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Targeted next generation sequencing for molecular diagnosis of Usher syndrome

María J Aparisi^{1,2}, Elena Aller^{1,2}, Carla Fuster-García¹, Gema García-García^{1,4}, Regina Rodrigo^{1,2}, Rafael P Vázquez-Manrique^{1,2}, Fiona Blanco-Kelly^{2,3}, Carmen Ayuso^{2,3}, Anne-Françoise Roux⁴, Teresa Jaijo^{1,2†} and José M Millán^{1,2,5*†}

Abstract

Background: Usher syndrome is an autosomal recessive disease that associates sensorineural hearing loss, retinitis pigmentosa and, in some cases, vestibular dysfunction. It is clinically and genetically heterogeneous. To date, 10 genes have been associated with the disease, making its molecular diagnosis based on Sanger sequencing, expensive and time-consuming. Consequently, the aim of the present study was to develop a molecular diagnostics method for Usher syndrome, based on targeted next generation sequencing.

Methods: A custom HaloPlex panel for Illumina platforms was designed to capture all exons of the 10 known causative Usher syndrome genes (MYO7A, USH1C, CDH23, PCDH15, USH1G, CIB2, USH2A, GPR98, DFNB31 and CLRN1), the two Usher syndrome-related genes (HARS and PDZD7) and the two candidate genes VEZT and MYO15A. A cohort of 44 patients suffering from Usher syndrome was selected for this study. This cohort was divided into two groups: a test group of 11 patients with known mutations and another group of 33 patients with unknown mutations.

Results: Forty USH patients were successfully sequenced, 8 USH patients from the test group and 32 patients from the group composed of USH patients without genetic diagnosis. We were able to detect biallelic mutations in one USH gene in 22 out of 32 USH patients (68.75%) and to identify 79.7% of the expected mutated alleles. Fifty-three different mutations were detected. These mutations included 21 missense, 8 nonsense, 9 frameshifts, 9 intronic mutations and 6 large rearrangements.

Conclusions: Targeted next generation sequencing allowed us to detect both point mutations and large rearrangements in a single experiment, minimizing the economic cost of the study, increasing the detection ratio of the genetic cause of the disease and improving the genetic diagnosis of Usher syndrome patients.

Keywords: Usher syndrome, Molecular diagnosis, Next generation sequencing, Point mutations, Large rearrangements

Background

Usher syndrome (USH) is an autosomal recessive disease characterized by the association of sensorineural hearing loss, retinitis pigmentosa (RP) and, in some cases, vestibular dysfunction. Its prevalence ranges from 3 to 6.2 per 100,000 [1-3] and it is the most common form of hereditary syndromes combining hearing loss and retinitis pigmentosa.

USH has a remarkable clinical and genetic heterogeneity. According to the severity and progression of the disease, three clinical types are distinguished. Type I (USH1) is defined by profound congenital hearing loss, onset of RP usually within the first decade of life and an absence of vestibular function. Usher syndrome type II (USH2) patients display congenital moderate-severe hearing loss, onset of RP around or after puberty and normal vestibular function. Usher syndrome type III (USH3) is characterized by postlingual progressive hearing loss, RP with a variable age of onset and variable vestibular response [4]). However, in some patients the disease does not fit into any of these three subtypes, and they are classified as 'atypical Usher syndrome'.

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To date, 10 genes and three additional loci have been associated with the disease. For USH1 six genes have been identified: *MYO7A* (USH1B), *USH1C* (USH1C), *CDH23* (USH1D), *PCDH15* (USH1F), *USH1G* (USH1G) and *CIB2* (USH1J); and two additional loci have been described: USH1E [5] and USH1H [6]. Three genes have been found to cause USH2: *USH2A* (USH2A), *GPR98* (USH2C) and *DFNB31* (USH2D). For USH3 only mutations in *CLRN1* have been described, and a second locus was proposed, USH3B. Furthermore, most of these genes are also responsible for non-syndromic hearing loss or isolated RP [4,7].

In addition to these 10 genes, three other genes have been associated with USH. *PDZD7* was proposed as a contributor of digenic inheritance with *GPR98*, and as a retinal disease modifier in *USH2A* patients [8]. Recently, a novel biallelic missense variant in *HARS* was identified in two patients with a phenotype compatible with USH3. This gene encoding an aminoacyl tRNA synthetase was proposed as a novel gene causative of USH3 [9]. More recently, *CEP250* has been associated with atypical Usher syndrome [10].

Proteins encoded by Usher genes belong to different classes: actin-based motor protein, scaffolding proteins, cell-adhesion molecules or transmembrane proteins, some of them with very large extracellular domains. These proteins through their protein-protein interacting domains are integrated in a network known as "Usher-interactome". The main sites of colocalization of Usher proteins are the stereocilia or hair bundle of the inner ear hair cells, and the synaptic and periciliary areas of the photoreceptors [4,11].

In the stereocilia, two different USH protein networks are known to be involved in the development and maintenance of the inner ear hair cells. One of them is composed of the USH2 proteins, myosin VIIA (encoded by MYO7A), Vezatin and PDZD7 [12,13]. Vezatin is encoded by VEZT and it is required for the sound resilience of cochlear hair cells. In addition, it has been proposed that it is involved in the maturation steps of these cells or in the maintenance of junction integrity between hair cells and supporting cells in the inner ear [14]. The second stereocilia interactome is composed of the USH1 proteins [15]. In this network, several unconventional myosins, as myosin XVa, have also been found [16-18]. Mutations in MYO15A, the gene encoding Myosin XVa, are responsible for non-syndromic autosomal recessive hearing loss DFNB3 [19]. In addition, Myosin XVa interacts with whirlin in stereocilia, considered a key event in hairbundle morphogenesis [20], and this direct interaction at the stereocilia tip is likely to control the elongation of stereocilia [21].

In the retina, two similar molecular networks composed of USH2 and USH1 proteins have also been described.

The USH2 protein network, localized at the periciliary ridge complex region, contributes to the trafficking of cargos moving from the inner segment to the outer segment of the photoreceptor cells through the connecting cilium. The USH1 interactome has been reported to localize at the membrane-membrane connection site between the outer segment of the photoreceptor cells and the calyceal processes in primates. This USH1 protein complex is thought to form an adhesion belt and would contribute to the daily renewal of photoreceptors outer segments [22-24].

Traditionally, the molecular diagnosis of Usher syndrome has been mainly based on Sanger sequencing [25,26]. However, the large size of most USH genes (above 350 exons in total) makes this technique expensive and time-consuming. Array-based mutation screenings (arrayed primer extension (APEX) technology) have become a rapid and efficient technique to detect previously described mutations, and a specific APEX-microarray for USH was developed [26]. However, most of USH mutations are private, so the low detection rate of the APEXmicroarray for Usher syndrome hampers the use of this technology [27,28]. Furthermore, Copy Number Variants (CNVs) have been identified as an important cause of the disease in Usher syndrome. Deletions and duplications are screened by Multiplex Ligation-dependent Probe Amplification (MLPA), for which commercial probemix is only available for USH2A and PCDH15, or by array-Comparative Genomic Hibridization (a-CGH) [29-31].

In recent years, next generation sequencing (NGS) techniques have been developed further, permitting the whole genome, whole exome and targeted gene sequencing to be more feasible, and hence making the identification of diseased genes and the underlying mutations easier, rapid and cost-effective. The improvement of NGS has been especially useful in the diagnosis of genetically heterogeneous diseases, such as hearing loss or retinal dystrophies [32-35].

For Usher syndrome, several NGS methods have been developed. Licastro et al. [36] used two different approaches: whole exome sequencing with the use of the SOLiD system and long-PCR sequencing on nine USH genes with two different platforms, Illumina (Genome Analyzer II) and the Roche 454 (GS FLX). Recently, Besnard et al. [37] developed a targeted NGS approach. They included 9 USH genes, 2 candidate USH genes (VEZT and PDZD7), seven hearing-loss genes and the choroideremia-causative gene CHM and the sequencing was carried out on Roche GS Junior sequencer. Yoshimura et al. [38] has also applied a massively parallel DNA sequencing methodology, but only for USH1 patients. Recently, Rong W et al. [39] used the NGS approach to identify three new alleles and one known mutation in MYO7A in three Chinese families.

We developed a molecular diagnosis of Usher syndrome based on a targeted NGS technique, HaloPlex (Agilent) gene target enrichment for Illumina platforms, including the ten USH known genes and four candidate genes that allowed us to identify not only point mutations but also CNVs, implementing a platform for the genetic diagnosis of this disease.

