

ARTICLE

# USH3A transcripts encode clarin-1, a four-transmembrane-domain protein with a possible role in sensory synapses

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Usher syndrome type 3 (USH3) is an autosomal recessive disorder characterised by the association of post-lingual progressive hearing loss, progressive visual loss due to retinitis pigmentosa and variable presence of vestibular dysfunction. Because the previously defined transcripts do not account for all USH3 cases, we performed further analysis and revealed the presence of additional exons embedded in longer human and mouse *USH3A* transcripts and three novel *USH3A* mutations. Expression of *Ush3a* transcripts was localised by whole mount *in situ* hybridisation to cochlear hair cells and spiral ganglion cells. The full length *USH3A* transcript encodes clarin-1, a four-transmembrane-domain protein, which defines a novel vertebrate-specific family of three paralogues. Limited sequence homology to stargazin, a cerebellar synapse four-transmembrane-domain protein, suggests a role for clarin-1 in hair cell and photoreceptor cell synapses, as well as a common pathophysiological pathway for different Usher syndromes.

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## Introduction

Combined deafness and blindness in adults is most frequently caused by Usher syndromes (USH). These account for more than half of the dual sensory deficit, with a prevalence of 1/10000 in the age group of 30–50.<sup>1</sup> Sensorineural hearing loss and retinitis pigmentosa (RP) characterise this group of autosomal recessive hereditary disorders.

Three USH types are distinguished clinically. Postlingual progressive hearing loss and variable vestibular function characterise USH3 (MIM 276902).<sup>2</sup> Progressive RP with variable age of onset occurs in all USH types. While USH1 and USH2 map to at least 10 distinct loci,<sup>3–11</sup> only one locus for USH3 has been reported so far.<sup>12,13</sup> Four of the *USH1* and the *USH2A* genes have been identified. Mutations in the unconventional myosin *MYO7A* cause USH1B (MIM 276903<sup>14</sup>) and in two unique cases also atypical USH phenotype similar to that of USH3.<sup>15</sup> Mutations in the PDZ-domain-containing protein, harmonin, cause

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USH1C (MIM 276904)<sup>16,17</sup>. PDZ-domains are protein–protein interaction modules that allow the binding to and clustering of specific membrane-associated proteins, such as receptors and ion channels.<sup>18</sup> Cadherin23 and protocadherin15 mutations have recently been shown to underlie USH1D (MIM 601067) and USH1F (MIM 602083), respectively.<sup>19–22</sup> Most cadherins are integral membrane glycoproteins that mediate the calcium-dependent formation of cell–cell adhesion,<sup>23</sup> while protocadherins are thought to be involved in a variety of functions, including neural development, neural circuit formation and formation of the synapse.<sup>24</sup> Finally, mutations in Usherin, a gene that encodes a basement membrane protein, cause USH2A (MIM276901)<sup>25</sup>.

Joensuu *et al.*<sup>26</sup> have recently identified three mutations (a missense, a nonsense and a 3 bp deletion) in exons 2 and 3 of a newly identified gene that maps to 3q24, approximately 100 kb proximal to the previously defined USH3A linkage interval in 3q25.1. These mutations established this gene as the *USH3A* gene. Northern blot analysis and reverse-transcription PCR indicated the expression of the *USH3A* gene in several tissues, including retina. Two predicted transmembrane domains were identified in its deduced protein product, which at the time did not show similarity to any known protein.<sup>26</sup>

In this study, we characterise new human and mouse *USH3A* transcripts (AF495717; AF495718; AF495719 and AF495720), identify three additional USH3 mutations, analyse the *USH3A* expression pattern and by *in situ* hybridisation localise transcripts of this gene to mouse cochlear hair cells and spiral ganglion cells. We redefine the *USH3A* protein, name it clarin-1 and affiliate it to a new four transmembrane domain (4TM) vertebrate-specific protein family. Furthermore, based on sequence similarities to stargazin, a well-studied member of this hyperfamily, we suggest a role for clarin-1 in the hair cells synaptic junctions.

## Materials and Methods

### USH3 families and controls

Blood samples were drawn by venipuncture after obtaining informed consent in accordance with the guidelines of the Tel Aviv University Helsinki Committee. Three of the Jewish Ashkenazi USH3 families included in this study, were referred to us through the Center for Deaf–Blind Persons in Tel Aviv as part of a larger study on the genetics of Usher syndromes in Israel. The fourth Jewish Ashkenazi USH3 family living in the US has one affected and two unaffected siblings whose grandparents originate from Eastern Europe. The family was ascertained by one of us (JGF). The USH3 Jewish Yemenite and the Spanish families have been described elsewhere.<sup>27,28</sup> Clinical diagnosis of affected members in these families is compatible with the USH3 phenotype and haplotype segregation analysis does not exclude linkage to the USH3A locus. Control DNA samples from unrelated Jewish Ashkenazi and Yemenite individuals,

were provided by the National Laboratory for the Genetics of Israeli Populations. A total of 51 grandparents in families from the Centre d’Etude du Polymorphisme Humain (CEPH) were also studied as controls.

### RNA expression

The expression of *Ush3a* was determined by *in situ* hybridisation and reverse-transcription (RT)–PCR in mice at embryonic day (E) 16, postnatal day (P)0, 5, 10, 15, 20, and 30. For RT–PCR, cochleae were dissected out of the temporal bones. Total RNA was prepared from cochleae with TRI Reagent (Sigma). Genomic DNA was removed from all RNA samples using DNA-free (Ambion). Total RNA was purified using phenol-chloroform extraction (PCI; Gibco) and phase lock gel tubes (Eppendorf), followed by an isopropanol precipitation. RT reactions were performed using Expand Reverse Transcriptase (Roche) with Homo-Oligomeric DNA d(T)12–18 and Random Hexamer (Amersham Pharmacia Biotech). cDNA was amplified from the RT reactions using primers (F) 5′-GGTCCAAGC-CATCCCCGTA-3′ AND (R) 5′- CCTCCTGCTTCTGTTATTTT-CC-3′. As a control, primers spanning the last intron of *Myo6* (F) 5′-CTGGTGGTATGCCCATTTTGA-3′ and (R) 5′-TCGCTTTCATAAGGCATTCTA-3′ were used.<sup>29</sup>

For *in situ* hybridisation, cochleae were dissected from surrounding tissue and fixation was performed by immersion in 4% paraformaldehyde in PBS for 12–16 h at 4°C. The samples were processed without decalcification until P10. From P10 onwards, the samples were dehydrated in graded concentrations of methanol, from 25–100%, followed by decalcification (to facilitate dissection) in 8% formic acid in 100% methanol for up to 4 days. We generated digoxigenin-labelled cRNA probes in a standard transcription reaction (Roche Molecular Biochemicals). We performed whole mount *in situ* hybridisation as described,<sup>30</sup> with modifications. Finally, we visualised *in situ* reaction product by cryoprotecting whole cochleae with sucrose, embedding the tissue in OCT, and cryostat sectioning (10 μm). Antisense and sense probes were *in vitro* transcribed from a linearised vector, pPCR-Script AMP SK(+) cloning vector (Stratagene), containing a partial mouse *Ush3a* cDNA (corresponding to exons 2 and 3) by using either T3 or T7 RNA polymerase. No *in situ* hybridisation signal was detected with the control sense RNA probe. Our use of animals was approved by the Tel Aviv University Animal Care and Use Committee (11-00-65).

