OFD1 Is Mutated in X-Linked Joubert Syndrome and Interacts with LCAS-Encoded Lebercilin

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We ascertained a multi-generation Malaysian family with Joubert syndrome (JS). The presence of asymptomatic obligate carrier females suggested an X-linked recessive inheritance pattern. Affected males presented with mental retardation accompanied by postaxial polydactyly and retinitis pigmentosa. Brain MRIs showed the presence of a “molar tooth sign,” which classifies this syndrome as classic JS with retinal involvement. Linkage analysis showed linkage to Xpter-Xp22.2 and a maximum LOD score of 2.06 for marker DXS8022. Mutation analysis revealed a frameshift mutation, p.K948NfsX8, in exon 21 of OFD1. In an isolated male with JS, a second frameshift mutation, p.E923KfsX3, in the same exon was identified. OFD1 has previously been associated with oral-facial-digital type 1 (OFD1) syndrome, a male-lethal X-linked dominant condition, and with X-linked recessive Simpson-Golabi-Behmel syndrome type 2 (SGBS2). In a yeast two-hybrid screen of a retinal cDNA library, we identified OFD1 as an interacting partner of the LCAS-encoded ciliary protein lebercilin. We show that X-linked recessive mutations in OFD1 reduce, but do not eliminate, the interaction with lebercilin, whereas X-linked dominant OFD1 mutations completely abolish binding to lebercilin. In addition, recessive mutations in OFD1 did not affect the pericentriolar localization of the recombinant protein in hTERT-RPE1 cells, whereas this localization was lost for dominant mutations. These findings offer a molecular explanation for the phenotypic spectrum observed for OFD1 mutations; this spectrum now includes OFD1 syndrome, SGBS2, and JS.

Introduction

Joubert syndrome (JS [MIM 213300]) is characterized by a specific mid-hindbrain malformation, hypotonia, cerebellar ataxia, and developmental delay. Oculomotor apraxia and abnormalities in breathing patterns are frequently part of this condition as well. The typical cerebellar and brainstem malformations in JS result in a characteristic neuroradiological hallmark known as the “molar tooth sign.”1 The molar tooth sign is not restricted to classical JS but has also been observed in disorders in which more organ systems are affected. This group of syndromes was termed Joubert syndrome and related disorders (JSRDs)2, and a classification into six subgroups has been proposed.3 Apart from the classical form, JS can occur in combination with progressive retinal degeneration,4 renal abnormalities (nephronophthisis, NPHP [MIM 256100]),5 both retinal and renal involvement (cerebello-oculo-renal syndromes, CORS [MIM 608091]),6 ocular colobomas and liver abnormalities (COACH [MIM 216360]),7 and both orofacial and digital signs (OFDVI [MIM 277170]).8 These additional features are characteristic of disorders caused by primary ciliopathy and/or basal-body dysfunction,9 indicating that the pathogenic mechanism in JSRDs probably involves defective ciliary and/or basal-body function. So far, nine autosomal JSRD loci have been mapped, and causative genes have been identified for eight of these.3,10 All eight JSRD genes encode proteins that localize to cilia or centrosomes, reiterating ciliary dysfunction as a key factor in the molecular pathogenesis of JSRDs. The AHI1 (MIM 608894) gene is most frequently associated with a combination of JS and retinopathy and encodes jouberin, which physically interacts with the NPHP1-encoded nephrocystin-1 protein (MIM 607100).11–13 ARL13B (MIM 608922) encodes a small GTPase that belongs to the Arf/Arl class of the Ras GTPase family.10 Animal ARL13B-null models emphasize the ciliary importance of this gene in that they reproduce cilia-related phenotypes such as disruption of neural-tube development, renal cysts, and overall defective ciliophylogenetic morphology.14,15 The protein product encoded by the CC2D2A (MIM 612013) gene functions in close association with the centrosomal protein CEP290.16 A nonsense mutation in the zebrafish CC2D2A ortholog (sentinel) results in the formation of pronephric cysts. The most recently identified JSRD gene is INPP5E, which is the causative gene for the JBTS1 locus. The encoded protein functions in phosphatidylinositol signaling, linking this pathway to the ciliopathies.17 The remaining four JSRD genes have also been implicated in other overlapping human ciliary disorders: NPHP1 in isolated NPHP18.

