

ORIGINAL ARTICLE

Update of the *GJB2*/DFNB1 mutation spectrum in Russia: a founder Ingush mutation *del(GJB2-D13S175)* is the most frequent among other large deletions

Elena A Bliznetz¹, Maria R Lalayants², Tatiana G Markova², Oleg P Balanovsky^{1,3}, Elena V Balanovska¹, Roza A Skhalyakho^{1,3}, Elvira A Pocheshkhova⁴, Natalya V Nikitina⁵, Sergey V Voronin⁶, Elena K Kudryashova⁷, Oleg S Glotov⁸ and Alexander V Polyakov¹

Although mutations in the *GJB2* gene sequence make up the majority of variants causing autosomal-recessive non-syndromic hearing loss, few large deletions have been shown to contribute to DFNB1 deafness. Currently, genetic testing for DFNB1 hearing loss includes *GJB2* sequencing and DFNB1 deletion analysis for two common large deletions, *del(GJB6-D13S1830)* and *del(GJB6-D13S1854)*. Here, we report frequency in Russia, clinical significance and evolutionary origins of a 101 kb deletion, *del(GJB2-D13S175)*, recently identified by us. In multiethnic cohort of 1104 unrelated hearing loss patients with biallelic mutations at the DFNB1 locus, the *del(GJB2-D13S175)* allele frequency of up to 0.5% (11/2208) was determined and this allele was shown to be predominantly associated with profound sensorineural hearing loss. Additionally, eight previously unpublished *GJB2* mutations were described in this study. All patients carrying *del(GJB2-D13S175)* were of the Ingush ancestry. Among normal hearing individuals, *del(GJB2-D13S175)* was observed in Russian Republic of Ingushetia with a carrier rate of ~1% (2/241). Analysis of haplotypes associated with the deletion revealed a common founder in the Ingushes, with age of the deletion being ~3000 years old. Since *del(GJB2-D13S175)* was missed by standard methods of *GJB2* analysis, *del(GJB2-D13S175)* detection has been added to our routine testing strategy for DFNB1 hearing loss.

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INTRODUCTION

Pathogenic variants in the DFNB1 locus (MIM#220290) are the most commonly identified cause of congenital, recessively inherited, sensorineural non-syndromic hearing loss. This locus contains the *GJB2* gene, encoding the connexin 26 protein. The mutation spectrum of *GJB2* varies in different populations. There are ethnic-specific frequent mutations in the *GJB2* gene as a result of the founder effect.¹ Although, more than 300 mutations in the *GJB2* gene sequence have been described and make up the majority of causative variants, six large deletions have been shown to contribute to DFNB1 deafness (The Human Gene Mutation Database or HGMD, Professional 2016.2). DFNB1 patients carry a large deletion in *trans* with a *GJB2* variant or they are homozygous (or double heterozygous) for the large deletion(s).^{2–10}

Two mechanisms of pathogenesis for the large DFNB1 deletions have been hypothesized. Four large deletions lie upstream of *GJB2* and do not affect *GJB2*, but were shown to reduce *GJB2* expression by disrupting unidentified *cis*-regulatory element. Presumably, *cis* regulatory element is located within the common 95.4 kb genomic interval that is deleted in all four mutant alleles (Figure 1).^{7,11–14} This interval encompasses the upstream *CRYL1* gene and extends telomerically. The most common deletions being 309 and 232 kb, *del(GJB6-D13S1830)* and *del(GJB6-D13S1854)*, respectively, truncate the upstream *GJB6* and *CRYL1* genes. Two private DFNB1 deletions of 131 and 179 kb involve *CRYL1* only.^{7,9} Two additional mutations directly affect the *GJB2* gene by deletion of the entire *GJB2* sequence. One of them is 920 kb in size involves seven more other genes and was found in only one deaf individual.⁷ The second deletion of 101 kb,

¹The Federal Agency for Scientific Organizations, Federal State Budgetary Institution, Research Centre for Medical Genetics, Moscow, Russia; ²Federal Medical and Biological Agency, Federal State-Funded Institution of Science, National Research Center for Audiology and Hearing Rehabilitation, Moscow, Russia; ³The Russian Academy of Sciences, Federal State-Funded Institution of Science, Vavilov Institute of General Genetics, Moscow, Russia; ⁴The Ministry of Health, Federal State-Funded Educational Institution of Higher Education, Kuban State Medical University, Krasnodar, Russia; ⁵The Ministry of Health, State Budgetary Healthcare Institution, Clinical-Diagnostic Center of Mother and Child Health Protection, Yekaterinburg, Russia; ⁶State Autonomous Healthcare Institution, Regional Clinical Center of Specialized Medical Care, Vladivostok, Russia; ⁷The Ministry of Health, State Budgetary Healthcare Institution, Leningrad Regional Clinical Hospital, Saint Petersburg, Russia and ⁸Federal State Budgetary Educational Institution of Higher Education, Saint Petersburg University, Saint Petersburg, Russia

Correspondence: Dr EA Bliznetz, Federal State Budgetary Institution 'Research Centre for Medical Genetics', Moskvorechie street, 1, Moscow 115478, Russia.
E-mail: bliznetzelena@mail.ru

