Cx26r as described in Table 1 for 30 cycles with an annealing temperature of 55°C for 30 s, extension at 72°C for 60 s and denaturation at 94°C for 30 s as described. 18 Direct DNA-sequencing on both strands was performed with the PCR primers and the

Table 1 Sequences of amplification primers for molecular beacon PCR-reactions and Connexin 26 whole sequence-

Primers	Sequence			
Cx26fw	5'-CCGCCCAGAGTAGAAGATGGAT-3'			
Cx26rv	5'-GATCTTTCCAATGCTGGTGGAG-3'			
Cx26f	5'-TCTTTTCCAGAGCAAACCGCC-3'			
Cx26r	5'-TGAGCACGGGTTGCCTCATC-3'			
Sequences o	f molecular beacons			
Molecular				
beacon	Sequence			
WT-24-41	41 HEX-5'-GGCTGG TTCACACCCCCAGGATC CCAGCC-DABCYE			
MT-35delG	FAM-5'-GGCTGG TTCACACCCCAGGATC CCAGCC-3'-DABCYL			

Molecular beacon stem regions (underlined) and loop sequences (italics) are shown.

internal forward Cx26fi and reverse Cx26ri primers (see Table 1) on an ABI-377-DNA-Sequencer and aligned to the wild type sequence with GENERUNNER-Software.

Design of molecular beacons

Molecular beacon probes were designed following the guidelines described by Tyagi and Kramer²⁴ available at the related website (www.phri.nyu.edu/molecular beacons). The intramolecular configuration of molecular beacons and the amplicon probing were modelled using the DNA Mfold program (www.ibc.wustl.edu/-zuker/dna/form1.shtml) that uses thermodynamic parameters established by SantaLucia. 25 Melting temperatures of probe sequences were calculated using Primer software and molecular beacons were synthesised and purified as described.²⁶ The temperature at which there is no contribution of mutant molecular beacon to the fluorescence from wild type target and of wild type molecular beacon to mutant target was determined as described.²⁴

Thermal denaturation profiles

Fluorescence was measured as a function of temperature using an ABI Prism 7700 spectrofluorometric thermal cycler. For mutant (MUT-FAM) and wild type (WT-HEX) molecular beacons, three tubes were prepared. Two of the tubes contained target oligonucleotide sequence complementary to the loop, either wild type (WT-ODN) or mutant (MT-ODN). The third tube did not contain target oligonucleotide (NO-ODN). The temperature was increased from 15°C to 85°C with 1°C steps and holds for 60 s each. Laser light of 488 nm was used for excitation and the fluorescence was monitored during the second 30 s of each step (see Figure 3). Measurement data were normalised as described.²⁷

Molecular beacon-PCR conditions

Two-colour molecular beacon PCR reactions were carried out in the presence of 6-carboxyfluorescein (FAM)-labelled mutant and hexacholo-6-carboxyfluorescein (HEX)-labelled wild type molecular beacons. Amplification reactions were performed in an ABI Prism 7700 spectrofluorometric thermal cycler (Applied Biosystems) with the following cycle conditions: 95°C denaturation and enzyme activation step for 10 min followed by 40 cycles of 95°C denaturation for 15 s, 62°C annealing for 30 s and 72°C elongation for 30 s. Fluorescence spectra were recorded during the annealing phase of each PCR cycle. The Sequence Detection Software (SDS v1.7) of the ABI-Prism 7700 was used to generate the amplification curves. The threshold cycle number (Ct) was calculated with Sequence Detection System (SDS) software v1.7 (Applied Biosystems, The Netherlands) and an automatic setting of the baseline. The baseline value was the average fluorescence value of PCR cycles 3 to 15 plus 10 times its standard deviation (SD). Standard/calibration curves were generated by plotting the Ct values against the log of the input copy numbers.

Results

Mutation frequency

Analysis of the complete coding region of Cx26 of all individuals (n=57) showed that 29% of unrelated individuals had mutations in Cx26. In four families and four sporadic cases, the 35delG mutation was homozygous. Hearing impaired individuals with a 35delG mutation showed a non-progressive HL. In all cases 35delG associated deafness was severe to profound and the degree of HL in both ears was similar. The eight homozygous unrelated individuals represent 18.6% of the hearing impaired population studied and 35delG accounted for 76.9% of mutated Cx26 alleles. Five further mutations were detected (see Figure 1). The previously described mutation L90P^{28,29} was found in one allele in a case of sporadic deafness. The delE119 alteration, which was originally described in combination with a 167delT mutation,²⁸ was found in a sporadic case as a compound heterozygote with 35delG. The 71G→A mutation was also found as a homozygote in one family.8 457A→G, which has been reported as a polymorphic alteration was also found in one family³⁰ where the nonsymptomatic father and one of the two hearing impaired daughters were carriers.