Methods

Patients

A cohort of 44 patients diagnosed with Usher syndrome was selected for this study. Their clinical classification into USH1, USH2, USH3 or atypical USH was performed on the basis of their clinical history and ophthalmologic, audiometric and vestibular tests. Patients included in this work were divided into two groups: a test group and a group composed by USH patients without genetic diagnosis.

Whenever possible, samples from additional family members were used to perform segregation analysis of the sequence variants identified in the index patient.

All patients and relatives included in the present study signed authorizing their written informed consent. The study was approved by the institutional board of the Ethics Committee of the University Hospital La Fe.

Test group

The test group was composed of 11 patients: five USH1, four USH2, one USH3 and one atypical USH. These patients had been screened using a combination of

techniques as Sanger sequencing, APEX microarray for Usher syndrome or MLPA. Variants of different nature in six USH genes had been previously detected and were used as positive controls in the present study. Table 1 shows the previously identified changes in the test group.

Cohort of USH patients previously unscreened

A cohort of 33 USH patients without genetic diagnosis was enrolled in this study. It was composed of 13 USH1 patients, 17 USH2 and 3 USH3 cases.

DNA Samples

Genomic DNA from patients and relatives was extracted from EDTA blood using an automated DNA extractor (Magna Pure, Roche). DNA samples were purified with the "QIAqick PCR Purification Kit" following the manufacturer's instructions. The concentration of the genomic DNA was determined with the "Qubit dsDNA BR Assay Kit" in the Qubit 2.0 fluorometer.

Targeted next generation sequencing design

A custom HaloPlex panel was designed using Agilent's SureDesign tool (www.agilent.com/genomics/suredesign) to capture all exons and 25 bp of intronic flanking regions of 14 genes. The target genes included in our work were the 10 causative Usher syndrome genes (MYO7A, USH1C, CDH23, PCDH15, USH1G, CIB2, USH2A, GPR98, DFNB31 and CLRN1), two related USH genes (HARS and PDZD7) and two candidate genes VEZT and MYO15A, due to their involvement in the Usher interactome

Table 1 Test group: patients carrying sequence variants in USH genes previously detected

Patient	Clinical type	Gene	Type variant	Nucleotide	Protein	Classification
RP-808	USH1	CDH23	Intronic	c.6059-9G > A		Pathogenic. UV4
		USH1G	Missense	c.387A > G	p.Lys130Glu	UV1
		USH1G	Missense	c.423G > A	p.Glu142Lys	UV2
RP-1145	USH2	USH2A	Frameshift duplication	c.10272-10274dup	p.Cys3425Phefs*4	Pathogenic. UV4
		USH2A	Nonsense	c.7854G > A	p.Trp2618*	Pathogenic. UV4
		DFNB31	Isocoding	c.117G > A	p.Val39Val	UV1
RP-1286 [§]	USH1	PCDH15	Frameshift insertion	c.1304_1305insC	p.Thr436Tyrfs*12	Pathogenic. UV4
RP-1374	USH1	PCDH15 Nonsense c.7C > T		c.7C > T	p.Arg3*	Pathogenic. UV4
RP-1426	USH1	MYO7A	Missense	c.6610G > C	p.Ala2204Pro	Pathogenic
RP-1522	USH2	USH2A	Frameshift deletion	c.2299delG	p.Glu767Serfs*21	Pathogenic. UV4
		CDH23	Missense	c.1096G > A	p.Ala366Thr	UV2
RP-1608 [§]	USH3	USH2A	Missense	c.9799 T > C	p.Cys3267Arg	Pathogenic. UV4
RP-1614	USH1	MYO7A	Frameshift deletion	c.6025delG	p.Ala2009Profs*32	Pathogenic. UV4
RP-1637	USH2	USH2A	Frameshift duplication	c.5540dupA	p.Asn1848Glufs*20	Pathogenic. UV4
		USH2A	Large deletion	Del IVS4_IVS9	p.Gly262Valfs*2	Pathogenic. UV4
RP-1757 [§]	Atypical	MYO7A	In-frame deletion	c.655_660del	p.lle219_His220del	Pathogenic. UV4
RP-1760	USH2	USH2A	Missense	c.2296 T > C	p.Cys766Arg	UV3

Samples that failed in the amplification process are marked with §.

[14,20,21]. The novel *CEP250* [10] had not been associated with USH at the beginning of this study, so it was not included in our work.

The entire custom design was 481 targets with a total size of 132.763 Kb. The exons from the different isoforms included in this study are summarized in Table 2. The intronic region where the *USH2A* mutation c.7595-2144A > G [37] is located, was included in the design. The final probe design size was 296.74 Kb with a theoretical coverage of 99.88% for our targeted regions. The difference in size between our entire custom design and the final probe design was due to an intrinsic process of the engineering of probes by Agilent's Sure Design tool for the HaloPlex protocol.

Sequence capture and next generation sequencing

Sequence capture was performed according to the "HaloPlex Target Enrichment System" (Protocol Version

D.5, Agilent Technologies Inc, CA, USA) for Illumina Sequencing. Between 225 to 450 ng of gDNA were digested by 16 different restriction enzymes to create a library of gDNA restriction fragments. Four 12-reaction runs were performed including in each of them 11 gDNA samples and one Enrichment Control DNA sample. These enzymatic digestions were validated by electrophoretic analysis in a polyacrylamide gel. The gDNA restriction fragments were hybridized to the HaloPlex probe capture library. In this step, the Illumina sequencing motifs including index sequences were incorporated into the targeted fragments and the target DNA-HaloPlex probe hybrids were circularized. These hybrid molecules were captured using streptavidin beads. The DNA ligase was then added to close nicks in the circularized HaloPlex probe-target DNA hybrids and the captured DNA libraries were eluted with NaOH.

Table 2 Details about the exons studied in this targeted NGS analysis for the 14 genes analyzed

Chr	Gene	RefSec	Coding exons	Size (bp)	Coding Size (bp)		Alternative Ref Sec	Additional exons and the intronic <i>USH2A</i> region included in the study	Size (bp)	Number of exons analyzed
11	MYO7A	NM_000260.3 variant 1	48	7465	6648	2215				48
10	CDH23	NM_022124.5 variant 1	69	11134	10065	3354	NM_052836.3 variant 2	1	150	70
10	PCDH15	NM_033056.3 variant C	32	7021	5868	1955	NM_001142773.1 variant H	1	6	38
							NM_001142769.1 variant l	2	871	
							NM_001142771.1 variant K	2	4469	
11	USH1C	NM_153676 variant 3	27	3246	2700	899	NM_005709.3 variant 1	1	75	28
17	USH1G	NM_173477.4 variant 1	3	3565	1386	461				3
15	CIB2	NM_006383.2 variant 1	6	1580	564	187				6
1	USH2A	NM_206933.2 variant 2	71	18883	15609	5202	NC_000001.11	c.7595-2144A > G	152	71 + 1 intronic sequence
5	GPR98	NM_032119.3 variant 1	90	19333	18921	6307				90
9	DFNB31	NM_015404.3 variant 1	12	4079	2724	907	NM_001083885.2 variant 2	1	110	13
3	CLRN1	NM_174878.2 variant 1	3	2359	699	232	NM_052995.2 variant 4	1	20	4
10	PDZD7	NM_001195263.1 variant 1	17	4164	3102	1033	NM_024895.4 variant 2	1	285	18
5	HARS	NM_002109.5 variant1	13	2322	1530	509				13
17	MYO15A	NM_016239.3	65	11876	10593	3530				65
12	VEZT	NM_017599 variant 1	12	4580	2340	779	NR_038242.1 variant 2	1	74	13
Total targ	ets									481

A PCR amplification of the captured target libraries was performed following the manufacturer's instructions and, after its purification with the "AMPure XP beads" (BECKMAN CULTER Inc), the validation and quantification of the enriched target DNA in each library was performed using the 2100 Bioanalyzer system with the High Sensitivity DNA Kit and the 2100 Expert Software (Agilent Technologies Inc, CA, USA). The last step of the protocol was to pool samples for multiplexed sequencing in the Illumina sequencing platform MiSeq System (Illumina,Inc).

Variant analysis

Data was analyzed using the platform provided by DNA-nexus (www.dnanexus.com). Two different versions were used: DNAnexus Classic and the DNAnexus version recently implemented. Annotated variants were selected according to the following criteria: the quality value should be ≥250, a percentage of heterozygosity ≥30% of the reads, their annotation in the dbSNP (http://www.ncbi.nlm.nih.gov/SNP) and their description in the Usher syndrome mutation database (https://grenada.lumc.nl/LOVD2/Usher_montpellier). Variants should be observed in both direct and reverse strands.

Variants selected and suspected to be pathogenic were confirmed by Sanger sequencing. The DNA fragments containing the variants were amplified by PCR with specific primers and were sequenced on both strands using the Big Dye 3.1 Terminator Sequencing Kit. The purified sequence products were analyzed on a 3500xl ABI instrument (Applied Biosystems by Life Technologies, Thermo Fisher Scientific, Inc).