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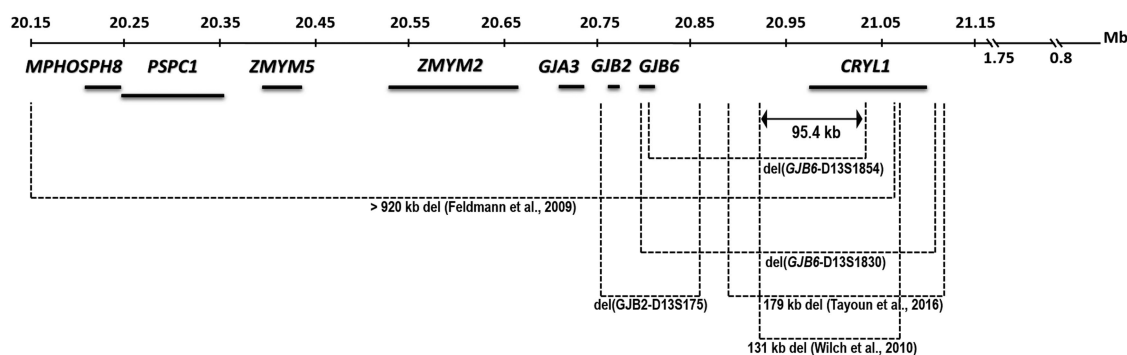


Figure 1 Schematic map of the six large deletions described at DFNB1 locus on chromosome 13q11-12. The DNA section encompasses eight genes (bold horizontal lines), the breakpoints of the deletions (dotted brackets). All elements are drawn approximately to scale. The del(*GJB2*-D13S175) mutation was investigated in this study. All genomic coordinates are based on Human Genome Build GRCh37.p13/hg19.

del(*GJB2*-D13S175), encompassing *GJB2* and *GJB6*, which we described recently, has been identified in three unrelated Russian patients.¹⁰

Currently, clinical DFNB1 testing for non-syndromic hearing loss patients includes *GJB2* sequencing and DFNB1 deletion analysis for only the two common deletions del(*GJB6*-D13S1830) and del(*GJB6*-D13S1854). These deletions account for 1–10% of all the DFNB1 alleles in most populations in Europe, America and Australia, but the del(*GJB6*-D13S1830) mutation is much more frequent than del(*GJB6*-D13S1854).^{5,6,15–17} All the studied chromosomes carrying the del(*GJB6*-D13S1854) mutation share a common founder, while two founders for del(*GJB6*-D13S1830) have been demonstrated.^{5,6}

The frequency of del(*GJB6*-D13S1830) observed among Russian patients is 0.3% of the DFNB1-mutant alleles, whereas del(*GJB6*-D13S1854) has not been found.^{18–21} Previously, we have published the spectrum of *GJB2*/DFNB1 in 705 Russian patients, but this study does not include the testing for recently found mutation del(*GJB2*-D13S175).¹⁹ Here, update findings of investigation of a larger cohort of hearing loss patients from Russia are present. We report the high frequency, clinical significance and the evolutionary origins of del(*GJB2*-D13S175). Additionally, eight novel *GJB2* mutations were found.

MATERIALS AND METHODS

Study cohort and population samples

We analyzed genomic DNA samples from 2569 unrelated individuals with non-syndromic sensorineural hearing loss that were referred for DNA analysis of *GJB2*/DFNB1 at the Laboratory of DNA Diagnostics, Research Centre for Medical Genetics, Moscow, between years 2000 and 2015, including the patients from cohort that has previously been reported.¹⁹ DNA analysis of *GJB2* was performed on all patients as previously described.¹⁹ Briefly, this is a two-step protocol starting with the common point mutations test, followed by exon 2 sequencing analysis in the cases when one or no common mutation has been revealed. Then samples with one heterozygous mutation were tested for del(*GJB6*-D13S1830). The common mutation test lay in two reactions, a multiplex PCR with primer pairs for five deletions (c.35delG, c.313_326del14, c.235delC, c.167delT and c.358_360delGAG) and restriction fragment length polymorphism (RFLP) analysis of c.-23+1G>A, followed by gel-based fragment analysis to differentiate wild-type and mutation alleles. The nomenclature of all *GJB2* sequence variants was based on complementary DNA reference sequence of the NM_004004.5 transcript.

The study cohort consisted of 1101 patients carrying biallelic mutations in the *GJB2* gene, 1465 individuals who carry one or no pathogenic *GJB2* variant, 3 patients in which the PCR product was not amplified with *GJB2*- and D13S175-specific primers. One patient 'homozygous' for c.35delG, but whose father had no *GJB2* mutations and two of three patients without sequences

of the *GJB2* and D13S175 were described in previous article about the del(*GJB2*-D13S175) identification.¹⁰ Among patients with homozygous *GJB2* mutation, there were 691 patients whose parents' DNA were not available.

