Mutations in the Cilia Gene ARL13B Lead to the Classical Form of Joubert Syndrome

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Joubert syndrome (JS) and related disorders are a group of autosomal-recessive conditions sharing the “molar tooth sign” on axial brain MRI, together with cerebellar vermis hypoplasia, ataxia, and psychomotor delay. JS is suggested to be a disorder of cilia function and is part of a spectrum of disorders involving retinal, renal, digital, oral, hepatic, and cerebral organs. We identified mutations in ARL13B in two families with the classical form of JS. ARL13B belongs to the Ras GTPase family, and in other species is required for ciliogenesis, body axis formation, and renal function. The encoded Arl13b protein was expressed in developing murine cerebellum and localized to the cilia in primary neurons. Overexpression of human wild-type but not patient mutant ARL13B rescued the Arl13b scorpion zebrafish mutant. Thus, ARL13B has an evolutionarily conserved role mediating cilia function in multiple organs.

Introduction

Joubert syndrome (JS [MIM 213300]) is characterized by congenital cerebellar ataxia, hypotonia, oculomotor apraxia, and mental retardation. It is the most common inherited cerebellar malformation syndrome and is part of the autosomal-recessive cerebellar ataxia group of disorders.1 The neuroradiological hallmark in JS is a peculiar malformation of the midbrain-hindbrain junction known as the “molar tooth sign” (MTS), consisting of cerebellar vermis hypoplasia or dysplasia, thick and horizontally oriented superior cerebellar peduncles, and an abnormally deep interpeduncular fossa.2 The MTS has subsequently been reported in a group of syndromes termed Joubert syndrome and related disorders (JSRD). These include (1) classical JS (MIM 213300), (2) JS plus Leber congenital amaurosis (LCA, i.e., congenital retinal blindness [MIM 204000]),3 (3) JS plus nephronophthisis (NPHP [MIM 256700]), (4) JS plus LCA plus NPHP (also known as cerebello-oculo-renal syndrome [CORS [MIM 608091]]),4 (5) cerebellar vermis hypo/aplasia, oligophrenia, congenital ataxia, ocular coloboma, hepatic fibrosis (COACH [MIM 216360]) syndrome,5 and (6) oral-facial-digital syndrome type VI6,7 (MIM 277170), with occurrence of other features such as encephalocoele and polydactyly variably found within each subgroup. Many of these disorders have previously been linked to genes encoding cilia-localized proteins, so JSRD was considered a candidate ciliopathy disorder. The recent discoveries that JSRD can be due to mutations in CEP2908,9 (encoding a known centrosomal/cilia protein), as well as mutations in NPHP110,11 (previously implicated in the NPHP ciliopathy), RPGRIP112–14 (encoding a binding partner of proteins implicated in the ciliopathy retinitis pigmentosa), and TMEM67 (a.k.a. MKS3)15 (previously implicated in the Meckel syndrome ciliopathy) have further strengthened the hypothesis that JSRD is due to defective ciliary function.

Here we report the identification of the JBTS8 locus and mutations in the ARL13B gene, encoding ADP-ribosylation factor-like protein 13B. ARL13B is a small GTPase belonging to the class of Arf/Arl family within the Ras superfamily of small GTPases involved in diverse cellular functions.

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ARL13B is thus the first gene encoding an enzymatically active protein implicated in JS, suggesting that its study may shed light on the essential signaling pathways. Furthermore, an Arl13b null mutation was recently identified as underlying the lethal hemin mouse mutant that displays a Sonic Hedgehog-like phenotype, as well as being identified in the zebrafish scorpion mutant that displays renal cysts and curved tail, both tied to impaired cilia function. The identification of ARL13B mutations in humans with JS, together with the previous work in mouse and zebrafish, indicates an extremely broad spectrum of phenotypes across species all related to defective cilia function. The results also suggest a role for cilia-mediated Sonic Hedgehog signaling in cerebellar development and the pathogenesis of JS in humans.