The pathogenicity of novel missense variants was analyzed with the SIFT (http://sift.bii.a-star.edu.sg/) and PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/) algorithms. Those putative variants that affect the splicing process were studied with the NetGene2 (http://www.cbs.dtu.dk/services/NetGene2), NNSPLICE v0.9 (http://www.fruitfly.org/seq_tools/splice.html), Human Splicing Finder (HSF; http://www.umd.be/HSF) and RESCUE-ESE (http://genes.mit.edu/burgelab/rescue-ese/) programs.

Novel variants were classified based on the classification system for Unknown Variants (UV) (https://grenada.lumc.nl/LOVD2/Usher_montpellier) as pathogenic (UV4), possibly pathogenic (UV3), possibly non-pathogenic (possibly neutral, UV2) and non-pathogenic (neutral, UV1) according to bioinformatics predictions and segregation analysis. This classification is in line with the guidelines published by the clinical and molecular genetics society (http://cmgdweb.shared.hosting.zen.co.uk/BPGs/Best_Practice_Guidelines.htm).

Nomenclature of variants was performed according to the reference sequences: *MYO7A* (NM_000260.3), *USH1C* (NM_153676), *CDH23* (NM_022124.5), *PCDH15*

(NM_033056.3), *USH1G* (NM_173477), *CIB2* (NM_006383.2), *USH2A* (NM_206933), *GPR98* (NM_032119.3), *DFNB31* (NM_015404), *CLRN1* (NM_174878), *HARS* (NM_002109), *PDZD7* (NM_001195263.1), *VEZT* (NM_017599) and *MYO15A* (NM_016239).

Segregation analysis was performed in cases where DNA samples of relatives were available.

Copy number variation analysis and validation

The coverage of every target region of the sample of interest was normalized and compared with average normalized data of all other samples of the same run to obtain the ratio relative coverage. Deletions and duplications were suspected of being present if this ratio fell below 0.7 or rose above 1.3 respectively.

Validation of rearrangements of *USH2A* and *PCDH15* was performed by MLPA analysis. For *USH2A* the SALSA MLPA probemixes P361 and P362 were used. To confirm the *PCDH15* copy number variations the SALSA MLPA probemix P292 was employed. The MLPA reactions were performed according to the manufacturer's recommendations (http://www.mlpa.com).

In the cases of putative rearrangements identified in *CDH23* and *GPR98*, their validation was carried out by an oligonucleotide array-CGH. The array-CGH chip included 77,366 probes covering the genes *MYO7A*, *CDH23*, *PCDH15*, *USH1C*, *USH1G*, *USH2A*, *GPR98*, *DFNB31*, *PDZD7* and *CLRN1* and 10.000 nucleotides of 5' and 3' untranslated regions [30].

Results

Next generation sequencing results of the USH panel

Capture of NGS using our customized USH panel was performed in a cohort of USH patients. High quality results were obtained. On average, a mean coverage of 1334x was obtained per sample and target region. Different average coverages were obtained for the fourteen analyzed genes, ranging from 935x (VEZT) to 1817x (CLRN1) (Figure 1).

Only one of the selected targeted regions showed coverage lower than 40x, the defined limit for proper validation and diagnostic procedures. This target was a coding region of 43 bp in the large exon 2 of *MYO15A*. A schematic representation of the coverage of all target regions is showed in Figure 2.

Test group: validation of the diagnostic strategy

Eleven USH patients were included in the test group to verify the reliability of our NGS custom panel. In these cases, 17 mutations and polymorphism had been previously detected in different USH genes by Sanger or by MLPA (Table 1).

During the experiment three control patients and one studied patient failed in the capture step of the

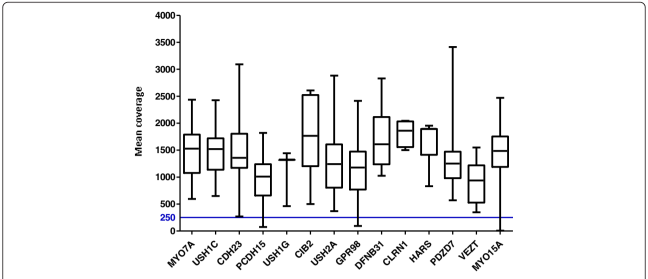


Figure 1 Mean coverage obtained for the different genes. The blue line shows the lower limit of coverage appropriate to perform CNV analyses (250x).

target DNA of the HaloPlex protocol. Due to a manual/technical problem with the PCR strips and magnetic rack Dynabeads, the circularized target DNA-HaloPlex probe hybrids, containing biotin, were not captured on streptavidin beads and these samples were lost.

The remaining eight patients were carriers of 14 different variants including missense, nonsense, point deletions or insertions and large deletions (Table 1). Our NGS approach allowed us to detect all previously detected changes in the testing cohort, both point mutations and CNVs.

The sequencing of the USH genes allowed the detection of a second pathologic allele in five patients in whom only one mutation was previously detected. In addition, in patient RP-1426, previously a carrier of a pathogenic

variant in *MYO7A*, we were able to identify a novel splice site mutation in *CDH23*. These results are summarized in Table 3.

Cohort of USH patients previously unscreened

In our cohort of 33 USH patients studied, mutations could be identified in the majority of analyzed cases. One of the USH cases failed in the capture and amplification of the library due to the same manual/technical error explained above. If we take into account the remaining 32 USH cases analyzed, we identified in 22 patients the two expected mutated alleles, in 5 cases only one pathogenic variant and in one case, RP-1950, two pathogenic variants were detected in two different USH2 genes. In four patients no mutation was found. In addition two patients

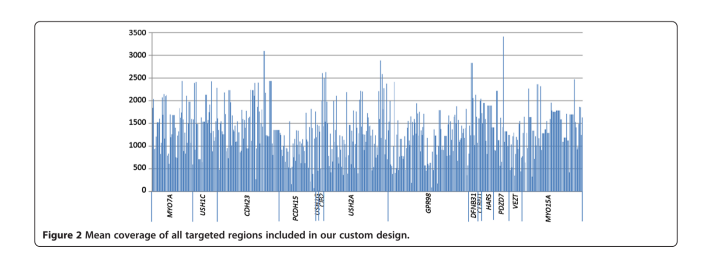


Table 3 Causative mutations and putative pathogenic variants identified in this study