The population samples have been selected from the Biobank of North Eurasia.²² The biobank's samples were collected in 1998–2011 during the extensive field trips to the rural areas. The samples analyzed in this study came from the two regions of Russia: Republic of Ingushetia and Chechen Republic. The sampled persons represented the corresponding indigenous rural populations up to the third generation, that is, all four grandparents belonged to the given ethnic groups and were born within the given region. The population samples included 241 persons from Ingushetia Republic (151 self-identified Ingushes and 90 self-identified Chechens), and 147 self-identified Chechens from Chechen Republic.

All written informed consent forms signed by the participants or the guardians of the under age participants involved in our study were obtained before the testing procedures. This study was approved by the local Ethics Committee of Federal State Budgetary Institution 'Research Centre for Medical Genetics', Moscow, Russia.

del(*GJB2*-D13S175) deletion test

The del(*GJB2*-D13S175) deletion test was performed using duplex PCR with deletion-specific breakpoint primers (F—5'-GCTCTGCCAGATGAAGATCTC-3', R—5'-CCTTCCAGGAGAGTTCACAACCTC-3') and *GJB2* exon 2 specific primer pair to amplify a PCR control fragment (F—5'-GTGATTCCTGTGTTGTGTGCATTC-3', R—5'-CCTCATCCCTCTCATGCTGTC-3'). The amplified products were separated by polyacrylamide gel electrophoresis with subsequent ethidium bromide stain and ultraviolet visualization.

STRs analysis and del(*GJB2*-D13S175) dating

Three short tandem repeats (STRs) flanking del(*GJB2*-D13S175) were genotyped by PCR amplification and size analyses of the amplified products by polyacrylamide gel electrophoresis. The STRs and the primers used for amplification are following: D13S1316F—5'-GATTGCACCACTACATACCA GC-3', D13S1316R—5'-CTTTGACTCTCCATGCTGCATTC-3', D13S1275F—5'-GCTAGTCTTCAGATTACCTTAGAATATACC-3', D13S1275R—5'-CAGCA TGAACTTTACCAGAATTCC-3', D13S232F—5'-GCTCACTGCTCTTGTTGATTCTG-3', D13S232R—5'-GGCACAGAAATAAATGTTGATGATGTAC-3'. The STR allele designation corresponds to appropriate repeat number, which was fixed by the control DNA samples, previously sequenced by using the ABI Dye Terminator, version 1 and run on a 3130 ABI genetic analyzer (Applied Biosystems, Tokyo, Japan). The STR genotyping was performed in 8 patients with del(*GJB2*-D13S175), in 6 their parents and in 90 healthy Ingush individuals. Haplotypes were imputed from family analysis data. To estimate the age of del(*GJB2*-D13S175), we followed the approach that have been offered and previously proven.^{23–26} STATISTICA 10 (StatSoft. Inc. (2011), Tulsa, OK, USA) was used for other calculations.

RESULTS

A novel *GJB2* mutation

A total of 39 pathogenic or likely pathogenic variants in *DFNB1* locus were found in Russian cohort of patients with non-syndromic sensorineural hearing loss, including previously published¹⁹ and novel samples collected between year 2000 and 2015 in the Laboratory of DNA Diagnostics. In Table 1, an allele frequency of these variants and references are present. Among the identified variants, eight mutations have for the first time been described in this work. These are five likely pathogenic missense substitutions, two pathogenic nonsense

Table 1 *DFNB1* alleles in Russian patients

Mutation	Effect	% (number) of alleles among 2208 chrs.	Reference ^a
c.35delG	p.Gly12Valfs*2	77.0 (1700)	CD972240
c.313_326del14	p.Lys105Glyfs*5	4.7 (103)	CD991732
c.-23+1G>A	IVS1+1G>A	4.4 (97)	CS991407
c.101T>C	p.Met34Thr	2.9 (65)	CM970679
c.235delC	p.Leu79Cysfs*3	2.3 (51)	CD991730
c.167delT	p.Leu56Argfs*25	1.5 (33)	CD972241
c.109G>A	p.Val37Ile	1.2 (27)	CM000016
c.269T>C	p.Leu90Pro	1.0 (23)	CM990691
c.358_360delGAG	p.Glu120del	0.9 (19)	CD993053
c.290dupA	p.Tyr97*	0.7 (15)	CI014787
del(<i>GJB2</i> -D13S175)	<i>GJB2</i> - <i>GJB6</i> deletion	0.5 (11)	CG145665
c.551G>C	p.Arg184Pro	0.5 (10)	CM992895
del(<i>GJB6</i> -D13S1830)	Putative abnormal regulation	0.3 (6)	CG024899
c.71G>A	p.Trp24*	0.3 (6)	CM970678
c.380G>A	p.Arg127His	0.2 (5)	CM980930
c.95G>A	p.Arg32His	0.2 (4)	CM013721
c.427C>T	p.Arg143Trp	0.2 (4)	CM000018
c.119C>A	p.Ala40Glu	0.1 (2)	CM041349
c.139G>T	p.Glu47*	0.1 (2)	CM970680
c.266T>C	p.Leu89Pro	0.1 (2)	This report
c.334_335delAA	p.Lys112Glufs*2	0.1 (2)	CD982678
c.559_561delGAG	p.Glu187del	0.1 (2)	CD042866
c.598G>A	p.Gly200Arg	0.1 (2)	CM0910079
c.632_633delGT	p.Cys211Leufs*5	0.1 (2)	CD982679
c.31_68del38	p.Gly11Leufs*24	<0.1 (1)	CG973465
c.94C>G	p.Arg32Gly	<0.1 (1)	This report
c.129delG	p.Trp44Glyfs*38	<0.1 (1)	Bliznetz et al. ^b
c.205T>C	p.Phe69Leu	<0.1 (1)	This report
c.245T>A	p.Ile82Asn	<0.1 (1)	This report
c.246C>G	p.Ile82Met	<0.1 (1)	CM021271
c.257C>G	p.Thr86Arg	<0.1 (1)	CM031189
c.385G>A	p.Glu129Lys	<0.1 (1)	CM014194
c.399G>A	p.Trp133*	<0.1 (1)	This report^c
c.402G>A	p.Trp134*	<0.1 (1)	This report
c.419T>G	p.Ile140Ser	<0.1 (1)	CM053901
c.502_601del10	p.Lys168Profs*5	<0.1 (1)	This report
c.532G>A	p.Val178Met	<0.1 (1)	This report
c.550C>T	p.Arg184Trp	<0.1 (1)	CM000709
c.614T>C	p.Leu205Pro	<0.1 (1)	CM055278