### Mutation detection and analysis

Using the primers detailed in Table 2 we amplified fragments encompassing *USH3A* exons from genomic DNA templates of USH3 patients and their family members. PCR products were gel-purified, sequenced using dye terminators of Big-Dyes kits (Perkin-Elmer/Applied Biosystems) and analysed on an ABI 3700 sequencer (Perkin-Elmer/

**Table 1** Mutation and polymorphism in the *USH3A* gene

Exon/Intron	Base change	Predicted AA change	Frequency	Origin & No. of families
<i>Disease causing mutations</i>				
Ex0	143T>G	N48K Asn>Lys (aat>aag)	1/221	Jew. Hungarian (1), Jew. Russian (3)
Ex0	189C>A	Y63X	0/135	Spanish (1)
Ex0	187-209del23bp	Frameshift at aa 63 and stop codon after 25 aa	0/199	Jew. Yemenite (1)
<i>Polymorphism</i>				
Ex0	-71A>G	UTR	10/33	Ashkenazi Jew. (7: Russian, Polish, Hungarian, Rumanian), Jew. Moroccan (1), Jew. Yemenite (2)
Ex0	+55A>T	Ala19Ala (gca>gct)	5/33	Jew. Moroccan (3), Jew. Russian (1), Jew. Yemenite (1)
IVS1	+112a>g	Silent	5/18	Jew. Hungarian (1), Jew. Moroccan (1), Jew. Russian (1), Jew. Yemenite (2)
IVS1	+135a>t	Silent	2/18	Jew. Moroccan (1), Jew. Yemenite (1)
Ex3(IVS3a)	963-1002 ATnGTn	UTR	5/14	Jew. Hungarian (1)*, Jew. Russian (2)*, Jew. Yemenite (1)*
Ex3(IVS3a)	1013T>C	UTR	5/14	Jew. Hungarian (1)*, Jew. Russian (2)*, Jew. Yemenite (1)*

Codon and nucleotide numbering starts from the codon of the first in-frame methionine of the *USH3A* ORF. The frequency column presents the number of chromosomes carrying the sequence change divided by the number of all tested chromosomes. All *USH3A* exons from individuals with *USH3* and other family members, were amplified from leukocyte genomic DNA and screened for mutations by direct PCR sequencing. An ATnGTn microsatellite at nucleotide positions 963–1002 that was identified in the 3' UTR of *USH3A* exon 3 can be used as a polymorphic genetic marker to test chromosomal segregation and linkage to the *USH3* locus. \*Carried on N48K chromosome.

Applied Biosystems). Sequence comparisons were performed by the use of Sequencher 4.1 software from GeneCodes Corporation. The Jewish Ashkenazi N48K mutation was screened by *StuI* digestion (16 h at 37°C) followed by agarose (2%) gel analysis. The 23 bp deletion (found in patients of Yemenite origin) was screened by PCR amplification of a 227 bp fragment followed by FMC's MetaPhor gel (4%) electrophoresis. The Y63X mutation (found in a patient of Spanish origin) was screened by SSCP; a 207-bp fragment of exon 0 was PCR amplified by primers ex0F2 (5-ATCAAAGCCACTGTCCTCTG-3) and ex0R (5-CTGGGAA-GAGTCTGCCTAAA-3), and the alleles were separated on 0.7 × MDE gels, for 18 h at 5 W, at room temperature and were visualised by silver staining.<sup>31</sup>

#### Computational analyses and sequence annotation

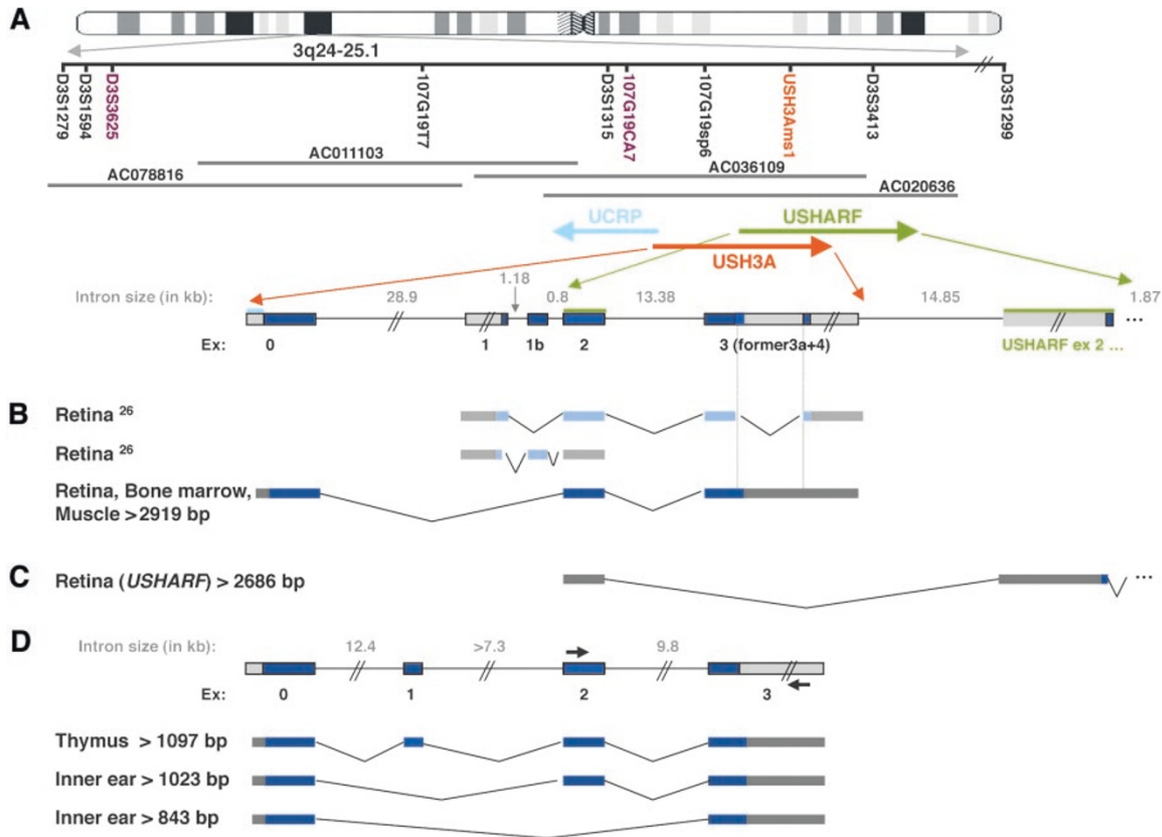
Genomic sequences were retrieved from four NCBI (National Center for Biotechnology Information) Human clones: AC020636, AC036109, AC011103, AC078816. Fragments were assembled by Sequencher 4.1 software from GeneCodes Corporation. Complete sequence annotation was performed using the GESTALT workbench.<sup>32</sup> Sequence similarity searches performed by the use of the BLAST<sup>33</sup> and Blimps<sup>34</sup> programs. Sequences were aligned using the BlockMaker<sup>34</sup> and MACAW<sup>35</sup> programs; PHYLIP ProtDist<sup>36</sup> PHYLIP – Phylogeny Inference Package (Version 3.2 Cladistics 5: 164–166) and CLUSTAL W programs<sup>37</sup> were used to compute trees and their significance (bootstrap) values. TM regions were predicted using the PHDhtm<sup>38</sup> and TMHMM programs.<sup>39</sup> Primers were designed using the Oligo primer analysis software (<http://medprobe.com/is/oligo/html>).