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TMEM67/MKS319 (MIM 609884), RPGRIP1L20,21 (MIM 610937), and also CC2D2A22 in Meckel syndrome (MIM 249000), and CEP290 (MIM 610142) in isolated Leber congenital amaurosis (LCA [MIM 204000]),24 Senior-Løken syndrome (SLs, a combination of retinal degeneration and NPHP [MIM 266900]),24 and Meckel syndrome.25 The exact biochemical pathway in which JSRD proteins play a role remains unclear. However, experimental evidence is pointing to cilia-mediated Sonic hedgehog (Shh) signaling as a likely candidate.15,20,26

In this study, we have identified mutations in exon 21 of the OFD1 (MIM 311200) gene in two families with classical JS, thereby defining OFD1 as the JBTS10 locus. In one family, patients also suffer from retinal pathology. To our knowledge, this is the first report of X-linked inheritance in JS and of mutations in OFD1 downstream of exon 17. The OFD1 gene has previously been associated with oral-facial-digital type 1 (OFD1) syndrome (MIM 311200),27 which is a male-lethal X-linked dominant condition involving malformations of the face, oral cavity, and digits in affected females. Budny and coworkers also described a single family in which OFD1 was found to be mutated in Simpson-Golabi-Behmel syndrome type 2 (SGBS2 [MIM 300209]).28

We show that OFD1 directly binds to the ciliary protein lebercilin, encoded by the LCA5 (MIM 611408) gene.29 Mutations in LCA5 cause LCA, an inherited condition of very early-onset childhood blindness that is due to retinal degeneration and also belongs to the ciliopathy spectrum. We also show that X-linked recessive mutations in OFD1 reduce, but do not eliminate, the interaction with lebercilin and do not affect ciliary localization in cell lines, whereas X-linked dominant OFD1 mutations completely abolish binding with lebercilin and disrupt ciliary localization as well. Our findings offer a molecular explanation for the phenotypic variability observed in OFD1-related disorders.

Material and Methods

Patients

In all patients from family W07-713, karyotypes at a resolution of 550 bands were normal, and expansions of the CGG repeat in the 3′-untranslated region of FMR1 were excluded. In patient IV-10, submicroscopic copy-number variations larger than 100 kb were excluded by Affymetrix 250k SNP array analysis in which was used as a restriction enzyme according to the manufacturer’s protocols (Affymetrix, Santa Clara, CA, USA). Copy-number estimates were made with the CNAG software package v. 2.0.30 In addition, probands from 84 JS families in which only males were affected were selected from a cohort of 250 families. All families showed a molar tooth sign on MRI scan and had developmental delay and ataxia. Known genes and loci for JSRD (AH11, AR1L13B, CC2D2A, CEP290, NHPH1, RPGRIP1L, TMEM67, and chromosomal loci 9q34 and 11p11.2–q12.1) were excluded in most cases by direct sequencing or segregation analysis. All DNA samples were isolated from whole blood by the salting-out method as described by Miller et al.31 Informed consent was obtained for all patients, and research was approved by the respective local ethics committees.

X-Chromosomal Linkage Analysis

Polyinform short-tandem-repeat markers on the X chromosome were selected with an average distance of 10 cM between the markers. Primers for amplifying these markers were designed with the Primer3 program.32 An M13 tail was added to the 5′ and 3′ ends of the primers. Markers were amplified with an M13 forward primer labeled with one of the fluorophores, FAM, VIC, NED, or ROX, at the 5′ end23 and an M13 reverse primer with a 5′-GTTCCTT-3′ added to its 5′ end so that tailing would be reduced.34 Primer sequences are given in Table S1. PCR conditions are available on request. Final PCR products were mixed with eight volumes of formamide and half a volume of Genescan 500–250 LiZ size standard (Applied Biosystems, Foster City, CA, USA) and analyzed with the ABI PRISM 3730 DNA analyzer (Applied Biosystems). The results were evaluated by GeneMapper (Applied Biosystems). Two-point LOD scores were calculated by Superlink,35 and exclusion mapping was performed with GeneHunter PLUS v. 1.236; both programs are integrated in the computer program easyLINKAGE.37 Inheritance model was set at recessive, and the frequency of the disease allele was set at 0.001. Full penetrance was assumed.

DNA Sequencing

Primers for amplification of all exons of OFD1 (GenBank ID NM_003588.3) were designed with the Primer3 program (Table S1).32 Of the last exon, 374 nucleotides of the 3′-untranslated region were analyzed. PCR conditions are available on request. The ABI PRISM BigDye Terminator Cycle Sequencing v. 2.0 Ready Reaction Kit was used for sequencing PCR products with the same forward and reverse primers as those used in the PCR, and the products were analyzed with the ABI PRISM 3730 DNA analyzer (Applied Biosystems).