^aA number following CD, CS, CM, CI or CG is the HGMD Professional 2016.2 accession number for the mutation.

^bPatient with the novel c.129delG allele is the same with that previously published.¹⁹

^cThe mutation c.399G>A has NCBI dbSNP accession number rs777225786, but for it, a population frequency and clinical significance are unknown and observation in hearing loss individuals is not described. The names of the novel mutations are highlight in bold type.

substitutions and one pathogenic frame shifting deletion according to the ACMG guidelines for classifying pathogenic variants²⁷ (Table 2).

The novel *GJB2* mutations have not been detected in 1465 individuals who carry one or no pathogenic/likely pathogenic *GJB2* variant and have not been registered as at January 2017 in the databases of 1000 Genomes Project (<http://browser.1000genomes.org>, <http://www.ensembl.org>), ExAC (<http://exac.broadinstitute.org>), NCBI dbSNP and HGMD Professional 2016.2. An exception is mutation c.399G>A having NCBI dbSNP accession number rs777225786, of which population frequency and clinical significance are unknown and observation in hearing loss individuals is not described. The use of the prediction programs has shown that all novel mutations are 'disease causing' by software Mutation testing (<http://www.mutationtaster.org>) and all five missense substitutions are 'deleterious/damaging' by PROVEAN, SIFT and PolyPhen-2 (<http://provean.jcvi.org>, <http://genetics.bwh.harvard.edu/pph2>). In this work, Sanger sequencing of the PCR product, which covers the entire *GJB2* coding region, was performed using one primer pair. Consequently, when a nucleotide substitution in combination with a deletion (or an insertion) in the coding region was observed in a compound heterozygous genotype, both of these variants were presented together in a single forward sequence chromatogram and in a single reverse sequence chromatogram. Therefore, analysis of both sequence chromatograms, along with testing parental samples, was performed to determine whether the variant occurs in *cis* or in *trans*. Additional information about novel mutations, including family history and DNA sequence chromatograms, is presented in Supplementary Figure S1.

Frequency of del(*GJB2*-D13S175) among the *DFNB1* alleles in Russia

Previously, we documented the novel 101 kb deletion, del(*GJB2*-D13S175), in three unrelated Russian patients with sensorineural hearing loss and identified its breakpoints (NC_000013.10: g.20 757 021_20 858 394del). This deletion encompasses the entire *GJB2* and *GJB6* genes, and so it has been observed to cause a false homozygosity of the *GJB2* c.35delG mutation in one compound heterozygous patient and the lack of the *GJB2* gene sequence in two patients homozygous for the large deletion.¹⁰ Thus, an apparent homozygosity of the *GJB2* mutation can conceal a hemizyosity, if samples are not available from the patient's parents, and segregation analysis cannot be performed. Since most of our hearing loss patients have been investigated without material from parents, we proposed that among patients homozygous for the *GJB2* mutation, there are carriers of del(*GJB2*-D13S175).