## Results

### Genomic analysis and transcripts

Since no mutations were identified within the described *USH3A* transcripts<sup>26</sup> in patients from other *USH3* families, we assumed that there might be additional uncharacterised exons of this gene. Therefore, we assembled and reconstructed a 500 kb genomic interval spanning the published *USH3A* gene and the previous *USH3A* linkage interval, defined by *D3S3413* and *D3S1279* (Figure 1A). Based on gene predictions and alignment with human and mouse ESTs, we designed primers that allowed the amplification of a new human *USH3A* transcript. This longer transcript begins with a newly identified exon 0, continues with exon 2, has exons 3 and 4 transcribed together with their intervening intron, but does not include the previously identified exons 1 and 1b<sup>26</sup> (Figure 1B).

The *USH3A* gene shows partial exon overlap (but no protein overlap) with the 5' end of the *UCRP* pseudogene<sup>26</sup> in an opposite polarity, and in the same polarity with the 5' end of a newly characterised gene, *USHARF* (*USH3A* Alternative Reading Frame, ms in preparation) (Figure 1C). By EST assembly and PCR amplification from mouse inner ear cDNA library we also defined three alternative transcripts of the *Ush3A* orthologue (Figure 1D). One of these transcripts (AF495719) has the same exon-intron structure as the longest human mRNA, and its protein product shares 88% identity with the product of the corresponding human transcript (AF495717). A longer mouse transcript (AF495718) has an extra exon coding for additional 18 amino acids (Figure 3C).



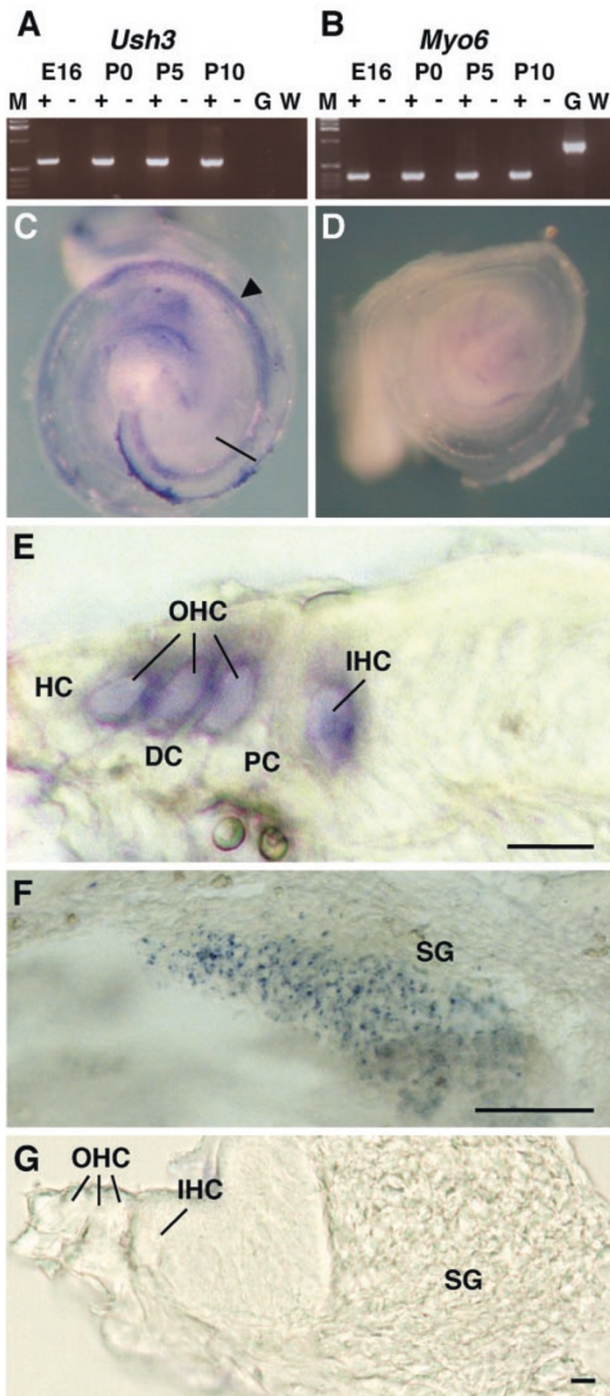
**Figure 1** (A) Schematic representation of physical and transcript maps covering the *USH3A* region comprised of partly overlapping NCBI (National Center for Biotechnology Information) clones. Polymorphic markers and sequence-tagged sites are indicated on the bar below the chromosome. Pink markers define the borders of the old *USH3A* linkage interval. The red marker is the newly identified polymorphic intragenic microsatellite located within the *USH3A*'s 3'-UTR. Colored arrows below the clones indicate the position and orientation of the *UCRP* pseudogene (light blue), the *USH3A* (red) and partially overlapping *USHARF* (green) genes. Below these arrows is an enlarged view of an approximate 66 kb segment, showing the genomic structure of the *USH3A* gene and the first two exons of the *USHARF* gene. Blue boxes and lines indicate coding exons, while grey boxes and lines indicate non-coding regions. Light blue and green lines above the exons indicate overlapping regions between *USH3A* and, respectively, *UCRP* and *USHARF*. (B) Alternative transcripts of the *USH3A* gene. The first two pale colored transcripts were defined previously by Joensuu *et al.* (2001).<sup>26</sup> (C) The first two exons of the *USHARF* gene. (D) Genomic structure of the mouse *Ush3a* gene and its three newly defined in frame alternative transcripts. Marked by black arrows is the position of the primers that were used for the amplification of the (RT)-PCR and the *in situ* hybridisation probe. Previously known *USH3A* human and mouse ESTs and mRNAs, amplified from several different tissues are listed in Table 3.