Amplification Refractory Mutation System

Amplification Refractory Mutation System (ARMS) primers for specific amplification of either the wild-type or mutant allele were designed with the Primer3 program (Table S1).32 The wild-type or mutant alleles were amplified from 50 ng DNA by the use of 10 U Taq polymerase (Invitrogen, Breda, The Netherlands) in Taq buffer, 2.0 mM MgCl2, 0.25 mM dNTPs (Invitrogen, Breda, The Netherlands), and 100 nM forward and reverse primers in a total volume of 25 μl. PCR cycling conditions consisted of (1) denaturation at 95°C for 10 min, (2) 40 cycles of amplification by denaturation at 95°C for 15 s, annealing at 58°C for 15 s, and elongation at 72°C for 15 s, and (3) final elongation at 72°C for 10 min. PCR products were analyzed on a 1.5% agarose gel.

X Inactivation

Skewing of X inactivation was investigated via analysis of the CGG repeat in the promoter region of FMR1, for which the tested carrier females had two different alleles. One microgram DNA from female carriers was ampliﬁed from 2.5 μl of the 5′-GTTTCTT-3′ end so that tailing would be reduced.34 Primer sequences are given in Table S1. PCR conditions are available on request. Final PCR products were mixed with eight volumes of formamide and half a volume of Genescan 500–250 LiZ size standard (Applied Biosystems, Foster City, CA, USA) and analyzed with the ABI PRISM 3730 DNA analyzer (Applied Biosystems). Two-point LOD scores were calculated by Superlink,35 and exclusion mapping was performed with GeneHunter PLUS v. 1.236; both programs are integrated in the computer program easyLINKAGE.37 Inheritance model was set at recessive, and the frequency of the disease allele was set at 0.001. Full penetrance was assumed.
LCLs from patients and controls were grown at 37°C for 10 min, (2) 32 cycles of amplification by denaturation at 95°C for 15 s, annealing at 64°C for 2 min, and elongation at 75°C for 2 min, and (3) final elongation at 72°C for 10 min. PCR products were analyzed as described in the X-Chromosomal Linkage Analysis section. Complete digestion of one of the two alleles was confirmed by the absence of a PCR product in the BamHI- and HhaI-digested DNA of the hemizygous male control, whereas in all other samples at least one allele was amplified.

Cell Culturing
Human B-lymphocytes were immortalized by transformation with Epstein-Barr virus according to established procedures. 38 EBV-LCLs from patients and controls were grown at 37°C and 7.5% CO2 in RPMI 1640 medium (GIBCO, Breda, The Netherlands) containing 10% (v/v) fetal calf serum (Sigma, Zwijndrecht, The Netherlands), 1% 10 U/μl penicillin and 10 μg/μl streptomycin (GIBCO), and 1% GlutaMAX (GIBCO). Twenty-four hours before emetin treatment, cells were centrifuged at 200 x g for 5 min at room temperature and resuspended in fresh medium to a density of 0.7 million cells per ml. Cells were treated with 100 μg/ml emetin for 10 hr. Subsequently, 5–10 million cells were harvested by centrifugation at 200 x g for 5 min at room temperature, washed with PBS, and pelleted by centrifugation at 200 x g for 5 min at room temperature. Pellets for RNA isolation were snap frozen in liquid nitrogen.

RNA Isolation and First-Strand Synthesis
Total RNA was isolated with the NucleoSpin RNA II kit (Macherey-Nagel, Düren, Germany) according to the manufacturer's protocol. For removing residual traces of genomic DNA, the RNA was treated with DNase I (Invitrogen, Breda, The Netherlands) while bound to the RNA-binding column. The integrity of the RNA was assessed on 1.2% agarose gel, and the concentration and purity were determined by optical densitometry. One microgram of total RNA was transcribed into cDNA with the iScript cDNA synthesis kit (Bio-Rad Laboratories, Hercules, CA, USA) according to the manufacturer's protocol. cDNA was purified with the NucleoSpin extract II kit (Macherey-Nagel) according to the manufacturer's protocol.

Quantitative PCR
SYBR Green-based real-time quantitative PCR (QPCR) analysis was performed on a 7500 Fast Real-Time PCR System (Applied Biosystems) with Power SYBR Green PCR Master Mix (Applied Biosystems) according to the manufacturer's instructions. Primers were developed by the Primer3 program 39 (Table S1) and validated as described before. 39 PCR products encompassed at least one exon-exon junction. GUSR was used as reference gene. QPCR quantifications were performed in duplicate on the equivalent of 400 pg/μl input of total RNA in the first-strand synthesis and included a reverse-transcriptase control. Values for experimental threshold cycles (Ct) were within the range of cDNA dilutions used for validating the primers. The melt curves of all PCR products showed a single PCR product. All controls were negative. Differences in the expression of a gene of interest between two samples were calculated by the comparative Ct or 2ΔΔCt.