Four (0.6%) carriers of del(*GJB2*-D13S175) have been discovered from the sample analysis of 691 patients, in whom previous testing at our laboratory detected 'homozygous' *GJB2* pathogenic variant but their parents' DNA were not tested. Among them, two patients were *GJB2* heterozygous for the c.35delG variant and the remaining two patients carried one of the c.290dupA or c.-23+1G>A variants (Table 3). Later, parental DNA has been tested to confirm *trans* configuration for the del(*GJB2*-D13S175) and c.290dupA alleles, while DNA from three patient's parents was not available. One more patient homozygous for del(*GJB2*-D13S175) was revealed using deletion-specific breakpoint primers, after PCR amplification with *GJB2*- and D13S175-specific primers was observed to be absent, in addition to two homozygous and one compound heterozygous for the deletion patients described previously.¹⁰ Thus, a total of 11 deletion-carrying chromosomes were detected in 8 unrelated patients (Table 3) that makes 0.5% of all *DFNB1* alleles in a study cohort of 1104 patients, carrying biallelic mutations in *GJB2/DFNB1*. In Table 1,

Table 2 Characteristics of the novel variants in the *GJB2* gene

Variant	Protein	Affected protein	Pathogenicity ^a		Genotype, number of family members with genotype	Severity of hearing loss	Ethnicity
	effect	domain	Pathogenicity ^a	Pathogenic criteria ^a			
c.94C>G	Missense	TM1	Likely pathogenic	PM2,3,5+PP1,2,3	c.[35delG]+[94C>G], 2	Moderately severe, severe	Tatar
c.205T>C	Missense	EC1	Likely pathogenic	PM2,3+PP2,3	c.[35delG]+[205T>C], 1	Severe	Unknown
c.245T>A	Missense	TM2	Likely pathogenic	PM2,3,5+PP2,3	c.[35delG]+[245T>A], 1	Severe	Tatar/Russian
c.266T>C	Missense	TM2	Likely pathogenic	PM2,3+PP1,2,3	c.[35delG]+[266T>C], 2	Severe	Russian
c.399G>A	Nonsense	TM3	Pathogenic	PVS1+PS1+PM2+PP3	c.[35delG]+[399G>A], 1	Severe to profound Unknown	Russian
c.402G>A	Nonsense	TM3	Pathogenic	PVS1+PS1+PM2,3,5+PP3	c.[35delG]+[402G>A], 1	Severe	Tajik
c.502_601del10	Frameshift	EC2	Pathogenic	PVS1+PM2,3+PP3	c.[35delG]+[502_601del10], 1	Severe	Russian
c.532G>A	Missense	EC2	Likely pathogenic	PM2,3,5+PP2,3	c.[35delG]+[457G>A,532G>A] ^b , 1	Severe to profound	Russian

^aPathogenicity was evaluated according to the ACMG guidelines for classifying pathogenic variants.²⁷ Each pathogenic criterion is weighted as very strong (PVS1), strong (PS1–4), moderate (PM1–6) or supporting (PP1–5).²⁷
^bc.457G>A is rare neutral polymorphism (rs111033186).

Table 3 Characteristics of patients with the del(*GJB2*-D13S175) mutation

Patient	Pathological genotype	Ethnicity (birth place)	Procedure and age at first hearing testing	Severity of hearing loss	Age at cochlear implantation	Outcome
A	del(<i>GJB2</i> -D13S175)/c.35delG	Ingush (Ingushetia)	CPA, 4 y	Moderate	Hearing aids were used	Good
B	del(<i>GJB2</i> -D13S175)/c. –23+1G>A	Ingush (Ingushetia)	OAE at birth, ABR, 12 m	Profound	31 months	Good
C	del(<i>GJB2</i> -D13S175)/c.35delG	Ingush (Ingushetia)	OAE at birth, ABR, 7 m	Profound	12 months	Good
D	del(<i>GJB2</i> -D13S175)/c.290dupA	Ingush (Ingushetia)	OAE at birth, ABR, 14 m	Profound	21 months	Good
E	del(<i>GJB2</i> -D13S175)/del(<i>GJB2</i> -D13S175)	Ingush (Ingushetia)	OAE at birth, ABR, 16 m	Profound	27 months	Good
F	del(<i>GJB2</i> -D13S175)/del(<i>GJB2</i> -D13S175)	Ingush (Ingushetia)	OAE at birth, ABR, 16 m	Profound	36 months	Good
G	del(<i>GJB2</i> -D13S175)/c.35delG	Ingush/Russian (Moscow)	OAE at birth, ABR, 5 m	Profound	11 months	Good
H	del(<i>GJB2</i> -D13S175)/del(<i>GJB2</i> -D13S175)	Ingush (Chechnya)	OAE at birth	Unknown	Unknown	Unknown

Abbreviations: ABR, auditory brainstem response audiometry; CPA, conditioned play audiometry; OAE, evoked otoacoustic emissions. By 'good outcome', we mean improvements in auditory skills as well as in the development of speech production.

Table 4 Number (%; 95% CI) of heterozygous carriers of del(*GJB2*-D13S175) or known *GJB2* mutations in apparently healthy populations

Mutation	Ingushes from Ingushetia, n = 151	Chechens from Ingushetia, n = 90	Chechens from Chechnya, n = 147
	del(<i>GJB2</i> -D13S175)	1 (0.7; 0.02–3.6)	1 (1.1; 0.03–6.0)
c.35delG	3 (2; 0.4–5.7)	0 (0; 0–4.0)	1 (0.7; 0.02–3.7)
c.358_360delGAG	0 (0; 0–2.4)	1 (1.1; 0.03–6.0)	2 (1.4; 0.2–4.8)
c.313_326del14	0 (0; 0–2.4)	0 (0; 0–4.0)	0 (0; 0–2.5)
c.235delC	0 (0; 0–2.4)	0 (0; 0–4.0)	0 (0; 0–2.5)

Abbreviation: CI, confidence interval.

a del(*GJB2*-D13S175) position from a list of all DFNB1 mutations in the study cohort is present.