### Expression patterns

By PCR amplification from cDNA or cDNA libraries we confirmed the previously reported expression of *USH3A* in retina and skeletal muscle. In addition, we have amplified *USH3A* transcripts from human testis and olfactory epithelium cDNA. The later observation might be related to previously reported expression of two other Usher genes, myosin VIIA and harmonin, in the olfactory tissue.<sup>40–43</sup> Yet, despite the observation of anomalies in olfactory cilia, no chemosensory deficits have been confirmed in Usher patients.<sup>44</sup>

By PCR amplification of mouse inner ear cDNA and by whole mount *in situ* hybridisation on mouse cochleae we

also demonstrate the expression of *Ush3a* transcripts in the auditory sensory organ. A developmental expression profile of *Ush3a* was performed both by PCR amplification of mouse inner ear cDNA and whole mount *in situ* hybridisation on mouse cochleae. *Ush3a* transcripts were detected as early as E16, the earliest age tested, and at all postnatal stages examined (P0, P5, and P10) (Figure 2A,B). Hybridisation with an *Ush3a* RNA probe on whole cochleae detected mRNA expression in the sensory epithelium at E16 (Figure 2C,D), P0, P5, P10, P15, P20 ( $n=4$  for each age) (data not shown). Sectioning through these cochleae revealed specific hybridisation in the inner and outer hair cells of the organ of Corti (Figure 2E,G). No hybridisation was detected in the



**Figure 2** Expression of *Ush3a* in mouse cochlea by reverse transcriptase (RT)–PCR and *in situ* hybridisation. (A) Expression of *Ush3a* in mouse cochlea by RT–PCR at E16 and P0, P5, and P10. RNA derived from cochlear ducts was used to amplify a 605 bp fragment spanning exons 2–4 of the *Ush3a* gene by RT–PCR. Amplifications were carried out with (+) and without (–) RT on cochlear RNA, on mouse genomic DNA (G), and a water control (W). (B) A known cochlear-expressing gene, myosin VI (*Myo6*),<sup>64</sup> amplified with primers spanning an intron from the same RNA preparation. (C–G) *In situ* hybridisation on

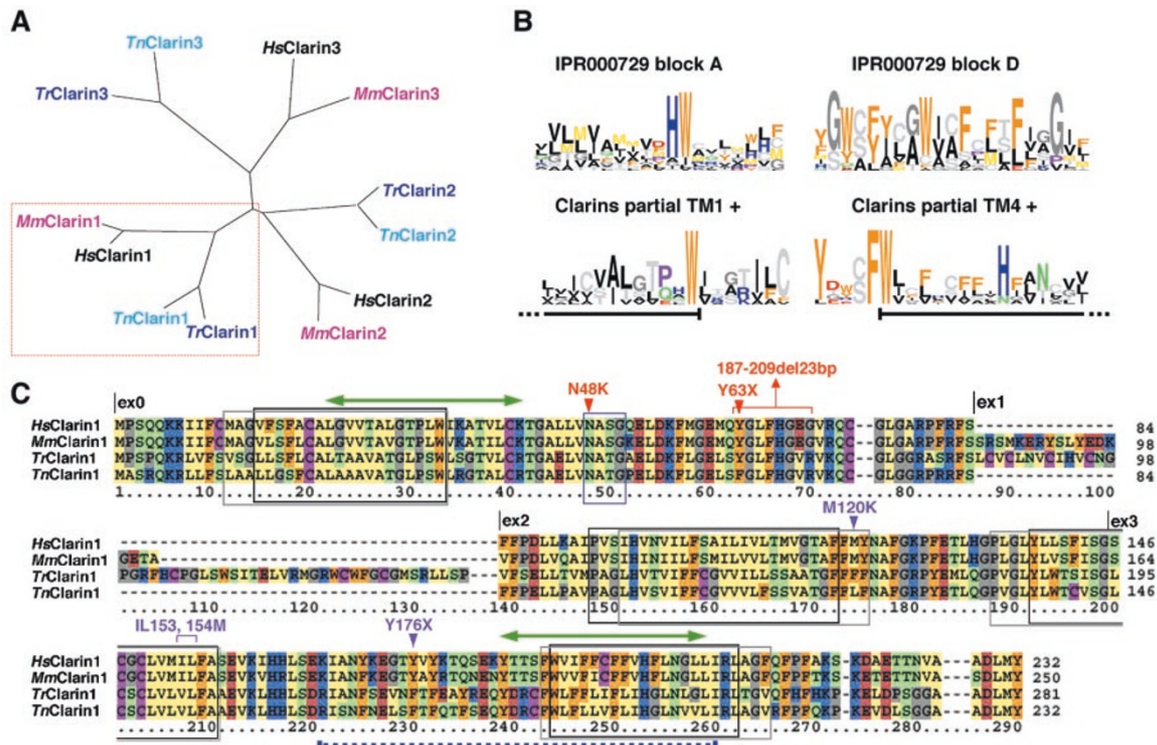
supporting cells of the organ of Corti, including the Deiters' cells, the pillar cells, and the Hensen cells. The only other hybridisation found in the cochlea was in the spiral ganglion cells (Figure 2F) containing the primary neurons that innervate the cochlear sensory epithelia.

#### Gene product and protein structure

The open reading frame of the newly defined *USH3A* transcript is expected to encode a 232 amino acid protein with four transmembrane domains, which we named clarin-1, after the clarity of sensory perception allowed by the intact protein (Figure 3C). Sequence database searches by BLAST<sup>33</sup> identified two additional clarin-1 human paralogues, clarin-2 and -3 (Figure 3A). The clarins were found to have respective orthologues in mouse as well as in fish (*Takifugu rubripes* [Fugu] and *Tetraodon nigroviridis*), all encoding small 224–284 amino acid proteins (Figure 3A). A partial clarin-1 mRNA was also identified in chicken. No orthologues were identified in the genomes of prokaryotes, yeast, plants, nematodes and insects. Four transmembrane-domains (TMs), conserved sequence motifs and a single glycosylation consensus site between TM1 and TM2 characterise the clarin family members (Figure 3C).