DNA Constructs
Constructs encoding the full-length OFD1 variant of 1012 amino acids (OFD1 33; NCBI Reference Sequence: NM_003611.2 [gene]; NP_003602.1 [protein]) were generated by PCR with an OFD1 cDNA clone that was kindly donated by Prof. Brunello Franco (TIGEM Telethon Institute of Genetics and Medicine, Naples, Italy). For this variant, the Coils2 program 42 predicts six coiled-coil domains. For the constructs encoding coiled-coil domains two/three (fusion of coiled-coil domains two and three) and four and coiled-coil domains five and six of OFD1, OFD1 cDNA IMAGE clone (GenBank BC096344) (imaGenes, Berlin, Germany) was used as a template in PCR. This clone contains an alternative splice variant that lacks exon 10 and thus encodes a 972 amino acid variant protein (OFD1 Δex10 35) that lacks the corresponding amino acids 313–353. For this splice variant, five coiled-coil domains are predicted because coiled-coil domains two and three are fused into one larger coiled-coil domain (coiled-coil domain two/three). The OFD1 constructs encoding amino acids 240–1012 and amino acids 356–1012 were generated by PCR on template DNA from the corresponding clones that were identified in the yeast two-hybrid assay, from which the first clone includes exon 10. Expression constructs were created with Gateway technology (Invitrogen, Leek, The Netherlands) according to the manufacturer's instructions. LCA5 constructs were generated as previously described. 29 Constructs for OFD1 mutants were generated by site-directed mutagenesis PCR. Primers are available upon request. The sequence of all entry clones was verified by nucleotide sequencing.

Yeast Two-Hybrid Assays
The GAL4-based yeast two-hybrid system (HybriZAP, Stratagene, La Jolla, USA) was used for identifying protein-interaction partners of lebercilin. The region encompassing the first 305 N-terminal amino acids of lebercilin, fused to a DNA-binding domain (GAL4-BD), was used as a bait for screening a human oligo-dT primed retinal cDNA library. The yeast strain PJ69-4A, which carries the HIS3 (histidine), ADE2 (adenine), MEL1 (α-galactosidase) and LacZ (β-galactosidase) reporter genes, was used as a host. Interactions were analyzed by assessment of reporter gene activation via growth on selective media (HIS3 and ADE2 reporter genes), α-galactosidase colorimetric plate assays (MEL1 reporter gene), and β-galactosidase colorimetric filter lift assays (LacZ reporter gene). For analysis of the binding capacities of OFD1 mutant proteins to lebercilin, expression constructs encoding amino acids 356–1012 of OFD1 as a GAL4-AD-fusion protein, either wild-type or containing the p.K948fs, p.K923fs, p.I784fs, p.E709fs, p.N630fs, or p.S586fs mutation, were cotransformed with a construct encoding the first two coiled-coil domains of lebercilin fused to GAL4-BD (pBD-lebercilinCC1-2) in PJ69-4A. As a negative control, the empty pAD vector was cotransformed with pBD-lebercilinCC1-2. Interactions were evaluated on the basis of growth on selective media and staining in α- and β-galactosidase activity assays. The interactions of wild-type and mutant OFD1 with wild-type lebercilin were semi-quantified with a liquid β-galactosidase assay. Liquid cultures of PJ69-4A, co-transformed with combinations of pAD-OFD1 and pBD-lebercilin (n = 4 for each combination), were grown for 24 hr at 30°C in SD medium lacking leucine and tryptophan. Subsequently, the optical density of the cultures was determined at a wavelength...
of 600 nm for data normalization. Cell lysis and colorimetric reactions were performed with the Yeast β-galactosidase assay kit (Pierce Biotechnology, Rockford, IL, USA) according to the manufacturer’s instructions. β-galactosidase activity was determined from absorbance at a wavelength of 420 nm. For the calculation of relative remaining reporter gene activity, values were corrected for background activity in the assay.

Colocalization in hTERT-RPE1 Cells

hTERT-RPE1 cells (kindly provided by Prof. Uwe Wolfrum, Johannes-Gutenberg Universität, Mainz, Germany) were cultured as previously described. This is a human retinal pigment epithelial (RPE) cell line that is immortalized by stable expression of human telomerase reverse transcriptase (hTERT) (Clontech, Saint-Germain-en-Laye, France). Cells were seeded on coverslips and were serum starved for 24 hr prior to transfection so that primary cilium formation would be induced. Subsequently, Effectene (QIAGEN, Venlo, The Netherlands) was used for transfecting cells. Single constructs encoding either wild-type or mutant OFD1 fused to enhanced cyan fluorescent protein (eCFP) or a combination of constructs encoding eCFP-OFD1 and constructs encoding full-length lebercilin fused to monomeric red fluorescent protein (mRFP) were transfected. After 24 hr, cells were fixed in icecold methanol for 10 min, treated with 1% Triton X-100 in phosphate-buffered saline (PBS) for 5 min, and blocked in 2% bovine serum albumine in PBS for 20 min. Incubation with the primary antibody GT335 (mouse monoclonal antibody, kindly provided by Carsten Janke, CNRS Centre de Recherches en Biochimie Macromoléculaire, Montpellier, France) was performed for 1 hr. After being washed in PBS, coverslips were stained with goat-anti-mouse Alexa 568 (Invitrogen, Leek, The Netherlands) for 45 min. Coverslips were washed again with PBS and briefly with mRFP before being mounted in Vectashield containing DAPI (Vector Laboratories, Burlingame, CA, USA). The localization of constructs was analyzed with a Zeiss Axio Imager Z1 fluorescence microscope equipped with a 63× objective lens.