Material and Methods

Genome Mapping
Family MTI-001 was recruited according to an approved Human Subjects Protocol at the University of Leeds and UCSD. DNA was extracted from peripheral blood leukocytes by salt extraction, genotyped with the Illumina Linkage IVb mapping panel, and analyzed with easyLINKAGE-Plus software, which runs Allegro version 1.2c in a PC Windows interface to calculate multipoint LOD scores. Parameters were set to autosomal recessive with full penetrance, and disease allele frequency of 0.001. Genomic regions with LOD scores over 2 were considered as candidate intervals. Linkage simulations were performed with Allegro 1.2c under regions with LOD scores over 2 were considered as candidate intervals. Linkage simulations were performed with Allegro 1.2c under the same parameters, with 500 markers at average 0.64 cM intervals, codominant allele frequencies, and parametric calculations. Additional subjects were enrolled according to approved protocols at University of Chicago, University of Michigan, University of Leeds, and Necker Hospital in Paris. Control samples were from healthy US adult subjects from a hypertension study, from healthy Arab controls from the United Arab Emirates, and healthy Pakistani controls from Leeds, UK.

Mutation Screening
We performed direct bidirectional sequencing of the ten coding exons and splice junction sites of ARL13B via BigDye Terminator cycle sequencing (Applied Biosystems). We screened 124 patients ascertained for the presence of nephropathis with or without features of Joubert syndrome (courtesy of Friedhelm Hildebrandt, U. Michigan) and 32 patients ascertained for the presence of Meckel syndrome with or without features of Joubert syndrome (courtesy of Tania Attie-Bitach, Necker Hospital, France) via bidirectional sequencing. We screened 44 patients ascertained for the presence of Meckel syndrome with or without features of Joubert syndrome (courtesy of Colin Johnson, U. Leeds) by LightScanner detection (Idaho Technologies), with subsequent bidirectional sequencing of aberrant bands. All patient samples were ascertained and managed according to approved human subjects protocols.

Candidate Genes and GenBank Accession Numbers
Human ARL6 has two GenBank numbers (NM_032146.1 and NM_177976.3) encoding the same protein, but the first encodes a longer transcript. Human ARL13B has two GenBank numbers (NM_144996.1 and NM_144996.2) differing in that the second entry is missing coding exons 2–3. Zebrafish arl13B has GenBank number NM_173272. Mouse Arl13b has GenBank number NM_026577.2.

Bioinformatics
Predicted folding and intermolecular interactions were predicted with Swiss-Model with crystallized ARL2-GTP as a template. The resulting ARL13B structure was manipulated and rendered with PyMOL software by importing amino acid sequence for wild-type and mutant and comparing the predicted hydrogen bonds.

GTP Binding Assay
Human wild-type and mutant ARL13B cDNA were cloned separately into the pGEX 6P1 vector (GE Lifesciences), and GST-tagged purified protein was isolated according to the manufacturer's recommendations. Protein was retrieved from the soluble phase, integrity verified by SDS-PAGE analysis followed by Coomassie staining, and analyzed by circular dichroism to validate that wild-type and mutant proteins had similar secondary structure as described. GTP binding was performed with a filter-based assay as described, with denatured (boiled for 15 min) Arl13b protein as negative control and Rac as a positive control, and analyzed with SigmaPlot Ver 10.0, with one-way ANOVA for statistical analysis.

Zebrafish Characterization
The arl13b<sup>−/−</sup> line was obtained from the Zebrafish International Resource Center (ZIRC) under an approved animal protocol at UCSD. One-cell embryos from arl13b<sup>+/−</sup> matings were injected with 50 pg in vitro transcribed capped open reading frame RNA (Ambion Message Kit) of human wild-type or mutant transcript. After 72 hr, embryos were phenotyped by an investigator that was blinded to the genotype, and then all embryos were subject to genotyping for correlation.

