

Utilizing Ethnic-Specific Differences in Minor Allele Frequency to Recategorize Reported Pathogenic Deafness Variants

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Ethnic-specific differences in minor allele frequency impact variant categorization for genetic screening of nonsyndromic hearing loss (NSHL) and other genetic disorders. We sought to evaluate all previously reported pathogenic NSHL variants in the context of a large number of controls from ethnically distinct populations sequenced with orthogonal massively parallel sequencing methods. We used HGMD, ClinVar, and dbSNP to generate a comprehensive list of reported pathogenic NSHL variants and re-evaluated these variants in the context of 8,595 individuals from 12 populations and 6 ethnically distinct major human evolutionary phylogenetic groups from three sources (Exome Variant Server, 1000 Genomes project, and a control set of individuals created for this study, the OtoDB). Of the 2,197 reported pathogenic deafness variants, 325 (14.8%) were present in at least one of the 8,595 controls, indicating a minor allele frequency (MAF) >0.00006. MAFs ranged as high as 0.72, a level incompatible with pathogenicity for a fully penetrant disease like NSHL. Based on these data, we established MAF thresholds of 0.005 for autosomal-recessive variants (excluding specific variants in *GJB2*) and 0.0005 for autosomal-dominant variants. Using these thresholds, we recategorized 93 (4.2%) of reported pathogenic variants as benign. Our data show that evaluation of reported pathogenic deafness variants using variant MAFs from multiple distinct ethnicities and sequenced by orthogonal methods provides a powerful filter for determining pathogenicity. The proposed MAF thresholds will facilitate clinical interpretation of variants identified in genetic testing for NSHL. All data are publicly available to facilitate interpretation of genetic variants causing deafness.

The advent of massively parallel sequencing has shifted the bottleneck in human genetics from data acquisition to variant interpretation. Accurate evaluation of genetic variants for pathogenicity is crucial to advance our understanding of disease processes and is a requirement for clinical diagnostics. Whole-exome sequencing and targeted gene panels based on targeted genomic enrichment and massively parallel sequencing are becoming commonplace and for some Mendelian diseases, including breast and ovarian cancer, degenerative eye disease, and hearing loss, they have become the ideal test and are now used routinely for clinical diagnostic testing.^{1–3} These tests regularly produce hundreds or thousands of variants that

require categorization and interpretation to assess their likelihood of causing disease. Correct interpretation is crucial when test results are used to direct clinical care.

Hearing loss (HL) is the most common sensory deficit in humans, affecting 1 in 500 children⁴ and 360 million people worldwide (World Health Organization Deafness Estimate Online Report). The majority of HL is genetic and nonsyndromic (NSHL, not associated with other clinical phenotypes). Genetic diagnosis for NSHL is particularly challenging given limited phenotypic variability and extreme genetic heterogeneity; increased use of massively parallel sequencing resulting in thousands of variants identified per individual has highlighted these challenges.

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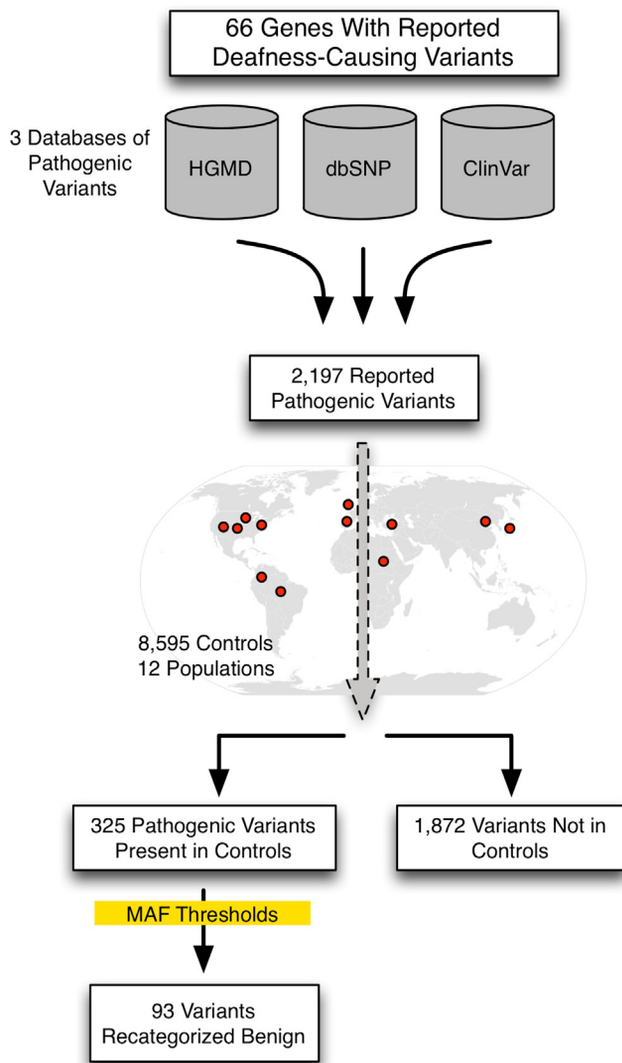