Patient	Clinical type	Gene	Exon	Nucleotide variant	Protein variant	Reference	Segregation analysis
Patients	with two path	ogenic m	utations	in the same gene			
RP-807	USH2	MYO7A	40	c.5516 T > C	p.Leu1839Pro	Novel. UV3	Yes
		MYO7A	27	c.3503G > A	p.Arg1168Gln	Novel. UV3	
RP-808 [¶]	USH1	CDH23	47	c.6059-9G > A	_	von Brederlow et al., (2002) [40]	No
		CDH23	10	c.871G > A	p.Gly291Arg	Novel. UV3	
RP-890	USH3	USH2A	26	c.5278delG	p.Asp1760Metfs*10	Garcia-Garcia et al., (2011) [41]	No
		USH2A	26	c.5278delG	p.Asp1760Metfs*10	Garcia-Garcia et al., (2011) [41]	
RP-1182	USH1	PCDH15	22_23	Duplication exons 22_23	_	Novel. UV4	No
		PCDH15	22_23	Duplication exons 22_23	_	Novel. UV4	
RP-1183	USH1	CDH23	26	c.3016G > A	p.Glu1006Lys	Schultz et al., (2011) [42]	No
		CDH23	26	c.3016G > A	p.Glu1006Lys	Schultz et al., (2011) [42]	
RP-1234	USH1	MYO7A	43	c.5884delTTCT	p.Phe1962Leufs*7	Novel. UV4	Yes
		MYO7A	43	c.5884delTTCT	p.Phe1962Leufs*7	Novel. UV4	
RP-1237	USH1	CDH23	46	c.6049G > A	p.Gly2017Ser	Roux et al., (2006) [43]	No
		CDH23	46	c.6049G > A	p.Gly2017Ser	Roux et al., (2006) [43]	
RP-1374 [¶]	USH1	PCDH15	2	c.7C > T	p.Arg3*	Ahmed et al., (2001) [44]	Yes
		PCDH15	27	c.3717 + 2dupT	_	Jaijo et al., (2012) [45]	
RP-1422	USH1	MYO7A	43	c.5944G > A	p.Gly1982Arg	Riazuddin et al., (2008) [46]	Yes
		MYO7A	43	c.5944G > A	p.Gly1982Arg	Riazuddin et al., (2008) [46]	
RP-1522 [¶]	USH2	USH2A	13	c.2299delG	p.Glu767Serfs*21	Liu et al., (1999) [47]	No
		USH2A	20	Deletion exon 20	_	Novel. UV4	
RP-1551	USH1	PCDH15	27	c.3511delA	p.Asp1172llefs*13	Novel. UV4	Yes
		PCDH15	27	c.3511delA	p.Asp1172llefs*13	Novel. UV4	
RP-1614 [¶]	USH1	MYO7A	44	c.6025delG	p.Ala2009Profs*32	Bharadwaj et al., (2000) [48]	No
		MYO7A	40	c.5537C > A	p.Pro1846His	Novel. UV3	
RP-1760 [¶]	USH2	USH2A	55	c.10888delA	p.Gly3631Valfs*43	Novel. UV4	No
		USH2A	13	c.2296 T > C	p.Cys766Arg	Glöcke et al., (2013) [35]	
RP-1781	USH2	CDH23	29	Duplication exon 29	_	Novel. UV4	No
		CDH23	68	c.9569C > T	p.Ala3190Val	Novel. UV3	
RP-1791	USH1	MYO7A	20	c.2283-1G > T	_	Roux et al., (2006) [43]	Yes
		MYO7A	28	c.3594C > A	p.Cys1198*	Roux et al., (2011) [43]	
RP-1802	USH2	USH2A	63	c.13811 + 2 T > G	_	Besnard et al., (2014) [37]	No
		USH2A	50	c.9799 T > C	p.Cys3267Arg	Aller et al., (2006) [49]	
RP-1835	USH2	USH2A	57	c.11065C > T	p.Arg3689*	Le Quesne Stabej et al., 2012 [50]	Yes
		USH2A	22	c.4758 + 3A > G	_	Novel. UV3	
RP-1864	USH2	MYO7A	6	c.494C > T	p.Thr165Met	Ouyang et al., (2005) [51]	No
		MYO7A	6	c.494C > T	p.Thr165Met	Ouyang et al., (2005) [51]	
RP-1895	USH2	GPR98	79_83	Duplication exons 79_83	_	Besnard et al., (2012) [52]	No
		GPR98	79_83	Duplication exons 79_83	_	Besnard et al., (2012) [52]	
RP-1904	USH2	GPR98	11	c.2145_2149delGTTTT	p.Leu715Phefs*6	Novel. UV4	Yes
		GPR98	14	c.2612delG	p.Gly871Glufs*8	Novel. UV4	
RP-1910	USH1	CDH23	60	c.8722 + 1delG	_	Oshima et al., (2008) [53]	Yes
		CDH23	60	c.8722 + 1delG	_	Oshima et al., (2008) [53]	

Table 3 Causative mutations and putative pathogenic variants identified in this study (Continued)

RP-1924	USH1	MYO7A	39	c.5392C > T	p.Gln1798*	Janecke et al., (1999) [54]	Yes
		MYO7A	27	c.3503G > A	p.Arg1168Gln	Novel. UV3	
RP-1927	USH2	USH2A	21	c.4474G > T	p.Glu1492*	Bernal et al., (2005) [55]	Yes
		USH2A	2	c.269A > G	p.Tyr90Cys	Novel. UV3	
RP-1948	USH1	MYO7A	7	C.707 T > A	p.Leu236Gln	Novel. UV3	No
		MYO7A	42	c.5749G > T	p.Glu1917*	Jacobson et al., (2009) [56]	
RP-1960	USH2	USH2A	25	c.5167G > C	p.Gly1723Arg	Novel. UV3	No
		USH2A	7	c.1214delA	p.Asn405llefs*3	Bernal et al., (2005) [55]	
Patients	with three pa	thogenic i	mutatio	ns in two different genes			
RP-1847	USH2	USH2A	62	c.12067-2A > C	_	Kaiserman et al., (2007) [57]	Yes
		USH2A	14	Deletion exon 14	_	Glöckle et al., (2013) [35]	
		USH1G	2	c.805C > T	p.Arg269*	Novel. UV4	
RP-1923	USH2	USH2A	62	c.12093delC	p.Tyr4031*	Garcia-Garcia et al., (2011) [41]	No
		USH2A	44	Deletion exon 44	_	Glöckle et al., (2013) [35]	
		DFNB31	9	c.2234G > A	p.Arg745His	Novel. UV3	
Patients	with only one	pathoge	nic muta	ation			
RP-1455	USH1	USH2A	28	c.5666A > G	p.Asp1889Gly	Novel. UV3	No
RP-1496	USH3	GPR98	19	c.3443G > A	p.Gly1148Asp	Novel. UV3	No
RP-1741	USH2	USH2A	PE40	c.7592-2144A > G	_	Vaché et al., (2012) [58]	No
RP-1929	USH2	GPR98	58	c.11974G > A	p.Asp3992Asn	Novel. UV3	No
RP-1953	USH2	USH1C	18	c.1859G > T	p.Arg620Leu	Ouyang et al., (2002) [59]	No
Patients	with patholog	gic mutatio	ons in d	lifferent genes			
RP-1426 [¶]	USH1	MYO7A	49	c.6610G > C	p.Ala2204Pro	Jaijo et al., (2007) [60]	Yes
		CDH23	39	c.5068-2A > T	_	Novel. UV4	
RP-1950	USH2	USH2A	70	c.2299delG	p.Glu767Serfs*21	Liu et al., (1999) [47]	No
		GPR98	70	c.14278C > T	p.Pro4760Ser	Novel. UV3	

Patients previously included in the test group are marked with $\P.$

Novel variants are marked in bold.

PE: Pseudoexon 40.

(RP-1847 and RP-1923) were carriers of three mutations in USH genes (Table 3).

In this study, we were able to identify 53 different mutations, of which 24 were novel. These variants included 21 missense, 8 nonsense, 9 frameshifts, 9 intronic mutations and 6 large rearrangements.

The DNA nexus program showed a high number of novel variants in each patient. These changes were selected according to their nature, or the presence of another pathogenic allele in the same gene in the same patient. The variants were confirmed by Sanger sequencing and missense, isocoding or intronic variants were analyzed by different *in silico* algorithms to predict their pathogenic effect. According to the bioinformatics predictions, 15 variants out of 25 studied variants were considered as pathogenic (UV4) or possibly pathogenic (UV3) (Table 4).

Segregation analysis of detected alleles with the respective family members was performed in 13 cases, and

cosegregation of the mutations with the disease was verified in all these cases (Table 3).

CNVs Detection

Next Generation Sequencing with the Illumina MiSeq system allowed us to perform qualitative and quantitative analysis. We could detect not only point mutations but also CNVs.

In our cohort of patients we performed CNV analysis in all patients where two point mutations had not been detected. This analysis allowed us to identify six large rearrangements: three deletions in *USH2A* comprising the exons 14, 20 and 44, and three duplications comprising the exon 29 in *CDH23*, exons 22_23 in *PCDH15* and exons 79_83 in *GPR98*. The *USH2A* and the *CDH23* rearrangements were detected in a heterozygous state, whereas the *PCDH15* and *GPR98* duplications were identified in a homozygous state.

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Table 4 Summary of putative pathogenic mutations and their bioinformatics predictions