Histology
Mouse tissue was fixed with 4% paraformaldehyde, then processed for 15 μm cryostat sections and immunostained as described. We used two different rabbit polyclonal antibodies for Arl13b, one that was recently created against a bacterially expressed GST fused to amino acids 208–428 of murine Arl13B, and one created against an internal polypeptide, which behaved similarly, so only the first was used for further experimentation. Neither antibody showed reactivity with the cilia in the hemin mutant (data not shown). Antibodies were used at the following concentrations: Arl13b, 1:2000; acetylated tubulin (Zymed 32-2700), 1:500; calbindin (Swant), 1:600. Cerebellar granule neurons were isolated with a cell-density gradient as described, fixed, and immunostained at various time points after plating.

Results
Identification of the JBTS8 Locus at the Centromeric Region of Chromosome 3
In our series, we excluded linkage to known loci in 27 consanguineous JSRD families, supporting evidence for further genetic heterogeneity. One of the largest families (MTI-001, Figure 1A), which is of Pakistani origin and the first to be recruited in this study, displayed a proven MTS on brain imaging (Table S1 available online), provided a maximum expected LOD score of 3.0, and did not display any mutations...
in any of the five known JSRD genes (NPHP1, AHI1, CEP290, MKS3, RPGRIP1L). We performed a 0.64 cM resolution genome-wide screen by the Illumina Whole Genome SNP Linkage IVb panel, which defined an interval of 107–112 cM (80.0–103.3 MB) on chromosome 3p12.3-q12.3, with peak multipoint LOD score of 3.0 termed JBTS8 (Figure 1B).

Patients from MTI-001 Have a Homozygous p.R79Q Change in the ARL13B Gene

This interval encompassed the centromere, with 41 genes, and markers rs1905343 and rs938988 formed the p- and q-arm breakpoints, respectively (Figure 1C). Four genes within this interval were listed within the cilia proteome.

![Diagram](https://via.placeholder.com/150)
to localize to the ciliary axoneme in C. elegans sensory neurons, so we considered this the next-most-likely candidate gene. Sequence analysis identified a homozygous c.G236A mutation in exon 3 that is predicted to lead to a p.R79Q (positively charged to noncharged) amino acid substitution (Figure S1). This mutation was not encountered in 288 individuals (96 of Pakistani origin, 96 individuals of Arab origin, 96 of mixed US origin), providing >80% power to distinguish this from a normal sequence variant, and suggesting that it does not represent a common allele. Genotyping of all members of MTI-001 showed that this homozygous mutation segregated with the phenotype in affecteds but not in unaffecteds or obligate carriers.

Impaired GTP Binding Associated with the R79Q Mutation
The p.R79Q mutation identified in family MTI-001 occurred within the highly conserved GTP-binding domain. Therefore, we questioned whether this mutation might affect GTP binding. Because ARL13B is 36% identical and 61% similar in amino acid sequence with ARL2, we used the crystal structure of ARL2 to model the effect of the p.R79Q mutation on intramolecular forces. This modeling showed that R79 (R74 in ARL2) of the switch II domain establishes a hydrogen bond with D30 (D25 in ARL2) of the P loop domain. This interaction is conserved in other ARF family proteins and is predicted to be disrupted with alteration in amino acid charge because of the p.R79Q substitution (Figures 1E and 1F), and as a result might disrupt GTP binding. To test this hypothesis, we cloned both wild-type and p.R79Q ARL13B human cDNAs into bacterial expression vectors (Figure S2), isolated recombinant proteins (Figure S3), verified that both were soluble based upon their isolation from the liquid phase and that they had similar secondary structure based upon similar circular dichroism analysis (not shown), and then tested their ability to bind GTP in an established in vitro binding assay. We found that the concentration for half-maximal GTP binding by ARL13B was \(7.2 \times 10^{-7}\) M. At this concentration, the amount of GTP bound to the mutant protein was about half of the amount bound to the wild-type protein \((p < 0.05\) for each of three comparisons), whereas negative control was just above background (Figure 1G). We conclude that the p.R79Q mutation interferes with GTPase binding activity of ARL13B and that this mutation could lead to impairment of its cilia function.