In a wider context the clarins appear to belong to a large hyperfamily of small integral membrane glycoproteins with four transmembrane domains. These include the tetraspannins<sup>45</sup> (InterPro families PR00218 and PS00930) as well as the PMP22/EMP/MP20 and claudin family (InterPro family IPR000729) that perform diverse membrane transport, transduction and cell–cell interaction and scaffolding functions. Yet, BLAST searches did not provide a sequence similarity basis for a relationship between clarin-1 and any members of this hyperfamily. To examine this question

mouse cochlea was done using digoxigenin-labelled sense and antisense riboprobes for *Ush3a*, followed by cryosectioning to confirm specific cell patterns. Expression was shown in the hair cells of the cochlea and in spiral ganglion cells at all ages tested. (C) Whole mount of mouse cochlea at E16 reveals specific labelling of sensory epithelium by the *Ush3a* antisense probe (arrow). Bar indicates area of cross-section for (E). (D) Absence of labelling with the *Ush3a* sense probe in a whole mount cochlea. (E) Cross section of cochlea in C showing a portion of the organ of Corti and specific labelling of the *Ush3a* probe only in the inner (IHC) and outer hair cells (OHC). Hybridisation is absent in the supporting cells, including the pillar cells (PC), Deiters' cells (DC), and Hensen's cells (HC). Scale bar, 10  $\mu$ m. Left corresponds to the peripheral end of the section bar in C. (F) The region containing the spiral ganglion (SG) cells is shown. No staining was seen in any other regions of the cochlea upon sectioning. Scale bar, 50  $\mu$ m. (G) Cross section of cochlea showing hair cells, supporting cells, and spiral ganglion in D demonstrating lack of hybridisation with sense probe. Scale bar, 10  $\mu$ m. All experiments with mice were carried out with the approval of the Tel Aviv University Animal Care and Use Committee (11-00-65).



**Figure 3** (A) Unrooted tree reflecting sequence relations between protein products of 12 of the 14 identified clarins. Sequence similarity searches revealed 14 members of the 4TM *USH3A* gene family, 12 of which, with the most completed sequences, are presented in this tree. These include *HsClarin-1* (*USH3A*) and its two predicted paralogues *HsClarin-2* (LOC166908 product of XM\_068256) and *HsClarin-3* (LOC119467 product of XM\_058398), that map to 4p15.33 and 10q26.2 respectively; *MmClarin-1* (*Ush3a*) and its two predicted paralogues *MmClarin-2*, deduced from TblastN results against Celera's mouse genomic assembly, and *MmClarin-3* (product of Celera's annotated transcript mCP9472) that map to mouse chromosomes 5 and 7 respectively; three predicted *Fugu* gene products, *TrClarin-1* (JGI\_31106), *TrClarin-2* (JGI\_10159) and *TrClarin-3* (JGI\_29906) that was corrected according to TblastN and Block analysis results; Three predicted *Tetraodon* genes that were also reconstructed from TblastN results against Genoscope *Tetraodon* genomic traces and named *TnClarin1*, *TnClarin-2* and *TnClarin-3*. Prediction of a partial mRNA of a bovine gene (BF044503) were also included in the analyses of the *USH3* gene family but are not shown in the figure. Block multiple alignment analysis<sup>34</sup> of the clarins identified several conserved sequence motifs covering about 95% of the sequence of clarin-1 and 80% or more of the sequences of other family members. Both PHYMLIP and CLUSTAL W, used to compute trees and their significance (bootstrap) values, programs gave identical topologies. Included branch points have 70% and higher bootstrap values. (B) Sequence logo of the two regions (indicated in C by green arrows) that are similar between the clarins and members of the IPR000729 protein family. In the logo, the height of each amino acid is scaled in bits of information and is proportional to its degree of conservation.<sup>34</sup> Black lines under the logos indicate TM prediction. (C) Alignment of clarin-1 and its three orthologues including *MmClarin1*, *TrClarin1* (JGI\_31106) and *TnClarin1*, which are boxed in red on the phylogenetic tree (A). Newly identified and previously described<sup>26</sup> *USH3A* mutations are indicated, respectively, by red and purple arrows. The starting position of each exon is indicated above the sequence. Black and grey boxes indicate TM positions according to, respectively, PHDhtm analysis of the four aligned proteins and TMHMM analysis of clarin-1. Boxed in blue is the putative N- glycosylation site. Green arrows indicate the positions of regions, where clarins share similarity with members of the IPR000729 protein family. Human and mouse clarin-1 harbour in their C-terminal region a TNV signature that might serve as a PDZ-binding motif. However this signature is not conserved among other members of the clarins. Gaps in the aligned sequence originate from the full alignment of all 12 clarins presented in the tree (A).

further, it was necessary to employ more sensitive search routines, capable of detecting subtle sequence similarities. First, we asked whether any of the clarins contained protein motifs already included in the BLOCKS database, but the results were negative. Therefore, using the newly discovered group of 12 clarin sequences, we defined novel clarin-specific Blocks motifs. These were subsequently analysed by the Local Alignment of Multiple Alignments (LAMA) method,<sup>46</sup>

looking for remote but statistically significant similarities between the clarin Blocks and all others. Two of the clarin Blocks, in and around the first and fourth putative TM domains generated a hit with two of the 4TM proteins Blocks (Figure 3B,C). Pairwise Smith-Waterman alignments<sup>47</sup> then allowed us to delineate the broad similarity relations between the clarins and representative members of several human 4TM families (Figure 5).

In order to find out which of the 4TM proteins is most highly related to the clarins, we performed further pairwise alignments with accurate statistical testing (Figure 6). This highlighted the calcium channel gamma subunit proteins (CACNG) as best clarin sequence matches. Of these CACNG2 (stargazin)<sup>48,49</sup> was the only protein showing significant matches ( $P < 0.001$ ) to all three clarins. It is noteworthy that like in some *USH3* cases,<sup>2</sup> stargazin mutation appears to affect also the inner ear vestibular function.<sup>48,49</sup>