Patient	Gene	Exon	Nucleotide Change	Amino Acid Change	Classification	SIFT (score)	PolyPhen-2 (score)	NetGene2	Human Splicing Finder	NNSPLICE	RESCUE-ESE
RP-1948	MYO7A	7	c.707T>A	p.Leu236Gln	UV3	deleterious (0)	probably_damaging (1)	Neutral	Neutral	Neutral	A new ESE site is created
RP-807 RP-1924	MYO7A	27	c.3503G>A	p.Arg1168Gln	UV3	deleterious (0)	probably_damaging (1)	Score for the main donor site decreases from 95 to 75	Score for the main donor site decreases from 90 to 80	The main donor site is not recognized	Neutral
RP-807	MYO7A	40	c.5516T>C	p.Leu1839Pro	UV3	deleterious (0)	probably_damaging (1)	Score for the main acceptor site decreases from 80 to 77	Neutral	Neutral	Neutral
RP-1614	MYO7A	40	c.5537C>A	p.Prp1846His	UV3	deleterious (0.01)	possibly_damaging (0.85)	Score for the main acceptor site decreases from 80 to 77 and one acceptor site is not recognized	Neutral	Neutral	Neutral
RP-808	CDH23	10	c.871G>A	p.Gly291Arg	UV3	deleterious (0)	probably_damaging (1)	Neutral	Neutral	Neutral	A new ESE site is created
RP-1426	CDH23	39	c.5068-2A>T	c.5068-2A>T	UV4	_	_	The main acceptor site is not recognized	Score for the main acceptor site decreases from 95 to 66	The main acceptor site is not recognized	_
RP-1781	CDH23	68	c.9569C>T	p.Ala3190Val	UV3	deleterious (0)	probably_damaging (1)	Neutral	Neutral	Neutral	Neutral
RP-1927	USH2A	2	c.269A>G	p.Tyr90Cys	UV3	tolerated (0.17)	possibly_damaging (0.796)	Score for acceptor site decreases from 82 to 80	Neutral	Score for acceptor site decreases from 75 to 69	A ESE site is not recognized
RP-1835	USH2A	22	c.4758+3A>G	c.4758+3A>G	UV3	_	_	The main donor site is not recognized	Neutral	The main donor site decreases from 98 to 73	_
RP-1960	USH2A	25	c.5167G>C	p.Gly1723Arg	UV3	deleterious (0)	probably_damaging (0.994)	The main donor site is not recognized	Score for the main donor site decreases from 86 to 75	The main donor site is not recognized	Neutral
RP-1455	USH2A	28	c.5666A>G	p.Asp1889Gly	UV3	deleterious (0)	probably_damaging (0.982)	The main donor and acceptor sites decrease from 82 to 80 and from 53 to 48 respectively and a new acceptor site is created	Neutral	Neutral	Two ESEs are not recognized
RP-1496	GPR98	19	c.3443G>A	p.Gly1148Asp	UV3	deleterious (0)	probably_damaging (0.999)	Neutral	Neutral	Neutral	Neutral
RP-1929	GPR98	58	c.11974G>A	p.Asp3992Asn	UV3	deleterious (0.01)	probably_damaging (0.999)	Neutral	Neutral	Neutral	A ESE is not recognized

RP-1950 <i>GPR98</i>	70	c.14278C>T	p.Pro4760Ser	UV3	deleterious (0)	probably_damaging (0.998)	Neutral	Neutral	Neutral	Neutral
RP-1923 <i>DFNB31</i>	9	c.2234G>A	p.Arg745His	UV3	deleterious(0.01)	probably_damaging (0.984)	Neutral	Neutral	The main donor site increases from	Neutral
									66 to 86	

SIFT: SIFT Score ranges from 0 to 1. The amino acid substitution is predicted to be damaging if the score is < 0.05, and tolerated if the score is > 0.05.

PolyPhen stablish three classifications: "Probably damaging" (it is believed most likely to affect protein function or structure), "Possibly damaging" (it is believed to affect protein function or structure), "Benign" (most likely lacking any phenotypic effect).

ESE: Exonic Splicing Enhancer.

Large rearrangements in *USH2A* and *PCDH15* were subsequently confirmed by MLPA, whereas patients that carried duplications affecting *CDH23* and *GPR98* were analyzed by a-CGH (data not shown). In 5 patients, the rearrangements detected by comparison of normalized coverage data were confirmed. The a-CGH technique to confirm the presence of the heterozygous duplication of *CDH23* (exon 29) in patient RP-1781 was not successful due to technical problems. Unfortunately, it was impossible to get a new DNA sample for this patient to repeat the experiment.

Three of the six rearrangements found in this study had been previously described. The *USH2A* deletions of exon 14 and exon 44 were described by Glockle et al. [35] and the *GPR98* duplication was previously described by Besnard et al. [52]. The deletion of exon 14 in *USH2A* has also been detected by our group, in one Spanish USH patient in a homozygous state (unpublished results).

In our cohort of patients previously unscreened we could detect 51 altered alleles, six of which corresponded to large rearrangements, i.e. 11.76% of the detected pathogenic alleles.

In one patient no point mutation was identified, and the quantitative analysis could not be performed (RP-531). The ratio between this sample and the normalized data was altered in a high number of target regions from most genes. In this case, the image obtained with the bioanalyzer in the validation and quantification of the enriched target DNA process was atypical (Figure 3).

In some cases, we could observe that the CNV analysis displayed doubtful results in the targeted regions. The analysis of those target regions revealed coverages lower than 250x.

Discussion

Resolved cases

Usher syndrome displays both genetic and allelic heterogeneity. The number of genes related to the disease and the large size of most of them contribute to the genetic heterogeneity. Furthermore, mutations causing disease include point mutations and large rearrangements, most of them, being private. Recently, the high-throughput sequence analysis and next generation sequencing, has

been proved to be very helpful to carry out genetic studies in heterogeneous diseases.

In this study we have developed a targeted method based on NGS for the molecular diagnosis of Usher syndrome. We evaluated our panel in a group of patients with known USH variants and applied it to a cohort of USH patients previously unscreened.

In our series of study, we were able to detect biallelic mutations in one USH gene in 22 out of 32 USH patients (68.75%) and to identify 51 out of the 64 expected mutated alleles (a detection ratio of 79.7%).

Uncovered regions

We obtained a highly heterogeneous coverage using a custom *HaloPlex Target Enrichment System* (see Figure 1). However, only one region included in the final design showed a mean coverage lower than 40x in our samples. It corresponded to a 43 bp sequence in the large exon 2 of *MYO15A*. That fragment shows a high percentage of CGs, 77%. It has been reported that most sequencing platforms, including Illumina sequencers, show a GC content-dependent bias coverage, and protocols should be modified to minimize these events [61].

Unresolved cases

Mutations were not detected in four patients of our cohort. In one USH1 case detailed clinical data could not be obtained, whereas in the remaining cases clinical information was revised and the USH diagnosis was confirmed. Five additional patients were carriers of only one pathogenic variant (Table 3), the second mutation remaining unidentified.

In these cases, in which the second mutated allele escapes detection, the region that contains the mutation may have been excluded from our study. It is possible that the gene responsible for the disease had not yet been identified, and therefore it could not be included in our panel. Furthermore, deep intronic regions, promoter regions or 5' and 3' untranslated regions were not systematically included in the panel design. The only intronic region targeted in our panel was designed to detect the *USH2A* c.7595-2144A > G mutation identified by Vaché et al. [58] in five French patients and four Spanish patients. Moreover, it was subsequently detected

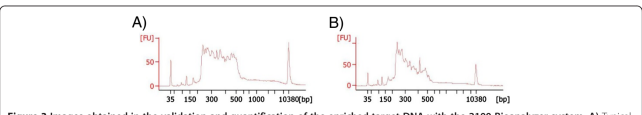


Figure 3 Images obtained in the validation and quantification of the enriched target DNA with the 2100 Bioanalyzer system. A) Typical image obtained in most patients. B) Atypical image obtained in the patient RP-531, in whom CNV analysis could not be performed.

in five additional families [31]. Subsequently, Steele-Stallard et al. [31] detected this intronic change in three UK and European USH families.

In two patients, two mutated alleles, each one in a different gene, were detected (Table 3). Patient RP-1426 was carrier of a missense MYO7A variant (p.Ala2204Pro). Using the NGS panel we could not detect a second mutation in MYO7A, but a splice acceptor site mutation (c.5068-2A > T) was found in CDH23. In the USH2 patient RP-1950, the USH2A mutation c.2299delG and a novel missense variant (p.Pro4760Ser) in GPR98 were detected. Bioinformatics algorithms predicted a pathogenic effect for these two novel variants (Table 4). In the former case, the two mutations were present in the same, clinically unaffected parent (data not shown), which does not support a digenic inheritance of USH. Unfortunately, segregation analysis could not be performed in the case of patient RP-1950. Perhaps a second mutation remains undetected in one of these genes in the patients RP-1426 and RP-1950. In two other patients (RP-1847 and RP-1923) with biallelic mutations in *USH*2A, we were able to detect additional mutations in USH1G and DFNB31, respectively. These cases show the high decisive power of massive parallel re-sequencing, detecting additional mutations in patients carrying the two mutations responsible for the disease in the same gene.

The programs used to analyze the data generated from NGS followed different algorithms. In fact, both the alignment and the base calling are two crucial steps in these analyses. It is possible that some mutations could escape detection due to the algorithms used. Their modification will improve the quality of generated data, avoiding false negatives and allowing detection of missed mutations in further analysis.

Four patients carried two mutations in genes typically responsible for another clinical subtype. Two USH2 patients carried missense mutations in *MYO7A* (RP-807 and RP-1864), one USH2 case in *CDH23* (RP-1781) and a USH3 patient carried a homozygous frameshift deletion in *USH2A* (RP-890) (Table 2). The situation of USH patients diagnosed with a given clinical subtype, who carry mutations in a gene involved in a different subtype, is not unprecedented, and was previously reported [25]. These findings further support the use of NGS techniques that allow for the study of all known causative genes for Usher syndrome in patients independent of their clinical subtype. It facilitates the detection of mutations in genes typically associated with other clinical types.