Novel mutation

In addition to previously described mutations, a novel alteration in Cx26 was found. A heterozygous 262G→T mutation, which would lead to the substitution of an alanine for a serine in the Cx26 protein sequence, was found in a family in which the mother is a non-symptomatic carrier and the male offspring is affected. The second offspring in this family is a non-symptomatic female with wild type genotype (Figures 1 and 2).

Determination of melting and optimal hybridisation temperature to detect the 35delG mutation

Molecular beacon probes fluorensce upon hybrid formation with target sequences. The maximally discriminating hybridisation temperature is dependent on several factors, such as GC content, probe (loop) and stem length. In order to determine the optimal temperature range at which a mole-

Mutation Name Kind	Position of nucleic acid	Unrelated familial cases Number of cases and genotype	Sporadic cases Number of cases and genotype	Total (% of all mutated alleles)
35delG Frameshift	Deletion of one G at 30-35	11/46 4 Ho: 35delG/35delG 3 Het: 35 delG /WT	9/40 4 Ho: 35delG/35delG 1 Het: 35delG/ DEL E 119*	76.9
W24X Stopcodon	71G>A	2/46 1 Ho: 71G>A/71G>A		7.7
A88S Missense	262 G>T	1/46 1 Het: 262 G>T/WT		3.8
L90P Missense	269 T>C		1/40 1 Het: 269 T>C/WT	3.8
DEL E 119 Deletion	Deletion of GAG at 355-357		1/40 1 Het: 35delG/ DEL E 119*	3.8
V153I Missense * Same inc	457 G>A	1/46 1 Het: V153I/WT		3.8

Figure 1 Mutations identified in the Cx26 gene in cases of non-syndromic hearing loss (n=57). The nucleotide and amino acid positions of all mutations found in familial and sporadic cases, the genotype and occurrence of these alleles are shown.

cular beacon probe discriminates between wild type and mutant targets, 'melt-measurements' of the probe/ target combinations were performed. The melt characteristics of duplexes between the molecular beacon and the

TCCAC GC CAKC GC TCCTAGT

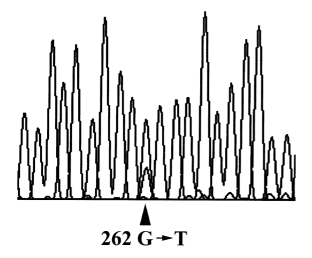


Figure 2 DNA sequence analysis of the Cx26 gene in the 262 G→T region is shown. The heterozygous sequence was confirmed by sequencing from both sides (reverse sequence not shown).

different target oligodeoxynucleotide sequences are shown in Figure 3.

On the basis of the melt measurements, we concluded that at 62°C there is a minimal contribution of mutant molecular beacon to the fluorescence from wild type target and of wild type molecular beacon in the case of mutant target (vertical arrow in Figure 3). Therefore, 62°C was used as the annealing temperature and primers were designed for optimal priming at this temperature. At the extension temperature of 72°C, the probe dissociates from the target preventing degradation of molecular beacons by the 5'-3' exonuclease activity of the Taq DNA polymerase. At higher temperatures, the fluorescence intensities equal those of the molecular beacon alone, which adopts a random coil configuration at about 75°C as a consequence of the melting of the stem.

Automated 35delG detection

To enable high throughput, routine diagnostic detection of the Cx26 35delG mutation, a simple and rapid method using molecular beacon based real-time fluorescence PCR was developed. Real-time PCR screening for the Cx26 35delG mutation in the study of individuals and their family members with real-time PCR produced identical results to those obtained by direct sequencing. A representative graphical illustration is shown in Figure 4 where circles represent genotypes; the position of each spot on the X;Y axes varies because of various amounts of DNA loaded. This method was then used to analyse 120 randomised samples for 35delG in the eastern Austrian population. Analysis demonstrated the presence of two

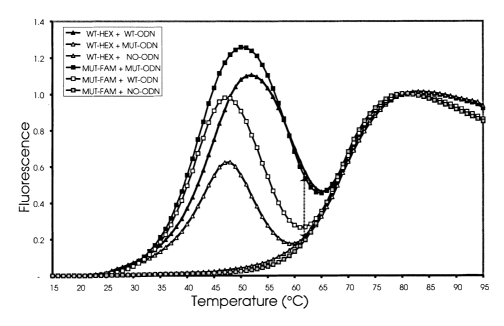


Figure 3 Thermal denaturation profiles of 35delG molecular beacons in the presence of perfectly matching, imperfectly matching and in the absence of complementary oligonucleotide target sequences. FAM labelled mutant and HEX labelled wild type molecular beacons.