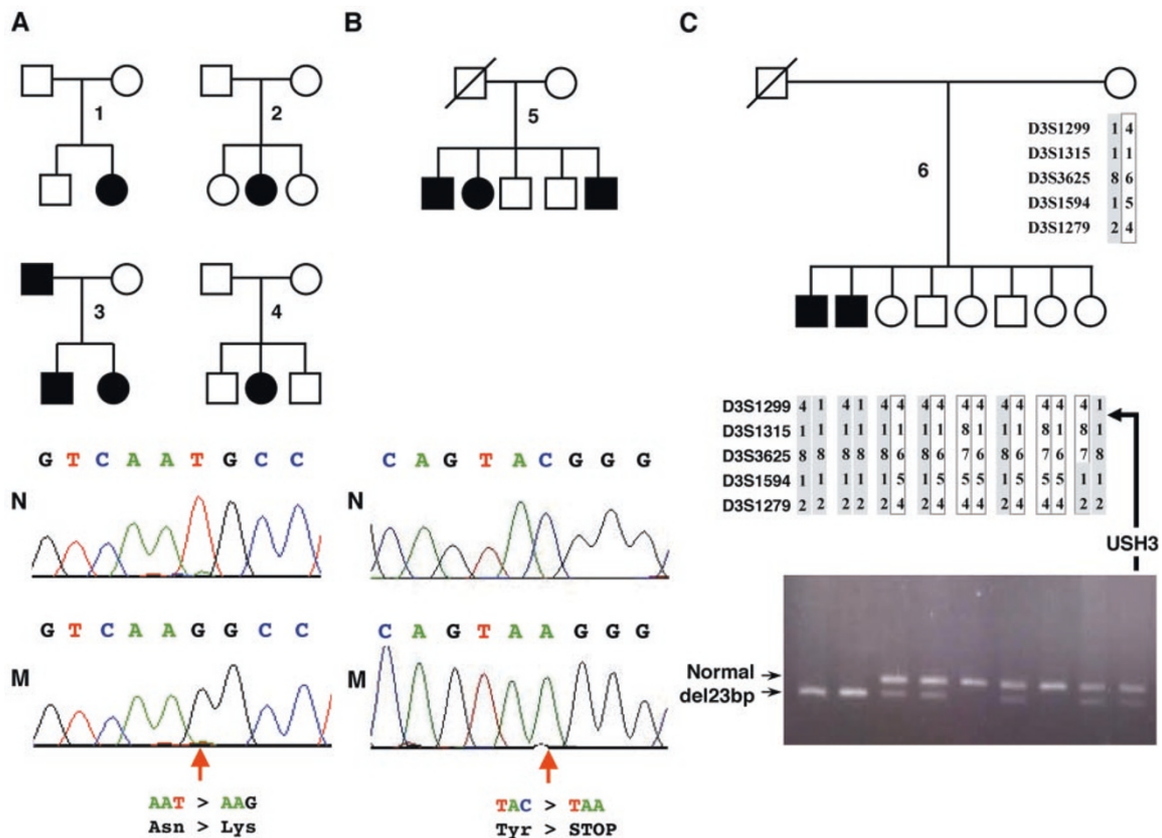
### USH3A mutations

Three novel *USH3* mutations were identified among 10 affected individuals from six *USH3A* families (Table 1 and Figure 4). A 143T>G substitution expected to cause a N48K missense mutation was found in six affected individuals from four unrelated families of Eastern European Jewish origin (Figure 4A). Shared microsatellite and SNP

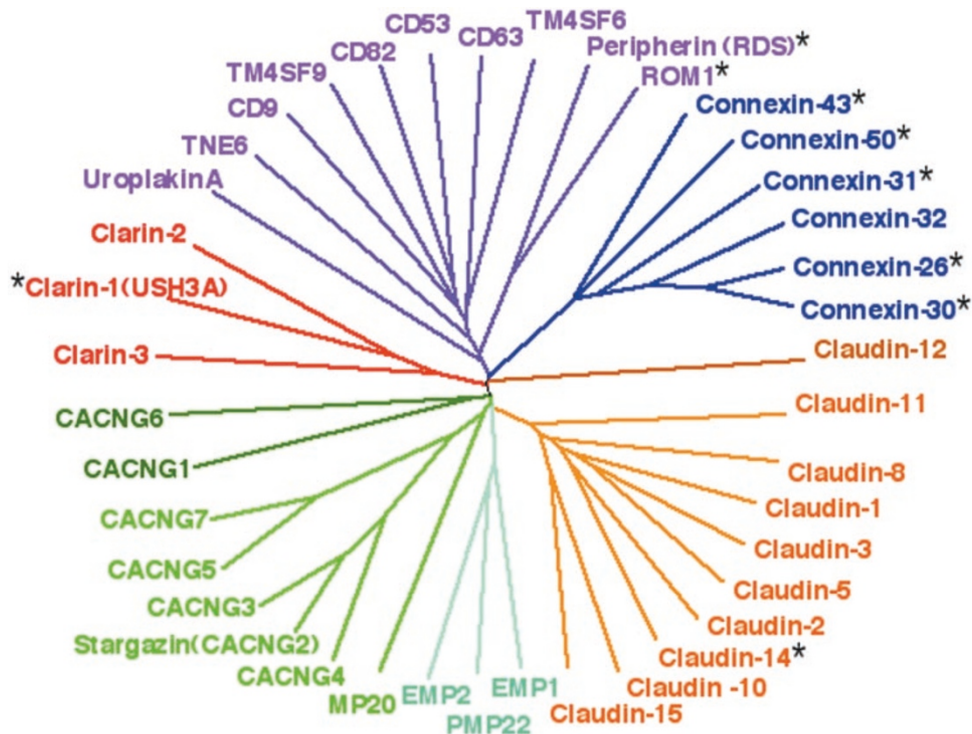
haplotypes on carrier chromosomes (data not shown) suggest the existence of a founder effect for this mutation. A 189C>A substitution, expected to cause a Y63X nonsense mutation, was found in a homozygous state in three affected individuals from a non-consanguineous Spanish family (Figure 4B; Table 1). A 23 bp deletion, spanning nucleotides 187–209 downstream from the first methionine codon, was found in a homozygous state in two affected individuals from a non-consanguineous family of Yemenite Jewish origin (Figure 4C).

### Discussion

Affiliation of clarin-1 to the newly defined clarin family and the definition of several motifs characterizing all members of this family indicate that the full-length coding sequence of the clarin-1 has now been determined. The additional 18aa encoded by the longest mouse transcript as well as



**Figure 4** *USH3* families and mutations. (A) Four Eastern-European Jewish *USH3* families. Below the pedigrees are parts of exon 0 chromatograms of DNA from non-carrier (top) and affected (bottom) individuals. Marked with an arrow is the 143T>G substitution, which is expected to cause an N48K missense mutation. Five patients from families 2–4 are homozygous for the 143T>G substitution while the patient from family 1 carries this substitution only on one allele. The second *USH3A* mutation of this individual was not detected. This result might be due to mutation in unexplored control regions or to a gross deletion, which in hemizygous state is hard to detect by the standard methods used in this work. (B) An *USH3* family of Spanish origin. Below are parts of exon 0 chromatograms of non-carrier and affected individuals. Marked with an arrow is the A 189C>A substitution that is expected to cause a Y63X nonsense mutation. (C) An *USH3* family of Yemenite Jewish origin. Below is FMC's MetaPhor agarose-gel electrophoresis of *USH3A* exon 0 fragments containing the 187–209 23 bp, which were amplified from genomic DNA samples of all family members.