GST Pull-Down Assay

Both OFD1 splice variants and OFD1 fragments (amino acids 240–1012 and amino acids 356–1012) were cloned into pDEST15 (Gateway cloning system, Invitrogen, Karlsruhe, Germany). For the creation of GST-fusion proteins, BL21-DE3 cells were transformed with pDEST15 constructs. Cells were induced overnight with 0.5 mM IPTG at 30°C and subsequently lysed in STE buffer (10 mM Tris-Cl [pH 8.0], 1 mM EDTA, and 150 mM NaCl) supplemented with 10 mg/ml Lysozyme, 0.5% Sarkosyl, 1% Triton X-100, and complete protease inhibitor cocktail (Roche Diagnostics, Mannheim, Germany). Lysates were incubated with glutathione-sepharose 4B beads (Amersham Biosciences, Freiburg, Germany). After incubation, beads were washed with STE and TBSTD (TBS with 1% Triton X-100 and 2 mM DTT). The amount of GST-fusion proteins bound to the beads was verified on a NuPAGE Novex 4%–12% Bis-Tris SDS-PAGE gel by staining with SimplyBlue SafeStain. The presence of 3xFLAG-lebercilin in complex with GST-OFD1 was assessed by immunoblotting, followed by staining with a monoclonal mouse α-FLAG primary antibody (Sigma-Aldrich, Zwijndrecht, The Netherlands) and goat-anti-mouse coupled to IRDye800 (Rockland Immunochemicals, Gilbertsville, PA, USA) as a secondary antibody. Fluorescence was analyzed on a LI-COR Odyssey 2.1 infrared scanner.

Coimmunoprecipitation

HA-lebercilin (full-length) and 3xFLAG-OFD1 (both splice variants and fragments encompassing amino acids 240–1012 and amino acids 356–1012) were coexpressed in HEK293T cells. As a negative control, HA-lebercilin was coexpressed with the functionally unrelated 3xFLAG-LRRK2 (Leucine-Rich Repeat Kinase2) protein. As a positive control, the previously described interaction between nephrocin4 and RPGRIP1 was used. After 48 hr of expression, cells were lysed on ice in lysis buffer (50 mM Tris–HCL [pH 7.5], 150 mM NaCl, 0.5% Triton X-100) supplemented with complete protease inhibitor cocktail. Lysates were incubated with anti-HA affinity matrix (Roche, Woerden, The Netherlands), or with anti-FLAG M2-agarose from mouse (Sigma-Aldrich, Zwijndrecht, The Netherlands), for 5 hr at 4°C. After incubation, beads with bound protein complexes were washed in lysis buffer. Then beads were taken up in 4× NuPage Sample Buffer and heated for 10 min at 70°C. Beads were precipitated by centrifugation, and supernatant was loaded on a NuPAGE Novex 4%–12% Bis-Tris SDS-PAGE gel. The presence of 3xFLAG-lebercilin in complex with GST-OFD1 was assessed by immunoblotting, followed by staining with a monoclonal mouse α-FLAG primary antibody (Sigma-Aldrich, Zwijndrecht, The Netherlands) and goat-anti-mouse RDIye800 as a secondary antibody. Fluorescence was analyzed on a LI-COR Odyssey 2.1 infrared scanner.

Immunofluorescence Labeling of Rat Retinas

Unfixed eyes of 20-day-old (p20) Wistar rats were harvested and frozen in melting isopentane. Seven micrometer cryosections were cut and treated with 0.01% Tween in PBS and subsequently blocked in blocking buffer (0.1% ovalbumin and 0.5% fish gelatin in PBS). Then the cryosections were incubated overnight with primary antibodies diluted in blocking buffer, either with affinity-purified α-lebercilin rabbit serum in combination with a mouse α-centrin monoclonal antibody (kindly provided by Prof. Uwe Wolfrum, Johannes-Gutenberg Universität, Mainz, Germany) or with affinity-purified α-OFD1 rabbit serum (kindly provided by Prof. E. Nigg, Max-Planck Institut für Biochemie, Martinsried, Germany) in combination with a mouse α-centrin monoclonal antibody. Secondary antibodies were also diluted in blocking buffer and incubated in a dark environment for 1 hr. Prolong Gold Antifade (Molecular Probes, Leiden, The Netherlands) was used for embedding the sections. Pictures were made with an Axioskop2 Mot plus fluorescence microscope (Zeiss, Oberkochen, Germany), which was equipped with an AxioCam MRC5 camera (Zeiss). Images were processed with Axiovision 4.3 (Zeiss) and Adobe Photoshop (Adobe Systems, San Jose, CA, USA). Procedures were in accordance with the ethical standards of the responsible committee on animal experimentation.