Figure 1. Study Overview

To date more than 70 genes and more than 2,000 causal variants have been implicated in NSHL. As with other genetic diseases, the vast majority of reported pathogenic NSHL variants were so designated based on five primary criteria: (1) cosegregation with the phenotype ideally with linkage analysis, (2) absence in 200 or more control chromosomes from the same ethnicity, (3) conservation of the affected nucleotide or amino acid through evolution, (4) predicted functional effect of the mutation, and (5) corroborative *in vitro* or *in vivo* functional data.

Absent, however, has been a sixth criterion—one that would require assessment of the variant in the context of a large number (thousands) of controls from varying ethnicities. This omission is noteworthy because the majority of human genetic variation is accounted for by ethnicity-specific differences.^{5,6} Our group and others have hypothesized that evaluation of variants in this context can reduce or eliminate false positives. Massively parallel sequencing techniques have for the first time delivered

the ability to sequence a heretofore-unobtainable number of control individuals from different ethnicities with data freely available to the scientific community.

In the first application of a large amount of control data to evaluate a human genetic disease, Norton et al. proposed a disease-specific iterative approach: use a large number of controls to determine the minor allele frequency (MAF) of known pathogenic variants and thereby determine a MAF cut-off that can be used for evaluation of future variants.⁷ The authors used data from the Exome Variant Server (EVS), which includes two ethnically distinct populations, to resolve false positives. Another recent study disqualified ten reported X-linked disability genes using similar methods and data from the EVS.⁸ Most recently, a gene with variants reported to cause NSHL, *MYO1A* (MIM 601478), has been disqualified through the use of EVS and lack of cosegregation of previously reported pathogenic variants in this gene.⁹

Incorrect classification of genetic variants as pathogenic is not limited to only one or a few genetic diseases—it is a systemic issue in human genetics. Cassa et al. evaluated all of the mutations present in the Human Gene Mutation Database (HGMD), the most comprehensive repository of pathogenic human mutations, in the context of control data from the 1000 Genomes Project.¹⁰ The authors found that 3.5% of variants present in HGMD have a MAF > 0.05, which is implausible for a highly penetrant Mendelian disease.

Our goal in this study was to evaluate all reported pathogenic deafness variants in the context of a large number of controls from multiple ethnicities in order to recategorize any clearly benign variants based on MAF in controls and establish MAF thresholds to aid interpretation of variants discovered in the future (Figure 1).

We sought to evaluate pathogenic variants present in 66 genes with variants reported to cause NSHL and NSHL-mimic syndromes (genetic causes of syndromic hearing loss that mimic NSHL at presentation such as Usher syndrome, the most common cause of deaf-blindness; see Table S1 [available online] for a complete gene list). Reported pathogenic deafness variants were obtained from three sources: HGMD (Professional v.2013.4), NCBI ClinVar, and dbSNP v.138 (the latter publically available databases were accessed in February 2014). When necessary, variants were converted to HGVS nomenclature using Mutalyzer to generate genomic coordinates.¹¹ The final comprehensive list of reported pathogenic deafness variants included all variants categorized as fully penetrant pathogenic or probable-pathogenic in any of the databases (DM or DM? in the case of HGMD).

In total, we obtained 2,197 variants designated as pathogenic or probable-pathogenic by any of three databases. As shown in Figure S1, the majority of these variants were identified in HGMD alone (1,572; 71.6%), seven were unique to ClinVar, and none were unique to dbSNP. The three databases shared 454 (20.7%) of the reported pathogenic variants.

Table 1. Previously Reported Pathogenic Deafness-Causing Variants

Variant Type	All	Autosomal Recessive	Autosomal Dominant	X-Linked
Nonsynonymous	1,165	972	173	20
Stop gain	350	336	7	7
Stop loss	3	2	1	–
Synonymous	31	28	3	–
Splice site	238	232	6	–
Frameshift deletion	330	311	9	10
Frameshift insertion	25	24	–	1
Nonframeshift deletion	46	35	9	2
Nonframeshift insertion	9	9	–	–
Total	2,197	1,949	208	40

There was disagreement between databases in the categorization of 96 variants. All differences were between HGMD and the other two databases, ClinVar and dbSNP, which agreed in all cases. HGMD categorized 87 variants pathogenic while the other two databases categorized the same variants as probable-pathogenic (41), probable-nonpathogenic (20), or nonpathogenic (26). HGMD categorized 9 variants as probable-pathogenic while the other two databases categorized these variants as pathogenic.