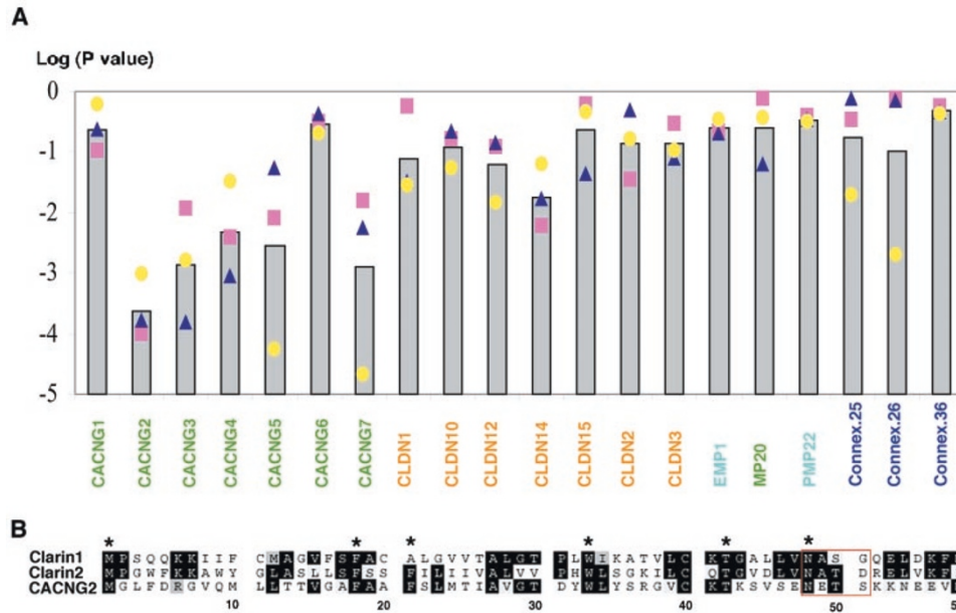
**Figure 5** Schematic relations of clarins to other four-transmembrane-domain proteins. Each sequence family is colour coded as follows: clarins: red, tetraspanins: purple, connexins: blue, claudins: orange and brown, EMPs and PMP22: turquoise, MP20 and calcium channel  $\gamma$  subunit-like proteins (CACNGs): green, CACNG1 and CACNG6: dark green. Claudins, EMPs, PMP22, MP20 and CACNGs all belong to the IPR000729 protein family. Marked with a black star are genes that were previously shown to be involved in forms of deafness or retinal diseases. Distances between all sequences were calculated by aligning each sequence pair. The presented unrooted tree was calculated from the resulting distance matrix using the Neighbor-Joining method.<sup>34</sup> Sequence SwissProt/TREMBL accessions are clarin-1 (*USH3A*): this work, clarin-2: XP\_068256.1, clarin-3: XP\_058398.1, CACNG1: CCG1\_HUMAN, CACNG2 (stargazin): CCG2\_HUMAN, CACNG3: CCG3\_HUMAN, CACNG4: CCG4\_HUMAN, CACNG5: CCG5\_HUMAN, CACNG6: CCG6\_HUMAN, CACNG7: CCG7\_HUMAN, MP20: LMIP\_HUMAN, PM22: PM22\_HUMAN, EMP1: EMP1\_HUMAN, EMP2: EMP2\_HUMAN, claudin 1: CLD1\_HUMAN, claudin 2: CLD2\_HUMAN, claudin 3: CLD3\_HUMAN, claudin 5: CLD5\_HUMAN, claudin 8: CLD8\_HUMAN, claudin 10: CLDA\_HUMAN, claudin 11: CLDB\_HUMAN, claudin 12: CLDC\_HUMAN, claudin 14: CLDE\_HUMAN, claudin 15: CLDF\_HUMAN, connexin 43: CXA1\_HUMAN, connexin 50: CXA8\_HUMAN, connexin 32: CXB1\_HUMAN, connexin 26: CXB2\_HUMAN, connexin 31: CXB3\_HUMAN, connexin 30: CXB6\_HUMAN, CD82: CD82\_HUMAN, CD53: CD53\_HUMAN, CD9: CD9\_HUMAN, CD63: CD63\_HUMAN, TM4SF6: T4S6\_HUMAN, TM4SF9: T4S9\_HUMAN, peripherin (RDS): RDS\_HUMAN, ROM1: ROM1\_HUMAN, TNE6: TNE6\_HUMAN, uroplakinA: UPKA\_HUMAN.

additional protein segments encoded by the long transcripts of *Trclarin-1* and *Hsclarin-2* elongate the putative first extracellular loop (Figure 3C). On the other hand, *Trclarin-3* and *Tnclarin-3* seem to be missing the C-terminal TM domain, indicated by a dashed blue line in Figure 3C. The fact that no orthologues were identified in the genomes of prokaryotes, yeast, plants, nematodes and insects, suggests that this gene family is limited to vertebrates.

The single missense *USH3A* mutation newly identified in this study, N48K, disrupts the molecule's only N-glycosylation consensus site (Figure 3C), and may render clarin-1 incapable of proper intracellular trafficking and plasma membrane insertion.<sup>50</sup> Both other newly identified mutations, the Y63X and the 187–209del23 bp mutations, are expected to result in truncated proteins. Thus, these three

mutations likely render clarin-1 functionally inactive, and account for the disease in most studied *USH3A* patients (Figure 4). Of the previously identified missense mutations,<sup>26</sup> one (M120K under the new enumeration) leads to the replacement of a hydrophobic methionine residue by a charged lysine inside or very near to the border of TM2, and another (IL153,154M) results in the shortening of TM3 by one residue. Because the four TMs are rather well conserved, such mutations may also be functionally deleterious.

*USH3* is progressive, i.e. at birth both vision and audition are much less severely impaired than later in life. This implies that the role played by clarin-1 in the hair cell and retina may display at least a measure of functional redundancy. The best candidates for serving as substitute for clarin-1 may be its two orthologues, clarin-2 and clar-



**Figure 6** (A) Significance values of sequence similarity scores (*P* values) between the three human clarin paralogues and some other four transmembrane-domain proteins. *P* values were calculated using the PRSS program<sup>47</sup> with 1000 shuffled sequences and the program's default parameters. Changing the shuffling method from uniform to window did not change the results significantly (not shown). Pink squares, blue triangles and yellow circles in this figure represent *P* values of comparisons with clarin-1, clarin-2 and clarin-3 respectively. Bars indicate the geometric means of the three. (B) Alignment of the first 58 amino acids of clarins-1 clarin-2 and stargazin. Boxed in red is the putative N-glycosylation site. Average positions of the inferred first extracellular loop and N-glycosylation site were found to be more similar between the clarins and the  $\gamma$  subunits calcium channel CACNG1-5, which include stargazin, than between the clarins and the claudins and other members of the IPR000729 protein family (data not shown).