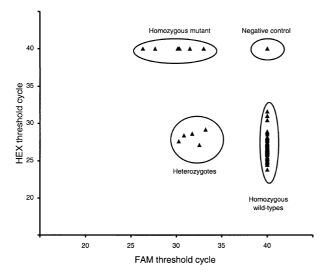


Figure 4 Spectral genotyping by molecular beacon based PCR for the 35delG mutation. Graphical representation of thresholdcycle values of HEX (wild-type molecular beacon, y) vs FAM (mutant molecular beacon, x) on a selected set of 96 samples analysed and including a negative control.

heterozygous samples in the sample population representing a relative carrier frequency of 1.7%.

Discussion

In this study we investigated the prevalence of Cx26 mutations in cases of familial and sporadic sensorineural HL and determined the allelic frequency of the 35delG mutation in the population of eastern Austria. Here we show, that mutations in the Cx26 gene are a main reason for inherited hearing impairment in eastern-Austrian individuals and Cx26 mutations were present in 29.1% of sufferers.

The 35delG frequency determined was similar to other closely related populations. 14,22 In this study, the 35delG mutation was responsible for 76.9% of changes in the Cx26 gene and the allelic frequency of this mutation in familial (23.9%) and sporadic cases (22.5%) was almost equal. Our findings suggest that the high prevalence of the 35delG mutation in the population of the eastern part of Austria may allow diagnosis and genetic counselling. The prevalence of 35delG can be compared with one of the most frequent disease mutations known so far, the delta F508 mutation of the CFTR gene, which leads to cystic fibrosis and for which a heterozygotic advantage has been suggested.³¹ These data show the importance of further investigations. Future genetic counselling for the 35delG mutation would require a simplified method for routine detection. The standard technique for the detection of 35delG is DNA sequencing, which is prone to sequencing errors due to the expansion of the same nucleotide (six guanine residues). In addition, this approach is cost and labour intensive. Several methods such as single-strand

conformational polymorphism (SSCP),²⁸ allele specific oligonucleotide (ASO) analysis³² or sandwich hybridisation³³ have been described for detecting 35delG. All these methods require further steps following PCR such as hybridisation, electrophoresis or the use of radioactive detection. Real time PCR with molecular beacons has the advantage of an easy and single semi-automated step following DNA isolation and has exceptional discriminatory power for single base mismatches due to the stem-loop structure of the molecule.²⁴ Handling is similar to common PCR-methods except for the addition of molecular beacons and a large amount of samples can be analysed simultaneously within a few hours. In addition, real time PCR is a closed tube format assay preventing carry-over contamination and provides a direct readout of results after PCR. In this study, we have shown that molecular beacons can reliably detect the 35delG mutation by comparison to the fully sequenced Cx26 gene. All results obtained from sequencing could be confirmed by this method even with variable amount of initial sample load which makes this newly established assay a highly sensitive, cheap and easy-to-use tool for the detection of 35delG.

In addition to the 35delG mutation, five different point mutations were detected of which one was novel. A $G \rightarrow T$ transition at nucleotide 262 would result in the substitution of an alanine for a serine at position 88 (A88S), a residue conserved in mouse, rat and sheep Cx26. A dominant mutation at the same position in the connexin 30 gene causes hidrotic ectodermal dysplasia which emphasises the importance of this region.34 A88S would lead to an additional hydroxyl group in the second transmembrane domain, which would increase hydrophilicity and reactivity in this region. This novel missense mutation could not be causally connected to deafness and therefore a simple, monogenetic disease model can be excluded. The delE119 mutation is a previously described deletion of a triplet of nucleotides (GAG) at positions 355-357, which leads to the deletion of glutamic acid at amino acid position 119.28 In this study, delE119 was found as a compound heterozygote with 35delG. The homozygous 71A→G point mutation was found in one family and leads to a premature stop codon as described.⁸ The L90P mutation was detected in only one sporadic case of deafness and was heterozygous.^{28,29} This finding contrasts to the high incidence of L90P (16.7%) which has been found in the Tyrolean population. 14 A G -> A transition which would lead to the substitution of a valine residue for isoleucine at position 153 (V153I), which is located in the second extracellular loop and is also highly conserved in mouse, rat and sheep Cx26 has been described as a polymorphic change.³⁰

This study emphasises the importance of Cx26 mutations in inherited cases of sensorineural deafness in eastern Austria. Six different mutations were identified of which one was a novel alteration and the 35delG mutation was found to occur in almost 77% of individuals suffering from

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deafness. We have developed a simple method to detect this most common disease causing Cx26 mutation, which will allow rapid high throughput screening to enable genetic counselling and the determination of allelic frequencies in populations.

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