Of the 2,197 reported pathogenic deafness variants, 1,165 were nonsynonymous, 350 were stop gain, 330 were frameshift deletions, 238 were splice site mutations, 46 were non-frameshift deletions, 31 were synonymous, 25 were frameshift insertions, 9 were nonframeshift insertions, and 3 were stop loss (Table 1). When divided by the mode of inheritance, there were 1,949 autosomal-recessive, 208 autosomal-dominant, and 40 X-linked recessive pathogenic variants.

In total, we ascertained 8,595 control individuals from 12 populations and 6 major phylogenetic groups of human evolutionary history using three orthogonal massively parallel sequencing methods. The populations ranged in size from 100 to 4,300 individuals (Table 2). Control data were obtained from three sources: (1) the NHLBI Go Exome Sequencing Project (ESP), made available on the Exome Variant Server (EVS) version ESP6500SI-V2 accessed 2/2014, (2) the 1000 Genomes Project (1000G) release 1000GApr2012, and (3) the OtoDB, the OtoSCOPE Control Database, data from 1,000 normal hearing controls from six populations, which we generated for this project by pooled targeted genomic enrichment (methods described in detail below). A summary of the controls used in this study is provided in Table 2.

The EVS database includes 6,503 individuals from two populations—4,300 European Americans (EA) and 2,203 African Americans (AA)—and was established to study individuals on the extremes of cardiovascular phenotypes as well as control individuals. Whole-exome sequencing

(WES) was performed on these samples with a target average coverage $>80\times$. Coverage statistics for the genes targeted in this study were calculated using BED files provided by the ESP.

The 1000G database includes 1,092 individuals from 4 superpopulations (large populations comprised of several subpopulations, generally from the same geographic region and shown to be closely related in evolutionary history): 246 samples from the African superpopulation (AFR), 181 samples from the American superpopulation (AMR), 286 individuals from the Asian superpopulation (ASN), and 379 individuals from the European superpopulation (EUR). Variant calls were obtained from a combination of WES and low-coverage whole-genome sequencing (WGS+WES). Coverage statistics for genes targeted in this study were calculated from BAM files for ten individuals.

The OtoDB includes 1,000 normal-hearing individuals from six populations sequenced using pooled targeted genomic enrichment and massively parallel sequencing (PTE+MPS). The populations included (1) 200 Ashkenazi Jews living in New York (AJ), (2) 160 Colombians originating from different cities around the country including Bogotá and the central region, the Atlantic coast, and the north, west, and southwest regions of the country (CO), (3) 200 Japanese students attending Shinshu University in Matsumoto (JP), (4) 160 European Americans from Iowa, USA (US), (5) 180 Spanish from Almería and Granada (ES), and (6) 100 Turks, primarily from the Southern and Mediterranean Sea regions (TR). All samples were from individuals with no self-reported hearing loss, and in the case of Japanese samples, subjects had audiometric testing showing no evidence of hearing loss. Individuals from the EVS and 1000G databases, in contrast, were not evaluated for hearing loss and therefore are not true “controls” but rather representative cross-sections of their derivative populations. Human ethics and institutional review board (IRB) policies were followed at each institution where samples were collected and informed consent was obtained. All methods were approved by the IRB at the University of Iowa.

To sequence the 1,000 OtoDB controls, we used a pooled targeted genomic enrichment method that entailed generating 50 pools, each of which included 20 DNA samples (40 alleles per pool). We then performed targeted genomic enrichment and massively parallel sequencing as described using automation equipment.³ We captured the exonic regions of 66 genes with variants causing NSHL, targeting in total 1,344 regions and 521,647 bp of the genome (Table S1). Molecular barcodes were added to each pool of 20 DNA samples. Five pools were then multiplexed and sequenced in a single flow-cell of the Illumina HiSeq using 100 bp paired-end reads (100 samples per flow-cell). We sought to generate a sequencing depth-of-coverage sufficient for variant calling for each allele present in the pool for all targeted regions (10 \times per sample or 200 \times per pool).