Clinical follow-up of the del(*GJB2*-D13S175)-carrying patients

All patients with the del(*GJB2*-D13S175) deletion were diagnosed as having bilateral sensorineural hearing loss (Table 3). Hearing impairment in patient A was identified first at the age of 4 years using conditioned play audiometry. Pure-tone audiometry between 4 and 10 years of age demonstrated bilateral moderate sensorineural hearing loss in this patient (Supplementary Figure S2(1)). Hearing loss in

patients B, C, D, E, F, G and H was identified during newborn hearing screening performed using transient-evoked otoacoustic emission and distortion-product otoacoustic emission, evoked otoacoustic emission from the patients was absent (Supplementary Figures S2(2), S2(3)). Additionally, patients B, C, D, E, F and G audiological examination included auditory brainstem response audiometry, which revealed bilateral profound deafness in these patients since auditory brainstem response audiometries were absent at 70, 90 and 100 dBnHL (Supplementary Figure S2(4)). Middle ear endoscopy, tympanometry and X-ray computed tomography of the temporal bone did not detect any middle or inner ear malformation in patients A, B, C, D, E, F and G. Audiological testing data from patient H were not available.

Patients A, B, C, D, E, F and G underwent medical genetic counseling and pediatrician, neurologists, ophthalmologist and dermatologist examination. Only data from medical genetic evaluation of patient H were available. Patient A was diagnosed with psoriasis and has a twin brother with the same severity of hearing loss and satisfying outcomes were derived from the use of hearing aids. Astigmatism and a quite dry skin was revealed in patient F.

Evolutionary origins and heterozygous carriage of del(*GJB2*-D13S175)

Seven patients with del(*GJB2*-D13S175) and their parents were the Ingushes. The patient G was of mixed ancestry (Table 3). He has inherited the deletion from his father who had heterozygous

Table 5 Haplotype analysis

Marker	D13S1316	del(<i>GJB2</i> -D13S175)	D13S175	D13S1275	D13S232
GRCh37, Mb	20.68	20.76–20.86	20.85	22.95	23.80
Marshfield, cM	0		6.03	6.99	6.99
Genethon, cM	0		7.4	8.8	No data
<i>Chromosome</i>					
Patient E	15	del	del	22	11
Patient H	15	del	del	22	11
Patient E	2	del	del	22	11
Patient D	15	del	del	22	13
Patient A	15/14	del	del	22/24	11/19
Patient F	15	del	del	23	11/13
Patient F	15	del	del	25	13/11
Patient H	15	del	del	23	19
Patient G	15	del	del	20	19
Patient B	15/14	del	del	23/26	13
Patient C	15/14	del	del	21/19	13/20
The healthy Ingush	15/18	del	del	25/26	13/14
Allele associated with del(<i>GJB2</i> -D13S175)	15	del	del	22	11
P_D	0.875 (0.917)			0.455 (0.417)	0.455 (0.417)
P_N	1/6 = 0.167 ^a (1/10 = 0.1) ^a			23/180 = 0.128	23/160 = 0.144
P -value ^b	0.0353 (0.0007)			0.015 (0.0202)	0.0290 (0.0390)
δ ; 90% CI	0.850; 0.612–1.00 (0.908; 0.761–1.00)			0.375; 0.042–0.708 (0.331; 0.062–0.601)	0.363; 0.024–0.702 (0.319; 0.044–0.594)
g^{Marsh} ; 90% CI	No linkage map data			102; 36–328 (114; 53–289)	105; 37–386 (118; 54–325)
g^{Gen} ; 90% CI	No linkage map data			70; 25–224 (78; 36–197)	No linkage map data

Abbreviation: CI, confidence interval.

^aThe frequency obtained using the normal chromosomes from patients and their parents. δ , the degree of linkage disequilibrium by Bengtsson and Thomson,²⁴ $\delta = (P_D - P_N)/(1 - P_N)$, where P_D is the frequency of associated allele on del(*GJB2*-D13S175) carrying chromosomes and P_N is the frequency of the same allele on chromosomes without del(*GJB2*-D13S175). g , the generation number by Risch et al.,²² obtained by use of θ values between D13S175 and D13S1275 or D13S232 from Marshfield map, g^{Marsh} , and from Genethon map, g^{Gen} .

^b P -value for Yates corrected χ^2 .