Copy number variation analysis

Studies performed in different populations have demonstrated that large rearrangements are an important cause of the disease in Usher syndrome. Le Guedard et al. [62] and Aller et al. [29] demonstrated that *PCDH15*

rearrangements are a significant cause of USH alterations. Large deletions involving *USH2A* have been reported [22,35], and large rearrangements in other USH genes as *MYO7A*, *CDH23* or *GPR98* have been detected [30,62]. These results indicate that the genetic analysis in USH patients should include the screening of rearrangements.

We have obtained with the MiSeq system a mean coverage in our USH panel of 1334x per sample and target region, allowing for CNV analysis of the selected genes. We could then detect in our series of patients six different rearrangements in *CDH23*, *PCDH15*, *USH2A* and *GPR98*, which represents 11.8% of the mutated detected alleles. We observed that when the coverage of regions fell below 250x, the results obtained in the CNV analysis were questionable and we did not take into account those regions. In order to perform quantitative analysis, we recommend, therefore, coverage of targeted regions above 250x.

In one patient (RP-531) the validation of the enrichment process gave an atypical image in the bioanalyzer (Figure 3). In this case the CNV analysis could not be performed. However, this does not affect the detection of point mutations. In fact, in another patient with a similar image in the validation (RP-808), two point mutations were detected.

Comparison with other NGS studies of USH

Traditionally, Sanger sequencing was used to identify the mutations responsible of the disease [25,50]. Nowadays different methods based on NGS have been used for Usher syndrome. Licastro et al. [36] employed two different approaches: the whole exome sequencing and long-PCR sequencing on nine USH genes. Whole exome sequencing has several problems: a low coverage of the interesting regions and the need of a correct interpretation of the high number of sequence variations identified. Regarding the long-PCR method, it displayed a better mean coverage, higher than 25x in 94% of regions. Taking together both approaches, the detection of pathogenic alleles was 62.5%.

Recently, Besnard et al. [37] developed a targeted NGS panel where 9 USH genes, 2 candidate USH genes (*VEZT* and *PDZD7*), seven hearing-loss genes and *CHM* were included. The overall depth of coverage obtained was estimated as 77x across the whole design ranging from 55.8x to 106.9x, with a detection ratio of 84.6% in the studied USH cohort.

In our series, the detection rate of mutated alleles was 79.7%. Our results are close to the Besnard et al. [37] study and higher than to the Licastro et al. [36] approaches. However, it is difficult to obtain conclusions due to the low number of patients included in those studies (thirteen and twelve USH patients respectively), whereas in our series 32 USH patients previously unscreened were sequenced.

Furthermore, in both previous studies, only point mutations were detected. The mean coverage obtained did not allow for the analysis of CNVs, that has been observed as an important USH cause. As has been pointed by Eisenberger et al.[63], high and extensive coverage allows for systematic analysis for CNVs and reduces the risk of mutations escaping detection because of their localization in regions with low coverage.

Conclusions

Our developed targeted NGS method based on HaloPlex gene target enrichment technology for the genetic diagnosis of Usher syndrome provides nearly complete coverage of all coding regions of the ten USH genes and four related and candidate USH genes. Furthermore, the high coverage obtained in our study allowed us to detect large rearrangements. It is important to detect both point mutations and large deletions or duplications with a single technique to minimize the economic cost of these studies, increasing the detection ratio of the genetic cause of the disease and improving the genetic diagnosis of Usher syndrome patients.

Availability of supporting data

LOVD http://www.lovd.nl/2.0/index_list.php

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

MJA performed the sequence capture, variant analysis and confirmation of point mutations by Sanger sequencing; EA contributed to the sequence capture, copy number variant analysis, MLPA and manuscript preparation; CFG contributed to variant analysis and confirmation of point mutations by Sanger sequencing; GGG performed a-CGH analysis; AFR contributed to a-CGH analysis and manuscript preparation, RR performed the functional in silico analysis of mutations; RPVM contributed to targeted next generation sequencing panel design and validation; FBK contributed to patients recruitment and DNA samples validation; CA contributed to patients recruitment and manuscript preparation; TJ designed the targeted next generation sequencing panel, supervised the experiments and redacted the manuscript JMM supervised the design and experiments and redacted the manuscript. All authors read and approved the final manuscript.

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References

- Espinos C, Millan JM, Beneyto M, Najera C: Epidemiology of Usher syndrome in Valencia and Spain. Community Genet 1998, 1:223–228.
- Keats BJ, Corey DP: The Usher syndromes. Am J Med Genet 1999, 89:158–166. Review.
- Cohen M, Bitner-Glindzicz M, Luxon L: The changing face of Usher syndrome: clinical implications. Int J Audiol 2007, 46:82–93.
- Millán JM, Aller E, Jaijo T, Blanco-Kelly F, Gimenez-Pardo A, Ayuso C: An update on the genetics of Usher syndrome. J Ophthalmol 2011, 2011:417217.
- Chaïb H, Kaplan J, Gerber S, Vincent C, Ayadi H, Slim R, Munnich A, Weissenbach J, Petit C: A newly identified locus for Usher syndrome type I, USH1E, maps to chromosome 21q21. Hum Mol Genet 1997, 6:27–31.
- Ahmed ZM, Riazuddin S, Khan SN, Friedman PL, Riazuddin S, Friedman TB: USH1H, a novel locus for type I Usher syndrome, maps to chromosome 15q22-23. Clin Genet 2009, 75:86–91.
- Riazuddin S, Belyantseva IA, Giese AP, Lee K, Indzhykulian AA, Nandamuri SP, Yousaf R, Sinha GP, Lee S, Terrell D, Hegde RS, Ali RA, Anwar S, Andrade-Elizondo PB, Sirmaci A, Parise LV, Basit S, Wali A, Ayub M, Ansar M, Ahmad W, Khan SN, Akram J, Tekin M, Riazuddin S, Cook T, Buschbeck EK, Frolenkov GI, Leal SM, Friedman TB, et al: Alterations of the CIB2 calcium- and integrinbinding protein cause Usher syndrome type 1 J and nonsyndromic deafness DFNB48. Nat Genet 2012, 44:1265–1271.
- Ebermann I, Phillips JB, Liebau MC, Koenekoop RK, Schermer B, Lopez I, Schäfer E, Roux AF, Dafinger C, Bernd A, Zrenner E, Claustres M, Blanco B, Nürnberg G, Nürnberg P, Ruland R, Westerfield M, Benzing T, Bolz HJ: PDZD7 is a modifier of retinal disease and a contributor to digenic Usher syndrome. J Clin Invest 2010, 120:1812–1823.
- Puffenberger EG, Jinks RN, Sougnez C, Cibulskis K, Willert RA, Achilly NP, Cassidy RP, Fiorentini CJ, Heiken KF, Lawrence JJ, Mahoney MH, Miller CJ, Nair DT, Politi KA, Worcester KN, Setton RA, Dipiazza R, Sherman EA, Eastman JT, Francklyn C, Robey-Bond S, Rider NL, Gabriel S, Morton DH, Strauss KA: Genetic mapping and exome sequencing identify variants associated with five novel diseases. PLoS One 2012, 7:e28936.
- Khateb S, Zelinger L, Mizrahi-Meissonnier L, Ayuso C, Koenekoop RK, Laxer U, Gross M, Banin E, Sharon D: A homozygous nonsense CEP250 mutation combined with a heterozygous nonsense C2orf71 mutation is associated with atypical Usher syndrome. J Med Genet 2014, Epub ahead of print.
- Kremer H, van Wijk E, Märker T, Wolfrum U, Roepman R: Usher syndrome: molecular links of pathogenesis, proteins and pathway. Hum Mol Genet 2006, 15. Spec No:R262-70.
- Michalski N, Michel V, Bahloul A, Lefèvre G, Barral J, Yagi H, Chardenoux S, Weil D, Martin P, Hardelin JP, Sato M, Petit C: Molecular characterization of the ankle-link complex in cochlear hair cells and its role in the hair bundle functioning. J Neurosci 2007, 13:6478–6488.
- Grati M, Shin JB, Weston MD, Green J, Bhat MA, Gillespie PG, Kachar B: Localization of PDZD7 to the stereocilia ankle-link associates this scaffolding protein with the Usher syndrome protein network. J Neurosci 2012, 32:14288–14293.
- Bahloul A, Simmler MC, Michel V, Leibovici M, Perfettini I, Roux I, Weil D, Nouaille S, Zuo J, Zadro C, Licastro D, Gasparini P, Avan P, Hardelin JP, Petit C: Vezatin, an integral membrane protein of adherens junctions, is required for the sound resilience of cochlear hair cells. EMBO Mol Med 2009, 1:155-138
- Kazmierczak P, Sakaguchi H, Tokita J, Wilson-Kubalek EM, Milligan RA, Müller U, Kachar B: Cadherin 23 and protocadherin 15 interact to form tip-link filaments in sensory hair cells. Nature 2007, 449:87–91.
- El-Amraoui A, Petit C: Cadherins as targets for genetic diseases. Cold Spring Harb Perspect Biol 2010, 2(1):a003095.
- Muller U: Cadherins and mechanotransduction by hair cells. Curr Opin Cell Biol 2008, 20:557–566.
- Sakaguchi H, Tokita J, Muller U, Kachar B: Tip links in hair cells: molecular composition and role in hearing loss. Curr Opin Otolaryngol Head Neck Surg 2009, 17:388–393.