**Table 2** *USH3A* Exon–Intron junctions and primers for genomic amplification of exons

Exon	5' Junction	3' Junction	Exon length	Primers for genomic amplification	PCR product
0	....	CGGTTCTCATgtaagtagcaattgc	> 290 bp	F: CAGAAAAGGAGAAAAGCCAAG R: CTGGGAAGAGTCTGCCTAAA	480 bp
1	....	CAGCAACCAAGtaggggtgcctgca	> 390 bp	F: TCACTATCTGAACTATCTTGTGT R: AAGCCCCTGAACCTTATAGG	910 bp
1b	tcactgagcacctaTATGTGCCAG	AGCTTATAACctaattggagaagac	87 bp	F: TTGTGGCCATTTTTGGAGAT R: CCCCAAACATGTATCAAGTGC	207 bp
2	cctttcggttctcaTTTTTCCAGA	TTCATTTACGtaagtacaaaattc	180 bp	F: TCAGAAGGATTTTGTAGTGATTTGA R: TCTTTTGGACATATTGAAAAGCACA	355 bp
3	tgagcttcattcagGCTCCTGTGG	....	> 1638 bp	F: ATGTCAATGGGGATGATGGT R: AGCATCTGGAACCTCGGTGT	1866 bp
3a	tgagcttcattcagGCTCCTGTGG	CTCATTCTGGgtcattttctttg	137 bp	F: ATGTCAATGGGGATGATGGT R: GGAGCCCATTCAGAAAATGA	411 bp
3b	caaattgatctcagCTGACTAAAG	TGTCCTTCAAgattctttccaata	3568 bp?	F: TTCCCCTGAATTACCCATCA R: AGCATCTGGAACCTCGGTGT	339 bp

in-3. However, it is also possible that other 4TM proteins might subserve this function. Future studies will be needed to clarify the mechanism of time-dependent loss of such potential redundancy.

Other small 4TM proteins have previously been shown to underlie deafness and retinal diseases. Mutations in four different connexins (26, 30, 31, and 43) and in claudin-14 underlie several forms of deafness (Hereditary Hearing Loss

Homepage: <http://www.uia.ac.be/dnlab/hhh/>). Mutations in connexin 50 cause congenital cataracts<sup>51</sup> and mutations in human peripherin/RDS and mouse ROM were shown to disrupt photoreceptor morphogenesis, leading to retinitis pigmentosa.<sup>45</sup> Also, mouse mutations in *Cacng2* (stargazin) cause defects not only in cerebellum but also in inner ear.<sup>52</sup> Yet, such information cannot help define more accurately the underlying mechanism for USH3, because the above

**Table 3** ESTs and mRNAs that align with the human and mouse *USH3A* transcripts

EST/mRNA	Specie	Align with	GenBank details	Organ/tissue & Dev. stage
<i>USH3A</i>				
AF388366	Hs		USH3A (USH3A) mRNA	Retina
AF388368	Hs		USH3A isoform b (USH3A) mRNA	Retina
W27577	Hs	2066-2473 (+)	cDNA randomly primed sublibrary	Retina
BB591018	Mm	1-210(+)	cDNA clone A830001O21 5'	10 days neonate cortex
BB639483	Mm	64-425; 660-977 (+)	cDNA clone A630099N24 5'	3 days neonate thymus
BB638319	Mm	65-340(+)	cDNA clone A630025E03 5'	3 days neonate thymus
BB689483	Mm	66-425; 660-977 (+)	cDNA clone 6820443B08 3'	12 days mullerian duct
BB630393	Mm	238-880 (+)	cDNA clone A130002D11 5'	16 days neonate thymus

Hs - Homo sapiens; Mm - Mus musculus.

listed genes participate in diverse cellular pathways, including the assembly and maintenance of gap junctions, tight junctions and synaptic junctions.<sup>53–55</sup>

Because stargazin has been shown to play a key role in the shaping and maintenance of cerebellar synapses,<sup>55</sup> our protein sequence analysis is suggestive of a synaptic role for clarin-1 as well. The case is strengthened by corroborative evidence: (a) Stargazin has been shown to interact with PDZ domain-containing proteins of the Membrane Associated Guanylate Kinase type (MAGUK), serving as intracellular anchors essential for the integrity of synaptic densities.<sup>55</sup> Intriguingly, one form of the Usher syndromes (USH1C) is caused by mutations in another PDZ protein harmonin. This may indicate that clarin-1 and harmonin are part of the same synapse-formation pathway, despite the absence of a clear PDZ-binding consensus<sup>56</sup> shared by all the clarins (Figure 3C). (b) Stargazin was shown to be involved in controlling the expression and mobilisation of amino-hydroxyl-methyl-isoxazole propionate (AMPA) glutamate receptors to the post-synaptic cleft in cerebellar granule cells. That cochlear hair cells also utilize glutamate (or a highly similar compound) as a rapid excitatory neurotransmitter<sup>57</sup> seems relevant. (c) Stargazin and other members of the CACNG family have been suggested to also play a role of forming protein–protein contacts across the synaptic cleft.<sup>58</sup> In bridging two cellular membranes they resemble other 4TM proteins such as the gap junction-forming connexins and the tight junction-forming claudins. That clarin-1 is expressed in inner ear, both pre- and post-synaptically, is consistent with possible clarin–clarin homophylic interactions, which might work in the hair cell synapse. Thus, we would like to propose that clarin-1 has a role in the excitatory ribbon synapse junctions between hair cells and cochlear ganglion cells,<sup>59</sup> and presumably also in analogous synapses within the retina.

We have previously reported a possible epistatic interaction between the *USH3A* locus and the *MYO7A* gene,<sup>27</sup> whereby two *USH3A* haploidentical patients of a Yemenite family showed different USH phenotypes, the patient with the more severe, USH1-like phenotype was also found to be a carrier for a complex rearrangement (3260T>C and 3266delG) in the tail of myosin VIIA.<sup>27</sup> Both patients are

shown here to be homozygous for the 23 bp deletion in *USH3A* exon 0 (Figure 4C). While the double heterozygous mother and siblings are healthy, in the context of a homozygous *USH3A* null mutation the presence of one null mutation in the *MYO7A* tail (the out of phase one base deletion in the complex rearrangement) mimics haploinsufficiency, and illustrates a departure from the monogenic model.

Since *MYO7A* expression in the inner ear is restricted to sensory hair cells,<sup>60–62</sup> including the sensory synaptic region,<sup>62</sup> the present observation that *Ush3a* is expressed in mouse sensory hair cells, and the implication of a synaptic role, are compatible with a clarin-1/*MYO7A* interaction. Furthermore, while myosin VIIA was localised only in pre-synaptic cells, MyRIP, a most recently defined *MYO7A*-interacting protein,<sup>63</sup> is present in both pre- and post-synaptic cells, similar to clarin-1. The proposed clarin-1/*MYO7A* interaction is also consistent with a pathophysiological continuum between *USH1* and *USH3A*, and is relevant to their treatment. Access to *Ush3* mutant models should allow testing of these hypotheses and contributing to a better understanding of the role of clarin-1 in the retina and inner ear.

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