At the top, the position of del(*GJB2*-D13S175) and four microsatellite markers studied. In the middle, haplotypes of del(*GJB2*-D13S175) carrying chromosomes. For patients A, B, C, F and the healthy Ingush, we were unable to phase mutation-carrying chromosomes into haplotypes by using other family members, therefore, two alleles are separated by slash. Below, the age of del(*GJB2*-D13S175) in the Ingushes is estimated. Within brackets, the calculations taking into account chromosomes of patients A, B, C, F and the healthy Ingush are present.

del(*GJB2*-D13S175) allele and whose father, in turn, was the Ingush. The Ingushes are a Caucasian native ethnic group of the North Caucasus, mostly inhabiting Russian Republic of Ingushetia. According to the 2010 Russian Census, Ingushes make up 94.1% of republic's population and 0.3% of Russia's population. The second ethnic group of Ingushetia includes Chechens (4.6%), genetically close to Ingushes. Chechens make up 95.3% of population of the Chechen Republic, geographically neighboring with Ingushetia.

We proposed a high frequency of del(*GJB2*-D13S175) in Ingush and Chechen populations, and investigated the population carrier frequency of del(*GJB2*-D13S175) and common *GJB2* mutations (c.35delG, c.313_326del14, c.235delC and c.358_360delGAG) among 151 Ingushes from Ingushetia, 90 native Chechens from Ingushetia and 147 Chechens from Chechnya, who had a normal hearing. The results of this investigation are presented in Table 4. The del(*GJB2*-D13S175) mutation was found in two individuals from Ingushetia while it was not observed among Chechens from Chechnya. It is the other way around, the c.358_360delGAG deletion was revealed in individuals from Chechnya only. The c.35delG mutation was found both in Ingushetia and Chechnya.

Age of del(*GJB2*-D13S175) estimation

Screening 3 STRs linked to *GJB2* in eight patients carrying del(*GJB2*-D13S175) and 2 of these STRs in 90 healthy Ingushes, we identified a common founder haplotype for the del(*GJB2*-D13S175)

allele and its decay derivatives (Table 5). The probable 'founder haplotype' is '15-22-11' at D13S1316-D13S1275-D13S232. To obtain an estimate of the age of del(*GJB2*-D13S175), we used calculations for allele '22' at D13S1275 and allele '11' at D13S232, which in a statistically significant linkage disequilibrium (LD) with del(*GJB2*-D13S175). The average spreading time (age) of del(*GJB2*-D13S175) was estimated to be 100 (90% confidence interval: 40–300) generations, when calculations from Table 5 were generalized. Applying an average generation time of 30 years,^{28–30} we estimated del(*GJB2*-D13S175) to be ~3000 years old (90% confidence interval: 1000–9000).

DISCUSSION

Among the *GJB2* variants reported previously, 252 missense substitutions, 54 small deletions, 26 nonsense substitutions, 18 small insertions, 7 small indels, 6 splice site substitutions, 5 large deletions, 3 substitutions in a regulatory sequence and 2 large insertions have been included in HGMD Professional 2016.2 as 'disease-causing' or 'probable pathological' mutations. Total of 100 and 39 null variants (nonsense, frameshift, canonical ± 1 or 2 splice sites) have been registered as at January 2017 in HGMD and dbSNP, consequently. Among them, 27 variants were contained in dbSNP and in HGMD simultaneously, and were labeled 'pathogenic' or 'likely pathogenic' in ClinVar database. Remaining 12 null variants from dbSNP were absent in HGMD or ClinVar, but were found in the collection of ExAC.

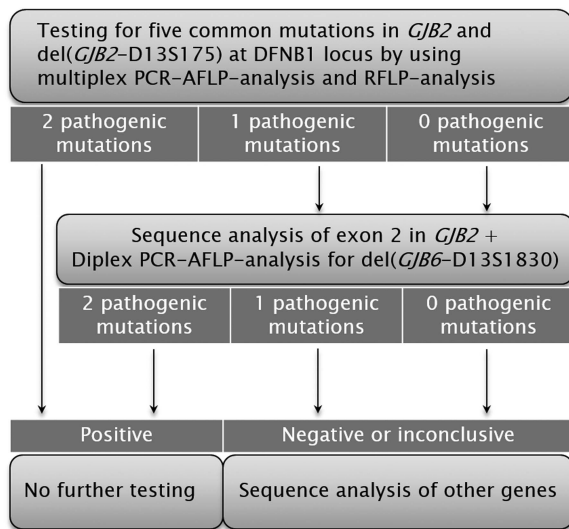


Figure 2 The cost-optimized DFNB1 testing strategy. The common mutation test lay in two reactions, a multiplex PCR with primer pairs for six deletions (c.35delG, c.313_326del14, c.235delC, c.167delT, c.358_360delGAG and *GJB2*-D13S175) and RFLP analysis of c.-23+1G>A, followed by gel-based fragment analysis to differentiate wild-type and mutation alleles.