We analyzed the pooled data sets using a local installation of the open-source Galaxy software¹² and the following

Table 2. Populations Used in This Study

Phylogenetic Group	Population	n Individuals	DB	
			Source	DB Type
African	AFR: African superpopulation	246	1000G	WGS+WES
	AA: African American	2,303	EVS	WES
American	AMR: American superpopulation	181	1000G	WGS+WES
	CO: Colombia	160	OtoDB	PTE+MPS
East Asian	ASN: Asian superpopulation	286	1000G	WGS+WES
	JP: Japan	200	OtoDB	PTE+MPS
European	EUR: European superpopulation	379	1000G	WGS+WES
	EA: European American	4,300	EVS	WES
	US: Europeans from USA	160	OtoDB	PTE+MPS
	ES: Spain	180	OtoDB	PTE+MPS
European, isolated	AJ: Ashkenazi Jewish	200	OtoDB	PTE+MPS
Mediterranean	TR: Turkey	100	OtoDB	PTE+MPS
Total		8,595		

Abbreviations are as follows: 1000G, 1000 Genomes Project; EVS, Exome Variant Server; OtoDB, OtoSCOPE Control Database; WGS, whole-genome sequencing; WES, whole-exome sequencing; PTE, pooled targeted enrichment; MPS, massively parallel sequencing.

steps: read mapping with Burrows-Wheeler Alignment (BWA),¹³ duplicate removal with Picard, local realignment with GATK,¹⁴ and variant calling with SAMtools.¹⁵ Coverage statistics were calculated with BEDTools.¹⁶ Variant frequency in each pool was assigned using vipR, a previously validated method that accurately calls variants in pooled data sets using skellam distributions.¹⁷ We ran vipR using default parameters except we enabled indel calling. Annotation of identified variants was performed with ANNOVAR.¹⁸

For the pooled OtoDB control set, the average pool depth of coverage was 7,846 for an average inferred per-sample depth of coverage of 392 \times . To validate pooled sequencing and the vipR analysis method, we used Sanger sequencing and genotyped 12 single-nucleotide changes in 100 persons comprising 5 pools including 7 variants showing the minimum frequency in the pool (1 allele in 40, frequency of 0.025). We found a sensitivity of 0.971 and specificity of 0.989 when using Sanger sequencing as the gold standard (Table S2).

As expected, coverage statistics for the 66 targeted NSHL genes varied by data source due to the different sequencing methods used (Table S3). Because coding variants comprise the majority of pathogenic mutations, we focused on exons and found that $\geq 1\times$ exonic coverage of these genes was 68.9%, 82.6%, and 99.8% with EVS, 1000 Genomes,

and OtoDB, respectively. When restricted to coding exons only, $\geq 1\times$ coverage was 88.7%, 93.8%, and 99.4% (Table S3). Average overall coding depth of coverage was 246, 90, and 475 for EVS, 1000 Genomes, and OtoDB, respectively. These data provided the source for annotation of the identified pathogenic variants.

Of the 2,197 pathogenic variants obtained from the three databases, we found 325 variants (14.8%) in controls. Fifty-seven of these variants (17.5%) were present in all three databases (Figure S2). By inheritance pattern, 296 and 29 pathogenic variants were associated with autosomal-recessive and autosomal-dominant NSHL, respectively. There were no reported X-linked NSHL variants in controls. In aggregate, of the 42 autosomal-recessive and 20 autosomal-dominant NSHL genes we screened, 24 recessive and 9 dominant genes contained pathogenic variants in controls (Table 3).

As shown in Table 3, the proportion of pathogenic variants in controls varied by gene and was highest at 86% for *MYO1A*. Variants in this gene were recently discounted as causing autosomal-dominant NSHL.⁹ Other genes with a high proportion of pathogenic variants found in controls include 50% for *DFNB31* (MIM 607084), *GJB3* (MIM 603324), *GRXCR1* (MIM 613283), *SLC26A5* (MIM 604943), and *DIAPH1* (MIM 602121). The minor allele frequencies (MAF) for these variants ranged from the limit of detection, which was 0.00006, to 0.72 for *GJB2* (MIM 121011; RefSeq accession number NM_004004; c.79G>A [p.Val27Ile]) in the Japanese population of OtoDB, a variant that is commonly considered nonpathogenic but is labeled as probable-nonpathogenic by ClinVar and dbSNP and probable-pathogenic by HGMD. We noted that several variants had MAF levels implausible for a fully penetrant Mendelian disease like deafness.

The remaining 1,872 variants (85.2%) were not identified in the 8,595 controls consistent with extreme rarity ($<1/17,190$ alleles or a MAF < 0.00006) and indicating an increased likelihood of pathogenicity. The distribution of pathogenic variants by MAF is shown in Figure 2. We evaluated variants using the maximum MAF (Max MAF), the highest MAF in any one of the 12 populations studied. Detailed data for each variant including MAF per population are provided online in full at the Deafness Variation Database (see Web Resources section).