- Wang A, Liang Y, Fridell RA, Probst FJ, Wilcox ER, Touchman JW, Morton CC, Morell RJ, Noben-Trauth K, Camper SA, Friedman TB: Association of unconventional myosin MYO15 mutations with human nonsyndromic deafness DFNB3. Science 1998, 29:1447–1451.
- Belyantseva IA, Boger ET, Naz S, Frolenkov GI, Sellers JR, Ahmed ZM, Griffith AJ, Friedman TB: Myosin-XVa is required for tip localization of whirlin and differential elongation of hair-cell stereocilia. Nat Cell Biol 2005, 7:148–156.
- Delprat B, Michel V, Goodyear R, Yamasaki Y, Michalski N, El-Amraoui A, Perfettini I, Legrain P, Richardson G, Hardelin JP, Petit C: Myosin XVa and whirlin, two deafness gene products required for hair bundle growth, are located at the stereocilia tips and interact directly. Hum Mol Genet 2005, 1:401–410.
- Maerker T, van Wijk E, Overlack N, Kersten FF, McGee J, Goldmann T, Sehn E, Roepman R, Walsh EJ, Kremer H, Wolfrum U: A novel Usher protein network at the periciliary reloading point between molecular transport machineries in vertebrate photoreceptor cells. Hum Mol Genet 2008, 17:71–86.
- Sahly I, Dufour E, Schietroma C, Michel V, Bahloul A, Perfettini I, Pepermans E, Estivalet A, Carette D, Aghaie A, Ebermann I, Lelli A, Iribarne M, Hardelin JP, Weil D, Sahel JA, El-Amraoui A, Petit C: Localization of Usher 1 proteins to the photoreceptor calyceal processes, which are absent from mice. J Cell Biol 2012, 199:381–399.
- 24. Cosgrove D, Zallocchi M: Usher protein functions in hair cells and photoreceptors. *Int J Biochem Cell Biol* 2014, **46**:80–89.
- 25. Bonnet C, Grati M, Marlin S, Levilliers J, Hardelin JP, Parodi M, Niasme-Grare M, Zelenika D, Délépine M, Feldmann D, Jonard L, El-Amraoui A, Weil D, Delobel B, Vincent C, Dollfus H, Eliot MM, David A, Calais C, Vigneron J, Montaut-Verient B, Bonneau D, Dubin J, Thauvin C, Duvillard A, Francannet C, Mom T, Lacombe D, Duriez F, Drouin-Garraud V, et al: Complete exon sequencing of all known Usher syndrome genes greatly improves molecular diagnosis. Orphanet J Rare Dis 2011, 6:21.
- Cremers FP, Kimberling WJ, Külm M, de Brouwer AP, van Wijk E, te Brinke H, Cremers CW, Hoefsloot LH, Banfi S, Simonelli F, Fleischhauer JC, Berger W, Kelley PM, Haralambous E, Bitner-Glindzicz M, Webster AR, Saihan Z, De Baere E, Leroy BP, Silvestri G, McKay GJ, Koenekoop RK, Millan JM, Rosenberg T, Joensuu T, Sankila EM, Weil D, Weston MD, Wissinger B, Kremer H: Development of a genotyping microarray for Usher syndrome. J Med Genet 2007, 44:153–160.
- Jaijo T, Aller E, García-García G, Aparisi MJ, Bernal S, Avila-Fernández A, Barragán I, Baiget M, Ayuso C, Antiñolo G, Díaz-Llopis M, Külm M, Beneyto M, Nájera C, Millán JM: Microarray-based mutation analysis of 183 Spanish families with Usher syndrome. *Invest Ophthalmol Vis Sci* 2010, 51:1311–1317.
- Vozzi D, Aaspöllu A, Áthanasakis E, Berto A, Fabretto A, Licastro D, Külm M, Testa F, Trevisi P, Vahter M, Ziviello C, Martini A, Simonelli F, Banfi S, Gasparini P: Molecular epidemiology of Usher syndrome in Italy. Mol Vis 2011. 17:1662–1668.
- Aller E, Jaijo T, García-García G, Aparisi MJ, Blesa D, Díaz-Llopis M, Ayuso C, Millán JM: Identification of large rearrangements of the PCDH15 gene by combined MLPA and a CGH: large duplications are responsible for Usher syndrome. *Invest Ophthalmol Vis Sci* 2010, 51:5480–5485.
- Roux AF, Faugère V, Vaché C, Baux D, Besnard T, Léonard S, Blanchet C, Hamel C, Mondain M, Gilbert-Dussardier B, Edery P, Lacombe D, Bonneau D, Holder-Espinasse M, Ambrosetti U, Journel H, David A, Lina-Granade G, Malcolm S, Claustres M: Four-year follow up of diagnostic service in USH1 patients. Invest Ophthalmol Vis Sci 2011, 52:4063–4071.
- 31. Steele-Stallard HB, Le Quesne SP, Lenassi E, Luxon LM, Claustres M, Roux AF, Webster AR, Bitner-Glindzicz M: Screening for duplications, deletions and a common intronic mutation detects 35% of second mutations in patients with USH2A monoallelic mutations on Sanger sequencing. Orphanet J Rare Dis 2013, 8:122.
- Choi BY, Park G, Gim J, Kim AR, Kim BJ, Kim HS, Park JH, Park T, Oh SH, Han KH, Park WY: Diagnostic application of targeted resequencing for familial nonsyndromic hearing loss. PLoS One 2013, 22:e68692.
- Mutai H, Suzuki N, Shimizu A, Torii C, Namba K, Morimoto N, Kudoh J, Kaga K, Kosaki K, Matsunaga T: Diverse spectrum of rare deafness genes underlies early-childhood hearing loss in Japanese patients: a crosssectional, multi-center next-generation sequencing study. Orphanet J Rare Dis 2013, 8:172.
- Fu Q, Wang F, Wang H, Xu F, Zaneveld JE, Ren H, Keser V, Lopez I, Tuan HF, Salvo JS, Wang X, Zhao L, Wang K, Li Y, Koenekoop RK, Chen R, Sui R: Next-generation sequencing-based molecular diagnosis of a Chinese patient cohort with autosomal recessive retinitis pigmentosa. *Invest Ophthalmol Vis Sci* 2013, 54:4158–4166.