Expression Analysis of OFD1 Isoforms

Total RNA from different human tissues was ordered from Stratagene (La Jolla, CA, USA) except for retina total RNA, which was
Clinical Descriptions

Family W07-713 consists of eight males with severe to profound mental retardation (Figure 1A). Family members IV-5, IV-6, and IV-10 also have postaxial polydactyly in all four limbs. The deceased males IV-2 and IV-4 shared this feature. Retinitis pigmentosa (RP) was noticed in family members III-9, III-10, and IV-10. All patients had recurrent infections, of which family members IV-2, IV-3, and IV-4 died. None of the patients were obese, and all had normal genitalia. No abnormalities were noticed by renal ultrasound scanning. None of the female family members had any symptoms similar to those of the male patients, indicating that the inheritance pattern is most likely X-linked recessive.

Patients IV-10 and III-9 were investigated in more detail. Patient IV-10 was born at term by a lower-segment cesarean section. His birth weight was 3050 g (50th percentile), his length was 48.5 cm (25th percentile), and his head circumference was 34.5 cm (25th percentile). Soon after birth, he presented with stridor and intermittent cyanosis and was thought to have central apnea. He also had feeding problems that required tube feeding. A delay in development was apparent: he was not able to sit or stand at the age of 3.5 years, although he could roll over at 6 months of age. All growth parameters were below –2 standard deviations (SD). The patient had apparent postaxial polydactyly of all four limbs, although central polydactyly could not be excluded (Figure 2B). He presented with mild hirsutism, had low-set ears, a broad nasal bridge, prominent philtrum and maxillary arch, and full lips (Figure 2A). Fingers and nails were normal, as were his genitalia. He could not speak, was unable to grasp or reach for objects, and responded minimally to eye contact or sounds. He was easily irritated and had frequent temper tantrums during which he bit his tongue and lips. In addition, every two weeks he had unexplained fevers that could be treated by antipyretics. There were no serious infections that required hospital admission. The patient

Results

Clinical Descriptions

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had retinal degeneration pointing to juvenile RP. He also had conductive hearing impairment that might be explained by the bilateral middle-ear effusion. Routine blood analysis showed no aberrations in hematologic values, liver function, or renal function; a renal sonography and echocardiogram were also normal. A brain MRI showed a molar tooth sign, hypoplasia of the cerebellar vermis, thickening of the superior cerebellar peduncles, and deep interpeduncular fossa (Figure 2C). Furthermore, patient IV-10 had a small pons and prominent medulla (Figure 2D).

Patient III-9 had severe mental retardation and was unable to care for himself. Breathing problems in early childhood were not noticed. He had low-set ears, a broad nasal bridge, a prominent philtrum and maxillary arch, and full lips, but no postaxial polydactyly (Figures 2E and 2F). He had recurrent respiratory infections during early childhood. By the age of 34, his vision was deteriorated to such an extent that he could only see shadows. Renal sonography showed no aberrations, but from the age of 31 onward, he had hypertension. An MRI scan of the brain also showed a molar tooth sign (Figure 2G), although it was more subtle than that of patient IV-10. Hypoplasia of the cerebellar vermis and thick superior cerebellar peduncles, as well as a cavum velum interpositum and mild volume loss in the cerebral and cerebellar hemispheres, were seen on a brain MRI, suggesting atrophy. The pons and medulla appeared normal in this patient (Figure 2H).