Based on these data, we sought to identify appropriate MAF cut-offs for autosomal-recessive and autosomal-dominant NSHL. For severe-to-profound autosomal-recessive congenital NSHL, there are a wealth of data showing that the most common genetic cause in most world populations are mutations in *GJB2*, with specific variants like c.35delG and c.167delT having exceptionally high frequencies in the European and Jewish populations, respectively. The aggregate *GJB2* mutation carrier frequency in these populations is typically 1%–2%.

We used these data to create an empirically based cut-off for autosomal-recessive NSHL by first positing a MAF of 0.01 for any autosomal-recessive NSHL-causing variant in

Table 3. Previously Reported Pathogenic Deafness-Causing Variants Identified in All Individuals Studied by Gene

Gene	n Reported Path. Variants	% of Path. Variants Identified in Individuals (n)	MAF Range	% of Path. Variants Recategorized Benign (n)
Autosomal Recessive				
CDH23	166	14% (24)	0.0001–0.1417	2% (4)
CLDN14	6	33% (2)	0.0001–0.0002	–
CLRN1	24	29% (7)	0.0001–0.0075	4% (1)
COL11A2	1	0 (0)	–	–
DFNB31	14	50% (7)	0.0001–0.01	7% (1)
DFNB59	13	8% (1)	0.0001–0.0003	–
ESPN	6	0 (0)	–	–
ESRRB	8	13% (1)	0.0002	–
GIPC3	9	0 (0)	–	–
GJB2	277	15% (42)	0.0001–0.7225	2% (6)
GJB3	6	50% (3)	0.0003–0.0307	33% (2)
GJB6	9	11% (1)	0.0003–0.0005	–
GPR98	51	18% (9)	0.0001–0.04	8% (4)
GPSM2	2	0 (0)	–	–
GRHL2	2	0 (0)	–	–
GRXCR1	4	50% (2)	0.0001–0.06	–
HGF	1	0 (0)	–	–
ILDR1	12	0 (0)	–	–
LHFPL5	5	0 (0)	–	–
LOXHD1	7	43% (3)	0.0007–0.06	14% (1)
LRTOMT	8	0 (0)	–	–
MARVELD2	6	0 (0)	–	–
MYO15A	52	17% (9)	0.0001–0.014	8% (4)
MYO3A	4	0 (0)	–	–
MYO6	2	0 (0)	–	–
MYO7A	257	11% (27)	0.0001–0.14	2% (6)
OTOA	2	0 (0)	–	–
OTOF	110	20% (22)	0.0001–0.03	5% (5)
PCDH15	41	15% (6)	0.0001–0.0945	10% (4)
RDX	3	0 (0)	–	–
SLC26A4	339	16% (53)	0.0001–0.1428	4% (12)
SLC26A5	2	50% (1)	0.002–0.0156	50% (1)
STRC	15	13% (2)	0.0002–0.0004	7% (1)
TECTA	10	0 (0)	–	–
TMC1	36	25% (9)	0.0001–0.0031	–
TMIE	10	0 (0)	–	–
TMPRSS3	27	33% (9)	0.0001–0.0483	7% (2)
TPRN	5	0 (0)	–	–
TRIOBP	9	11% (1)	0.0001	–

(Continued on next page)

Table 3. Continued

Gene	n Reported Path. Variants	% of Path. Variants Identified in Individuals (n)	MAF Range	% of Path. Variants Recategorized Benign (n)
<i>USH1C</i>	24	29% (7)	0.0001–0.06	8% (2)
<i>USH1G</i>	9	0 (0)	–	–
<i>USH2A</i>	354	13% (47)	0.0001–0.1743	5% (17)
Autosomal Dominant				
<i>ACTG1</i>	11	0 (0)	–	–
<i>COCH</i>	16	6% (1)	0.0001	–
<i>COL11A2</i>	2	0 (0)	–	–
<i>CRYM</i>	2	0 (0)	–	–
<i>DFNA5</i>	2	0 (0)	–	–
<i>DIAPH1</i>	2	50% (1)	0.0035	50% (1)
<i>DSPP</i>	2	0 (0)	–	–
<i>GJB2</i>	30	3% (1)	0.01	3% (1)
<i>GJB3</i>	9	44% (4)	0.0001–0.02	44% (4)
<i>GJB6</i>	3	33% (1)	0.0002–0.0006	33% (1)
<i>KCNQ4</i>	19	0 (0)	–	–
<i>MYH14</i>	6	17% (1)	0.0007–0.01	17% (1)
<i>MYH9</i>	1	0 (0)	–	–
<i>MYO1A</i>	7	86% (6)	0.0002–0.1125	57% (4)
<i>MYO6</i>	6	0 (0)	–	–
<i>MYO7A</i>	7	0 (0)	–	–
<i>POU4F3</i>	7	0 (0)	–	–
<i>TECTA</i>	39	18% (7)	0.0001–0.0031	10% (4)
<i>TMC1</i>	3	0 (0)	–	–
<i>WFS1</i>	34	21% (7)	0.0001–0.0232	12% (4)

Abbreviations are as follows: path., pathogenic; MAF, minor allele frequency.