Mutation Analysis in a JS Cohort
We tested for mutations in ARL13B within several cohorts of JSRD. By using bidirectional sequencing, we screened 182 patients from our cohort diagnosed with JSRD and identified two additional mutations in one family. Family MTI-423 has one affected female displaying a proven MTS on brain imaging (Figure S4). The affected female displayed a compound heterozygous mutation c.G246A (exon 3) resulting from a stop codon p.W82X from the mother and a c.C598T (exon 5) from the father that results in a positive charged to uncharged amino acid p.R200C missense mutation within the coiled-coil domain (Figure 2A; Figure S1). This family did not display mutations in any of the known JSRD genes (NPHP1, AHI1, CEP290, TMEM67, RPIP11L). Each of these ARL13B substitution mutations was fully conserved across evolution in all mammals as well as Xenopus tropicalis, Danio rerio, and Tetraodon nigroviridis (Figure 2B), and none of these mutations or any other predicted functional changes were identified by direct bidirectional sequencing of genomic DNA in a cohort of 182 healthy individuals (364 chromosomes). The early lethality of the hemnin mouse suggests a critical role for ARL13B in early embryonic development and may provide one explanation for the low percentage of ARL13B mutations in the JSRD cohort and the low representation of truncating mutations among the ARL13B patients.

Delineation of the Phenotypic Spectrum Associated with ARL13B Mutations
The phenotype observed in patients with identified mutations consists predominantly of classical Joubert syndrome. All patients displayed the MTS (Table S1). In MTI-001, two affecteds each displayed a small occipital encephalocele. In the other family, there was no occipital encephalocele, and there were no other supratentorial cerebral abnormalities and no renal abnormalities upon diagnostc ultrasound (performed on both families). Study of urinary concentration defects may be a more sensitive measure of early kidney involvement and was not available on any of these patients, so it is possible that renal symptoms might develop in the future. One affected from MTI-001 (now deceased) displayed evidence of mild nonspecific pigmentary retinopathy on clinical examination, although electroretinogram was normal on the affected siblings, so the involvement of ARL13B in retinal function is not yet clear. We also screened a cohort of 124 patients with JS with nephronophthisis and a cohort of 76 patients with overlapping Meckel syndrome with JS features. No
mutation in either of these cohorts was encountered, suggesting that mutations in ARL13B are restricted to the classical form of JSRD (which can include encephalocele and retinopathy) and are not identifiable in other related ciliopathies, at least at this level of detection.

Human Wild-Type but Not Mutated ARL13B Rescue the Zebrafish Scorpion Mutant

To test the evolutionary conservation of ARL13B, we utilized zebrafish (Danio rerio). Comparison of the human and zebrafish ARL13B shows 75% similarity and 59% identity in more than 90% of the predicted amino acid sequence. We took advantage of the scorpion zebrafish mutant in the arl13b locus. The arl13bsco allele is due to a retroviral insertion into the first exon, which inactivates the gene, as evidenced by the finding that morpholino antisense treatment against arl13b produced a phenocopy of the arl13bsco mutant. These mutants display fully penetrant curved tail and cystic kidney phenotypes, which are evident by 72 hr post fertilization (hpf) (n = 75, Figures 2C–F).
Arl13b protein was localized to cilia of all organs examined and we observed Arl13b staining of cilia structures within these tubules (Figure 3A). This staining might predict a role for Arl13b in kidney function, although none of our patients with ARL13B mutations displayed evidence of impaired kidney function. The retinal photoreceptor inner and outer segments, corresponding to the cell body and rhodopsin-containing disks, are bridged by a connecting cilium. We observed robust Arl13b staining of the connecting cilium at P10 (Figure 3B), similar to many of the proteins involved in LCA. It is interesting to note that one of the patients with an ARL13B mutation displayed evidence of retinopathy (Table S1). It will be important to follow the remaining patients to determine whether they develop progressive involvement of these organs in the future.