Patient UW87 is part of a cohort of 84 male JS patients who were screened for mutations after the identification of OFD1 as a JS gene. He was born to healthy nonconsanguineous parents at 41 weeks gestation by cesarean section for failure to progress, after a pregnancy complicated by polyhydramnios. His birth weight was 4090 g, his length was 53 cm (90th percentile), and his head circumference was 35.5 cm (50th percentile). As a newborn, he was noted to have tongue thrusting, distinctive craniofacial features, and postaxial polydactyly of both hands and the left foot. Brain MRIs performed at 3 weeks and 8 years of age demonstrated a molar tooth sign (Figure 2K), a small occipital encephalocele, and shortening of the corpus callosum (Figure 2L). A history of repeated middle-ear infections in early childhood required pressure equalization tubes, but no serious infections were reported. A gastrostomy tube was required because of feeding difficulties. On evaluation at 12 years of age, severe intellectual disability, ambulation difficulties, and hyperphagia with obesity were noted. He used a wheelchair and walker for mobility, lacked discernable words, and was not toilet trained. On examination, he had macrocephaly (head circumference >97th percentile) and generalized obesity (97.7 kg, >97th percentile), and his height could not be measured because of combativeness and stooping (Figures 2I and 2J). He had deep-set eyes, narrowed palpebral fissures, a left-sided preauricular pit, a flattened midface, a midline upper-lip notch, a deep midline groove of the tongue, a prominent jaw, and redundant neck tissue. His extremities were notable for relative brachydactyly, fetal finger pads, small feet (US size 5 shoes), and lateral deviation and contractures at the knees. He had normal male genitalia and Tanner stage 3-4 pubic hair. On neurological examination,
he had mild peripheral hypotonia, reduced tendon jerks, and mild dysmetria. Renal ultrasounds were normal, and serum transaminases were mildly elevated on at least two occasions. Ophthalmological examination was challenging, and apparent optic nerve atrophy was noted, but no pigment changes were seen.

**Linkage Analysis**

We performed linkage analysis with 16 markers on the X chromosome (Table 1). A maximum two-point LOD score of 2.07 was obtained for marker DXS8022. It was possible to exclude the remainder of the X chromosome from linkage (LOD score < -2). The interval is delimited by the pseudoautosomal region on the telomeric side and by DXS8036 on the centromeric side (Figure 1A). The 14 Mb interval contains 66 annotated genes (NCBI Map Viewer build 36.3), four of which, OFD1, NLGN4X, HCCS, and PRPS2, are included in the ciliary proteome database and thus might encode ciliary proteins. To our knowledge, ciliary localization has only been experimentally proven for the protein encoded by OFD1. Mutations in NLGN4X are involved in autism and/or mental retardation (MIM 300495), and those in HCCS are involved in microphthalmia and linear skin defects (MIM 309801). Mutations in OFD1 cause OFD1 syndrome and SGBS2. Mutations in PRPS2 have not been reported.

**Table 1. Two-Point LOD Scores for 16 X-Chromosomal Markers**

<table>
<thead>
<tr>
<th>Marker</th>
<th>Position (cM)</th>
<th>0 = 0</th>
<th>0 = 0.05</th>
<th>0 = 0.1</th>
<th>0 = 0.15</th>
<th>0 = 0.2</th>
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<td>DXS8022</td>
<td>27.46</td>
<td>2.07*</td>
<td>1.88</td>
<td>1.68</td>
<td>1.47</td>
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<td>0.51</td>
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<td>0.48</td>
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<td>0.47</td>
</tr>
<tr>
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<td>-0.02</td>
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<tr>
<td>DXS9907</td>
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<td>-0.75</td>
<td>-0.50</td>
<td>-0.35</td>
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<td>-0.64</td>
<td>-0.36</td>
<td>-0.18</td>
</tr>
<tr>
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<tr>
<td>DXS8092</td>
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<td>DXS990</td>
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<tr>
<td>DXS8096</td>
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<td>-0.89</td>
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<td>-1.19</td>
<td>-0.65</td>
<td>-0.36</td>
<td>-0.18</td>
</tr>
</tbody>
</table>

Significant LOD scores are marked with an asterisk. The genetic distances are given according to the Marshfield map.

we first analyzed OFD1 in family member III-10. Mutation analysis of all exons and splice sites of OFD1 showed a deletion of seven base pairs in exon 21: c.2841_2847 delAAAA GAC (p.K948NfsX8) (Figure 1B). The deletion cosegregated with the disease in the family as shown by direct DNA sequencing in all family members and was not present in 190 male controls as shown by ARMS, nor in an additional 250 control chromosomes as shown by direct sequencing (data not shown). Analysis of all OFD1 exons and splice sites in 65 males with JS and of exons 16–23 in 19 additional male JS patients revealed one de novo mutation, c.2767 delG (p.E923KfsX3), in exon 21 in patient UW87 (Figure 1C). This mutation was not present in 250 control chromosomes as shown by direct sequencing (data not shown) and supports the involvement of OFD1 in JS. Because OFD1 escapes X inactivation45, X inactivation should not be skewed in obligate carrier females. Indeed, in carrier females III-2, III-4, and III-8, there is no preferential expression of either X chromosome (data not shown). The deletion causes a frameshift in the OFD1 protein, resulting in a premature termination codon eight amino acids after I947. This termination codon should, in theory, give rise to nonsense-mediated decay of the OFD1 mRNA. QPCR analysis of OFD1 mRNA levels in EBV-LCLs from family member III-9 and isolated patient UW87 showed that 30% and 58%, respectively, of OFD1 expression remained (Figure S1).