GJB2. Our data, for example, gave a maximum MAF of 0.0108 and 0.01 for *GJB2* (RefSeq NM_004004): c.35delG and c.167delT, respectively. Next, we assumed the existence of a single mutation in a second gene that accounts for roughly one-quarter of all remaining recessive NSHL, a very liberal assumption given the documented genetic heterogeneity. The MAF of this variant would be 0.005. We then hand-curated the 83 recessive NSHL variants with MAFs > 0.005 for plausibility of pathogenicity and recategorized 73 variants as benign (Figure 2 and Table S4).

To evaluate the predicted effects of nonsynonymous changes, we used six pathogenicity prediction algorithms derived from dbNSFP v.2.3¹⁹ and annotated with ANNOVAR: SIFT, Polyphen2_HVAR, LRT, MutationTaster, MutationAssessor, and FATHMM. We used these methods to evaluate variants individually and as part of a composite pathogenicity score. For the composite score (path_composite), a score of 1 point was assigned for each tool indicating pathogenicity or likely pathogenicity. Because pathogenicity predictions were not available for

every variant for each tool, the sum was divided by the total number of predictions available for that variant to create a ratio from 0 to 1. We also used the MetaLR score, a radial support-vector machine (SVM) method trained with a hand-curated list of causative variants and scored using several available pathogenicity tools. MetaLR is part of the ANNOVAR suite and initial studies have shown that its performance is superior to other pathogenicity prediction tools.

To test the plausibility of this MAF threshold for autosomal-recessive NSHL in an outbred population, we used these methods to evaluate the recessive variants we recategorized as benign and compared this score to the pathogenicity score for pathogenic variants (Figure S3 and Table S5). Autosomal-recessive NSHL variants recategorized as benign had a significantly lower pathogenicity score for seven of the eight methods used including the composite score and SVM score ($p < 0.05$). These data support the recategorization of these variants as benign and the MAF threshold we propose.

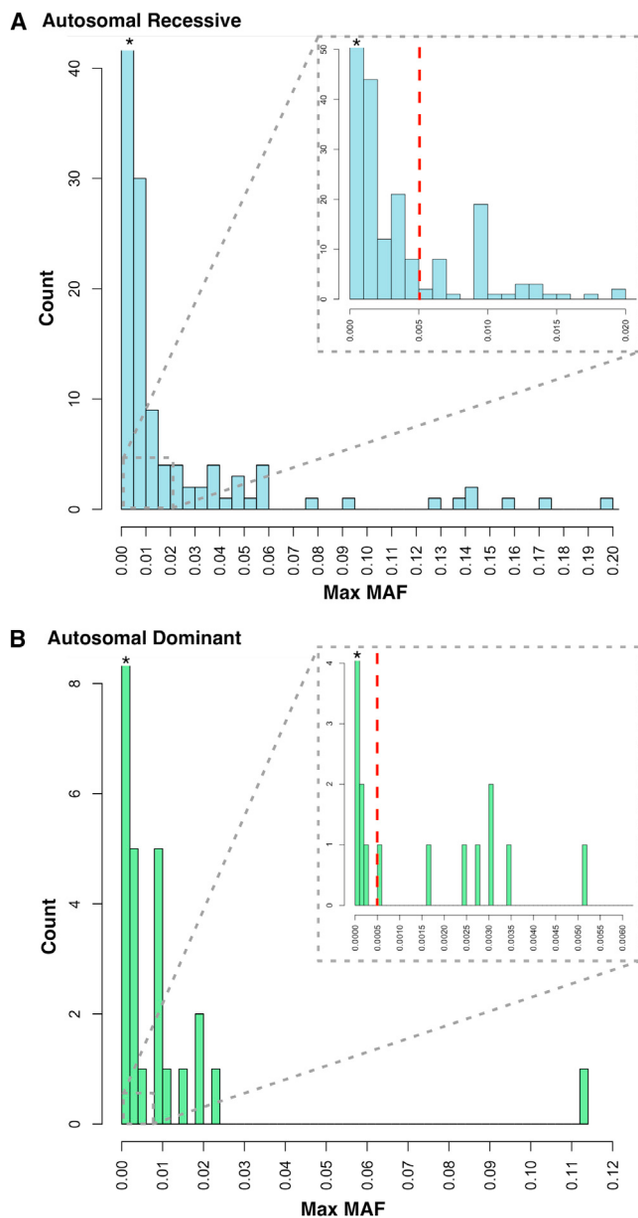


Figure 2. Distribution of Pathogenic Variants by Maximum Minor Allele Frequency in 12 Populations

Autosomal-recessive (A) and autosomal-dominant (B) variants. Dashed red line indicates MAF cut-offs for each inheritance mode. Asterisk (*) indicates that the count extends beyond the axis.