To test for expression in the developing cerebellum, the major site of involvement in JSRD, we examined Arl13b localization in postnatal mice. At the P0 time point, the cerebellum displays well-developed folia and consists of three well-defined layers: the external granule layer (EGL), Purkinje cell layer (PCL), and internal granule layer (IGL). Granule neuron precursors proliferate and migrate parallel to the surface of the cerebellum within the EGL. Postmitotic granule neurons then migrate radially into the cerebellar parenchyma and past the PCL to settle in the IGL. We detected strong staining of cilia-like structures on granule neurons both in the EGL and IGL (Figure 3C), which is consistent with previous serial electron microscopic sectioning of the cerebellum, as well as recent studies that have used staining for adenylate cyclase III to label cilia. Staining of Arl13b across the spectrum of cerebellar development in mouse demonstrated cilia-like structures in granule neuron populations, both in the EGL and IGL (Figure S5). There was a time dependence to the staining of cilia-like structures during cerebellar development, with about 40% of cells showing an immunopositive cilium at E16, dropping to less than 1% of cells by P21.

In order to be certain that these ARL13B-immunopositive structures represented cilia, we examined sections from the cerebellum at P0 in a transgenic mouse that expresses EGFP fused to Centrin 2, a constitutive centrosome/basal body marker that localizes to centrioles. As expected, we found that most cells in the EGL contained a readily identifiable pair of centrioles adjacent to the nucleus, representing the basal body. We found an immunopositive ARL13B structure emanating from the region surrounding the basal body in many of these cells (Figure 3C), thereby demonstrating the spatial relationship between the two structures and suggesting that ARL13B marks the cilium. However, in many cells, ARL13B was not identified near a pair of centrioles, suggesting the possibility that ARL13B stains the cilium in only a fraction of granule neurons or that only a fraction of granule neurons contain a cilium.

In order to differentiate between these possibilities, we isolated cerebellar granule neuron precursors from P5 mouse by using a cell-density gradient, cultured and fixed...
at various time points. We found that a fraction of these primary cultured neurons display a cilia, as evidenced by labeling of a structure protruding from the cell body that was positive for acetylated tubulin (Figure 3D). Although the acetylated tubulin staining was strongly positive in the cilia, it extended into many of the microtubules within the cell soma. All cells with a cilium based upon this staining also displayed immunopositivity for Arl13b, and Arl13b staining was largely restricted to the cilia. Thus, Arl13b appears to be a sensitive and specific marker for the cilium in many cells including in the developing cerebellum. The data also support a role for Arl13b in the pathogenesis of JSRD, and thus strongly links JSRD to other human ciliopathies.

Discussion

ARL13B encodes ADP-ribosylation factor-like protein 13B (previously known as ADP-ribosylation factor-like protein 2-like 1) and belongs to the small Ras GTPase superfamily. ADP-ribosylation factor (ARF) GTP-binding proteins are among the best-characterized members of the Ras superfamily of GTPases, with well-established roles in membrane trafficking pathways. However, the ARL proteins appear to be functionally distinct, and many have roles in microtubule function.176 A mutation in Arl13b was very recently identified as the cause for the murine hemin phenotype that displays a disrupted neural tube during embryogenesis.19 The encoded Arl13b protein is localized to the primary cilium where it presumably plays a role in microtubule assembly or transport akin to the roles of other Arl proteins. However, the Arl13b<sup>mm/mbm</sup> mutant is lethal at embryonic day 13.5–14.5 because of the severe cilia phenotype, thereby precluding a more detailed analysis of later functions.

Currently, mutations are identified in CEP290 in about 40% of patients that also display NPHP and LCA,50 in AHI1 in 10%–20% of patients mostly without NPHP,51,52 and in probably lower percentages in TMEM67, NPHP1, and RGRIP1L genes. Although there are two additional
Supplemental Data

Supplemental Data include a list of other members of the International JSRD Study Group, five figures, and one table and are available at http://www.ajhg.org/.

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Web Resources

The URLs for data presented herein are as follows:
Pfam database, http://www.sanger.ac.uk/Software/Pfam/
Human Genome Browser, http://www.genome.ucsc.edu/
Zebrafish database, http://zfinfo /
Pymol, http://www.pymol.org

References


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