Thus, currently there is not any benign (or likely benign) null variant of *GJB2*. Among 252 missense mutations registered in HGMD, 158 substitutions have not been presented in dbSNP or other databases. At the same time, total of 161 *GJB2* missense substitutions have been registered in dbSNP, which included 58 pathogenic (or likely pathogenic) and 11 benign (or likely benign) substitutions according to ClinVar. These data prove that missense variation in *GJB2*, as well as null variants, is a common cause of disease and there is little benign variation observed in the gene. Therefore, a novel missense variant in *GJB2* can be considered supporting evidence for pathogenicity, and a novel null variant can be weighted as a very strong pathogenic criterion, according to the ACMG guidelines for classifying pathogenic variants.²⁷ In this research, three unpublished pathogenic variants, namely, c.399G>A (p.Trp133*), c.402G>A (p.Trp134*), c.502_601del10 (p.Lys168Profs*5), and five novel likely pathogenic variants, c.94C>G (p.Arg32Gly), c.205T>C (p.Phe69Leu), c.245T>A (p.Ile82Asn), c.266T>C (p.Leu89Pro), c.532G>A (p.Val178Met), in the *GJB2* gene were detected in Russian patients with moderately severe or severe hearing loss. Each of these variants were observed in a single family, an exception is c.266T>C, which was found in two unrelated families.

The most remarkable updating of the DFNB1 mutation spectrum is the novel del(*GJB2*-D13S175) deletion, recently described by us. A deletion del(*GJB2*-D13S175) as it was shown in this study is the most frequent among other large deletions at DFNB1 locus in Russia and is predominantly associated with profound deafness. The frequency of del(*GJB2*-D13S175) in our patients with hearing loss is twice higher than frequency of the well-known del(*GJB6*-D13S1830) deletion. So del(*GJB2*-D13S175) detection is important, as well as del(*GJB6*-D13S1830) detection, and has been added to our testing strategy for DFNB1 hearing loss as outlined in Figure 2. The del(*GJB2*-D13S175) identification used to detect a heterozygous carriage in normal hearing individuals is especially necessary since del(*GJB2*-D13S175) is missed by standard methods of *GJB2* analysis, but it can be defined by using AFLP testing with deletion-specific breakpoint primers or copy-number analysis.

The del(*GJB2*-D13S175) mutation was originated from common founder in Ingush ethnic population. Among 241 normal hearing indigenous persons from Ingushetia, two heterozygous carriers for the deletion were observed (0.83% with 95% confidence interval 0.10%–2.97%), one can assume that there is unusually high carrier frequency of del(*GJB2*-D13S175) in Ingushetia. The similar frequency value of the c.35delG mutation was observed among normal hearing Ingushes. It is interesting that our cohort included a few patients with hearing loss from Ingushetia in addition to the patients with del(*GJB2*-D13S175). Among them, one out of two Ingush patients was homozygous for the c.35delG *GJB2* variant, while the remaining one Ingush patient has no mutation in *GJB2*. Therefore we assume that among Ingushes, a frequency of the del(*GJB2*-D13S175) allele is close to the c.35delG frequency, both among normal hearing and deaf individuals.

The *GJB2*-D13S175 deletion was found in the population samples both among Ingushes and Chechens from Ingushetia, while this deletion was not observed among Chechens from Chechnya. The other c.358_360delGAG mutation tested here was observed among Chechens, both from Chechnya and Ingushetia, but was not revealed among Ingushes. Probably, spectrum and frequency of *GJB2* mutations in Ingushes differ from that in Chechens from Chechnya, but Chechens from Ingushetia have mixed genetic material for this locus of both populations. This assumption agrees with findings, reported by Balanovsky *et al.*,³¹ which showed a high degree of genetic subdivision in the Caucasus populations resulting from genetic drift, probably due to isolation in the extremely mountainous landscape.

Study by Balanovsky *et al.*³¹ presents the most extensive survey of Y chromosomal variation in the Caucasus. Authors concluded that the Caucasus male lineages originated from a subset of the Near Eastern gene pool due to an Upper Paleolithic (or Neolithic) migration, followed by high levels of isolation, differentiation and genetic drift *in situ*. This process would result in the loss of some haplogroups and the increased frequency of others. Despite the fact that Ingushes and Chechens are nearest neighbors and had a shared ancestry for a long period, their languages and gene pools have been split ~2000 years ago, to what a linguistic history and age of a few specific haplotype clusters for Y chromosomal STRs in these populations testify.³¹ Our results demonstrate that the del(*GJB2*-D13S175) allele was spread in Ingushes after they separated from Chechens. In accordance with our average age estimates, the beginning of the del(*GJB2*-D13S175) spread among Ingushes occurred 1000 years before the linguistic division of Ingushes took place or the Ingush-specific Y chromosome haplotype clusters were originated.

It should be noted that at present over 25% of Ingushes live outside Russian Federation, mainly in Turkey, Syria, Jordan, Lebanon and Kazakhstan.³² The first mass migration of Ingushes and other North Caucasian ethnic groups to Ottoman Empire was happening during the second half of the nineteenth century after Caucasian War. During World War II, the whole of the Ingush and Chechen populations of the North Caucasus was temporarily deported to Kazakhstan and Central Asia, where they have partly remained and live to this day.³³ Thus, the Ingush deletion can even quite possibly be found in countries mentioned above. It is possible also that the deletion has been already present among Ingush ancestors, for example, among Neolithic/Bronze age migrants from the Near East, before the Ingush people were isolated. Therefore, to establish whether the deletion is unique for the Ingushes, additional researches are necessary. It is important both for clinical genetics and for the Caucasus history study.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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