Of the 177 dominant pathogenic deafness variants, 77 were present in controls. To establish a threshold for autosomal-dominant NSHL, we used a hypothesis-based method because no single mutation comprises the majority of autosomal-dominant NSHL in any given population. We assumed Hardy-Weinberg equilibrium as well as the following generally accepted values: 1 in 500 prevalence of NSHL, 80% genetic, 70% nonsyndromic, 20% dominant, and 56 dominant loci. These assumptions predict that if there is only a single pathogenic variant in a given autosomal-dominant NSHL gene, its MAF can be no higher than 1/2,000 or 0.0005. We then used this threshold to

reclassify 20 variants as benign (Table S4). Pathogenicity prediction scores were lower for 7 of 8 methods, and significantly lower for 4 of 8 methods including the composite score and SVM score, again supporting this reclassification of the variants (Figure S3 and Table S5).

Of the 93 variants we reclassified as benign, HGMD contained calls for all but four (89) and classified 100% of these as pathogenic or probable-pathogenic (Table S4 and Figure S4). In comparison, ClinVar and dbSNP contained 41 of these variants but classified only 19 (46%) as pathogenic. Our data indicate that although the HGMD database contains more variants, it is not editable by users and therefore may contain more false positives.

The importance of filtering against a large and diverse control population is illustrated by our ability to reclassify variants based on the populations used. Every population used in this study, regardless of size, contained variants above the MAF thresholds and therefore contributed to reclassification of variants (Figure 3). The contribution of each population to reclassification of variants varied from 38 variants above MAF threshold for the EVS African American population to 10 variants above MAF threshold for the OtoDB Japanese population (Figure 3). In addition, the number of unique variants above MAF varied from 0 to 5, with all populations but one (OtoDB_JP) contributing at least one unique ethnicity-specific variant above the set MAF threshold (Figure 3). The size of the control population did not correlate with the number of variants above MAF threshold ($r^2 = 0.15$).

Of the 93 reclassified variants in this study, only 26% would have been found to be above the proposed thresholds when using a single population database—EVS European Americans, which contains 4,300 individuals, the highest number of any database used in this study. However, by including other individuals of European descent sequenced with the other two orthogonal platforms (1000G EUR and OtoDB ES and US), 44% of the variants rose above the cut-offs, reflecting differences in coverage metrics and platform-specific weaknesses. As we added individuals from other ethnicities, we further improved our ability to detect and therefore reclassify variants from 52% (after addition of Turkish and Ashkenazi Jewish OtoDB individuals) to 64% after addition of Asian individuals (1000G ASN and OtoDB JP) to 73% after addition of American individuals (1000G AMR and OtoDB CO) and finally to 100% after addition of African individuals (EVS AA, 1000G AFR). These data support the need to generate multiple ethnically discrete population databases on a variety of different platforms and indicate that it may be more important to have a large number of populations from disparate ethnicities sequenced with orthogonal platforms rather than a few populations of enormous size.

Interpretation of genetic variants is of primary concern for clinical diagnosis and human genetics research, and the MAF thresholds we propose—0.005 and 0.0005 MAF for autosomal-recessive and autosomal-dominant NSHL variants, respectively—provide thresholds for interpreting