- Glöckle N, Kohl S, Mohr J, Scheurenbrand T, Sprecher A, Weisschuh N, Bernd A, Rudolph G, Schubach M, Poloschek C, Zrenner E, Biskup S, Berger W, Wissinger B, Neidhardt J: Panel-based next generation sequencing as a reliable and efficient technique to detect mutations in unselected patients with retinal dystrophies. Eur J Hum Genet 2014, 22:99–104.
- Licastro D, Mutarelli M, Peluso I, Neveling K, Wieskamp N, Rispoli R, Vozzi D, Athanasakis E, D'Eustacchio A, Pizzo M, D'Amico F, Ziviello C, Simonelli F, Fabretto A, Scheffer H, Gasparini P, Banfi S, Nigro V: Molecular diagnosis of Usher syndrome: application of two different next generation sequencing-based procedures. PLoS One 2012, 7:e43799.
- Besnard T, García-García G, Baux D, Vaché C, Faugère V, Larrieu L, Léonard S, Millan JM, Malcolm S, Claustres M, Roux AF: Experience of targeted Usher exome sequencing as a clinical test. Mol Genet Genomic Med 2014, 2:30–43.
- Yoshimura H, Iwasaki S, Nishio SY, Kumakawa K, Tono T, Kobayashi Y, Sato H, Nagai K, Ishikawa K, Ikezono T, Naito Y, Fukushima K, Oshikawa C, Kimitsuki T, Nakanishi H, Usami S: Massively parallel DNA sequencing facilitates diagnosis of patients with Usher syndrome type 1. PLoS One 2014. 9:e90688
- Rong W, Chen X, Zhao K, Liu Y, Liu X, Ha S, Liu W, Kang X, Sheng X, Zhao C: Novel and recurrent MYO7A mutations in Usher syndrome type 1 and type 2. PLoS One 2014, 9(5):e97808. doi:10.1371/journal.pone.0097808. eCollection 2014.
- von Brederlow B, Bolz H, Janecke A, La O, Cabrera A, Rudolph G, Lorenz B, Schwinger E, Gal A: Identification and in vitro expression of novel CDH23 mutations of patients with Usher syndrome type 1D. Hum Mutat 2002, 19:268–273.
- Garcia-Garcia G, Aparisi MJ, Jaijo T, Rodrigo R, Leon AM, Avila-Fernandez A, Blanco-Kelly F, Bernal S, Navarro R, Diaz-Llopis M, Baiget M, Ayuso C, Millan JM, Aller E: Mutational screening of the USH2A gene in Spanish USH patients reveals 23 novel pathogenic mutations. Orphanet J Rare Dis 2011. 6:65.
- Schultz JM, Bhatti R, Madeo AC, Turriff A, Muskett JA, Zalewski CK, King KA, Ahmed ZM, Riazuddin S, Ahmad N, Hussain Z, Qasim M, Kahn SN, Meltzer MR, Liu XZ, Munisamy M, Ghosh M, Rehm HL, Tsilou ET, Griffith AJ, Zein WM, Brewer CC, Riazuddin S, Friedman TB: Allelic hierarchy of CDH23 mutations causing non-syndromic deafness DFNB12 or Usher syndrome USH1D in compound heterozygotes. J Med Genet 2011, 48:767–775.
- Roux AF, Faugère V, Le Guédard S, Pallares-Ruiz N, Vielle A, Chambert S, Marlin S, Hamel C, Gilbert B, Malcolm S, Claustres M, French Usher Syndrome Collaboration: Survey of the frequency of USH1 gene mutations in a cohort of Usher patients shows the importance of cadherin 23 and protocadherin 15 genes and establishes a detection rate of above 90%. J Med Genet 2006, 43:763–768.
- Ahmed ZM, Riazuddin S, Bernstein SL, Ahmed Z, Khan S, Griffith AJ, Morell RJ, Friedman TB, Riazuddin S, Wilcox ER: Mutations of the protocadherin gene PCDH15 cause Usher syndrome type 1 F. Am J Hum Genet 2001, 69(1):25–34.
- Jaijo T, Oshima A, Aller E, Carney C, Usami S, Millán JM, Kimberling WJ: Mutation screening of the PCDH15 gene in Spanish patients with Usher syndrome type I. Mol Vis 2012, 18:1719–1726.
- Riazuddin S, Nazli S, Ahmed ZM, Yang Y, Zulfiqar F, Shaikh RS, Zafar AU, Khan SN, Sabar F, Javid FT, Wilcox ER, Tsilou E, Boger ET, Sellers JR, Belyantseva IA, Riazuddin S, Friedman TB: Mutation spectrum of MYO7A and evaluation of a novel nonsyndromic deafness DFNB2 allele with residual function. Hum Mutat 2008, 29:502–511.
- Liu XZ, Hope C, Liang CY, Zou JM, Xu LR, Cole T, Mueller RF, Bundey S, Nance W, Steel KP, Brown SD: A mutation (2314delG) in the Usher syndrome type IIA gene: high prevalence and phenotypic variation. Am J Hum Genet 1999, 64(4):1221–1225.
- Bharadwaj AK, Kasztejna JP, Huq S, Berson EL, Dryja TP: Evaluation of the myosin VIIA gene and visual function in patients with Usher syndrome type I. Exp Eye Res 2000, 71:173–181.
- Aller E, Jaijo T, Beneyto M, Nájera C, Oltra S, Ayuso C, Baiget M, Carballo M, Antiñolo G, Valverde D, Moreno F, Vilela C, Collado D, Pérez-Garrigues H, Navea A, Millán JM: Identification of 14 novel mutations in the long isoform of USH2A in Spanish patients with Usher syndrome type II. J Med Genet 2006, 43:e55.
- Le Quesne SP, Saihan Z, Rangesh N, Steele-Stallard HB, Ambrose J, Coffey A, Emmerson J, Haralambous E, Hughes Y, Steel KP, Luxon LM, Webster AR, Bitner-Glindzicz M: Comprehensive sequence analysis of nine Usher

- syndrome genes in the UK National Collaborative Usher Study. *J Med Genet* 2012, **49**(1):27–36.
- Ouyang XM, Yan D, Du LL, Hejtmancik JF, Jacobson SG, Nance WE, Li AR, Angeli S, Kaiser M, Newton V, Brown SD, Balkany T, Liu XZ: Characterization of Usher syndrome type I gene mutations in an Usher syndrome patient population. *Hum Genet* 2005, 116(4):292–299.
- Besnard T, Vaché C, Baux D, Larrieu L, Abadie C, Blanchet C, Odent S, Blanchet P, Calvas P, Hamel C, Dollfus H, Lina-Granade G, Lespinasse J, David A, Isidor B, Morin G, Malcolm S, Tuffery-Giraud S, Claustres M, Roux AF: Non-USH2A mutations in USH2 patients. Hum Mutat 2012, 33:504–510.
- Oshima A, Jaijo T, Aller E, Millan JM, Carney C, Usami S, Moller C, Kimberling WJ: Mutation profile of the CDH23 gene in 56 probands with Usher syndrome type I. Hum Mutat 2008, 29(6):E37–46.
- Janecke AR, Meins M, Sadeghi M, Grundmann K, Apfelstedt-Sylla E, Zrenner E, Rosenberg T, Gal A: Twelve novel myosin VIIA mutations in 34 patients with Usher syndrome type I: confirmation of genetic heterogeneity. Hum Mutat 1999. 13:133–140.
- Bernal S, Medà C, Solans T, Ayuso C, Garcia-Sandoval B, Valverde D, Del Rio E, Baiget M: Clinical and genetic studies in Spanish patients with Usher syndrome type II: description of new mutations and evidence for a lack of genotype—phenotype correlation. Clin Genet 2005, 68:204–214.
- Jacobson SG, Aleman TS, Sumaroka A, Cideciyan AV, Roman AJ, Windsor EA, Schwartz SB, Rehm HL, Kimberling WJ: Disease boundaries in the retina of patients with Usher syndrome caused by MYO7A gene mutations. Invest Ophthalmol Vis Sci 2009, 50:1886–1894.
- Kaiserman N, Obolensky A, Banin E, Sharon D: Novel USH2A mutations in Israeli patients with retinitis pigmentosa and Usher syndrome type 2. Arch Ophthalmol 2007, 125:219–224.
- Vaché C, Besnard T, le Berre P, García-García G, Baux D, Larrieu L, Abadie C, Blanchet C, Bolz HJ, Millan J, Hamel C, Malcolm S, Claustres M, Roux AF: Usher syndrome type 2 caused by activation of an USH2A pseudoexon: implications for diagnosis and therapy. Hum Mutat 2012, 33:104–108.
- Ouyang XM, Xia XJ, Verpy E, Du LL, Pandya A, Petit C, Balkany T, Nance WE, Liu XZ: Mutations in the alternatively spliced exons of USH1C cause non syndromic recessive deafness. Hum Genet 2002, 111(1):26–30.
- Jaijo T, Aller E, Beneyto M, Najera C, Graziano C, Turchetti D, Seri M, Ayuso C, Baiget M, Moreno F, Morera C, Perez-Garrigues H, Millan JM: MYO7A mutation screening in Usher syndrome type I patients from diverse origins. J Med Genet 2007, 44(3):e71.
- Ross M, Russ C, Costello M, Hollinger A, Lennon NJ, Hegarty R, Nusbaum C, Jaffe DB: Characterizing and measuring bias in sequence data. *Genome Biol* 2013. 14:R51.
- Le Guédard S, Faugère V, Malcolm S, Claustres M, Roux AF: Large genomicrearrangements within the PCDH15 gene are a significant cause of USH1F syndrome. Mol Vis 2007, 13:102–107.
- 63. Eisenberger T, Neuhaus C, Khan AO, Decker C, Preising MN, Friedburg C, Bieg A, Gliem M, Charbel Issa P, Holz FG, Baig SM, Hellenbroich Y, Galvez A, Platzer K, Wollnik B, Laddach N, Ghaffari SR, Rafati M, Botzenhart E, Tinschert S, Börger D, Bohring A, Schreml J, Körtge-Jung S, Schell-Apacik C, Bakur K, Al-Aama JY, Neuhann T, Herkenrath P, Nürnberg G, et al: Increasing the yield in targeted next-generation sequencing by implicating CNV analysis, non-coding exons and the overall variant load: the example of retinal dystrophies. PLoS One 2013, 8:e78496.

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