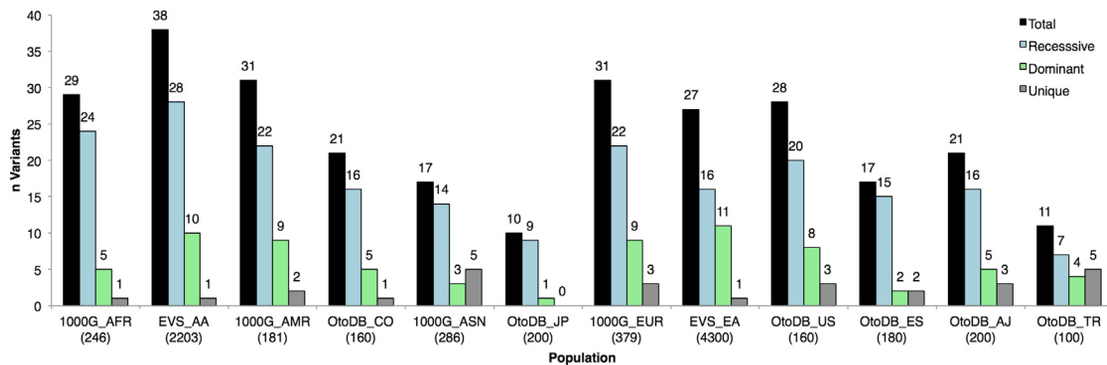


Figure 3. Contribution of Each Population to Recategorization of 93 Previously Reported Pathogenic Variants

The count for each population is the number of variants found in each population above MAF threshold. Values for total, recessive, and dominant variants are shown along with the number of variants above the MAF unique to that single population (Unique). Number of individuals per population are listed in parenthesis on the x axis, also see Table 1.

newly identified variants. Although these thresholds are conservative, in clinical testing each variant should be interpreted in the context of the phenotype and family history and with knowledge of the contribution of specific deafness-causing variants to disease in the particular population being studied.

Our study confirms the findings recently disqualifying variants in *MYO1A* as a cause of deafness,⁹ as four of the seven variants (57%) in *MYO1A* exceeded the MAF cut-off we propose for autosomal-dominant NSHL and two more fall immediately below this cut-off. *MYO1A* had more variants recategorized as benign than any other gene with variants reported to cause NSHL (Table 3). Our data also suggest that particular attention should be paid to two other genes with variants reported to cause NSHL that we believe are questionable, namely *GJB3*²⁰ and *SLC26A5*.²¹ These genes should be evaluated further to determine whether variants in these genes truly cause fully penetrant deafness.

This study has two important limitations. First, CNVs were not included, and as we recently showed, their contribution to NSHL is significant.²² A better understanding of the pathogenicity of CNVs is needed. However, a large number of controls must be analyzed using a reliable and sensitive method to detect CNVs before this question can be answered. Second, we did not ascertain all possible pathogenic NSHL-causing variants. Although we did attempt to collate a comprehensive list of reported pathogenic deafness variants, invariably some variants have been missed. For example, in the case of the Leiden-Open Variant Database (LOVD), we could not reliably retrieve variants in bulk from this database. This limitation underscores the importance of standardizing the recording of genetic data and making it freely accessible to improve clinical care and research.

Ideally, each variant identified by genetic testing should be evaluated in the context of the ethnic-specific prevalence of NSHL in the derivative population and also the variant-specific MAF in a large number of ethnically diverse populations. This study represents an important

step toward this goal but it is beyond the scope of this paper to provide prevalence data for NSHL in the different populations we studied. Although 1,872 variants (85.2%) were not identified in the populations we studied, supporting their role in human deafness, we cannot unequivocally ensure their pathogenicity as they could be present at high MAF in populations we have not queried.

In conclusion, our results validate the use of multiple platforms and multiple ethnicities in filtering and ascribing significance to genetic variants associated with NSHL. We recategorized 93 previously reported pathogenic deafness variants as benign. The MAF thresholds we propose will facilitate the interpretation of newly discovered variants in the context of NSHL. To further this goal, all data presented here are provided in the Deafness Variation Database (see Web Resources), a freely available and regularly updated database of all variants in known deafness genes.

Supplemental Data

Supplemental Data include four figures and five tables and can be found with this article online at <http://dx.doi.org/10.1016/j.ajhg.2014.09.001>.

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Web Resources

The URLs for data presented herein are as follows:

1000 Genomes, <http://browser.1000genomes.org>

ANNOVAR, <http://www.openbioinformatics.org/annovar/>
 ClinVar, <https://www.ncbi.nlm.nih.gov/clinvar/>
 dbSNP, <http://www.ncbi.nlm.nih.gov/projects/SNP/>
 Deafness Variation Database, <http://deafnessvariationdatabase.org>
 Galaxy, <https://usegalaxy.org/>
 Human Gene Mutation Database, <http://www.hgmd.org/>
 Mutalyzer, <https://mutalyzer.nl/index>
 NHLBI Exome Sequencing Project (ESP) Exome Variant Server,
<http://evs.gs.washington.edu/EVS/>
 Online Mendelian Inheritance in Man (OMIM), <http://www.omim.org/>
 RefSeq, <http://www.ncbi.nlm.nih.gov/RefSeq>
 World Health Organization Deafness Estimate, <http://www.who.int/pbd/deafness/estimates/en/>

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