**OFD1 Interacts with Lebercilin**

OFD1 was identified as an interacting partner of the LCA5-encoded protein lebercilin by a GAL4-based yeast two-hybrid screen of a retinal cDNA library: the first 305 N-terminal amino acids of lebercilin were used as a bait. Two independent prey clones that encoded largely overlapping OFD1 fragments, amino acid residues 356–1012 and 240–1012, respectively, were identified. The fragments interacting with lebercilin cover five of the six predicted coiled-coil regions of OFD1 and stretch as far as the C-terminus (Figure 3A).46 Apart from the full-length OFD1 protein of 1012 amino acids (OFD1FL), there exists an alternative splice variant that lacks exon 10 and thus encodes a protein (OFD1Δex10) lacking the corresponding 40 amino acids encoded by this exon. Both OFD1 splice variants appear to be of similar biological importance because mRNA of both isoforms is equally expressed in several human tissues and protein expression of both isoforms is detected in bovine retinal extracts (Figure S2).

The lebercilin-OFD1 interaction was confirmed in a dedicated yeast two-hybrid assay, in which both OFD1 fragments (amino acid residues 356–1012 and 240–1012) were coexpressed with full-length lebercilin. In order to determine which coiled-coil regions of lebercilin are responsible for OFD1 binding, we coexpressed different combinations of lebercilin deletion constructs fused to GAL4-BD with OFD1 fused to GAL4-AD in yeast, and we assessed protein-protein interactions by monitoring reporter gene activation (Figure 3A). The region containing...
Figure 3. OFD1 Interaction with Lebercilin

(A) OFD1 and lebercilin protein structures. The protein domains of the two OFD1 isoforms (OFD1FL and OFD1Δex10) and the OFD1 fragments identified in the yeast two-hybrid screen are depicted, along with the protein domains of the lebercilin fragments expressed by the deletion constructs that were tested for interaction.

(B) Yeast two-hybrid assays confirmed binding of both OFD1 fragments, spanning amino acid residues 240–1012 and 356–1012, to full-length lebercilin and the region containing the first two predicted coiled-coil domains of lebercilin. OFD1 fragments containing coiled-coil domains two/three and four (amino acids 240–614) and coiled-coil domains five and six (amino acids 615–1012) both interacted with full-length lebercilin and with the lebercilin fragment spanning amino acids 96–305, although the OFD1 fragment containing coiled-coil domains two/three and four showed the highest affinity. NA = not analyzed; activation of the HIS3 and ADE2 reporter genes is qualitatively indicated by – (no growth on selective media), +/– (slow growth), + (growth), or ++ (fast growth). The activation of the MEL1 (α-galactosidase activity) and the LacZ (β-galactosidase activity) reporter genes was in line with these results.
the first two coiled-coil domains of lebercilin (amino acid residues 96–305, lebercilin\(^{CC1-2}\)) was identified as the region with the highest OFD1 binding affinity on the basis of growth on selective media (activation of the HIS3 and ADE2 reporters) and both \(\alpha\) and \(\beta\)-galactosidase activity assays (Figure 3B).

Similarly, OFD1 constructs encoding coiled-coil domains two/three and four (amino acid residues 240–614 \(\Delta\)ex10 (\[\Delta\text{aa313–353}\]) and coiled-coil domains five and six (amino acid residues 615–1012) were cloned from the OFD1\(^{\Delta\text{ex10}}\) splice variant and tested in a yeast two-hybrid assay (Figure 3A). Both regions of OFD1 interacted with full-length lebercilin and with amino acid residues 96–305 of lebercilin, but the binding affinity of coiled-coil domains two/three and four was the highest (Figure 3B).

To validate the interaction between OFD1 and lebercilin, we performed both GST pull-down and communoprecipitation assays. For the GST pull-down assay, both OFD1 variants (OFD1\(^{\Delta\text{ex10}}\) and OFD1\(^{\Delta\text{ex10}}\)), as well as the OFD1 fragments encompassing amino acids 240–1012 (OFD1\(^{\Delta\text{ex10}–1012}\)) and 356–1012 (OFD1\(^{\Delta\text{ex10}–1012}\)), were expressed as GST fusion proteins. All of these fragments pulled down 3xFLAG-tagged full-length lebercilin from COS-1 cell lysates (Figure 3C). In a control experiment for specificity, unfused GST failed to pull down lebercilin completely. To assess communoprecipitation of OFD1 with lebercilin, we expressed HA-tagged lebercilin in HEK293T cells together with the previously mentioned OFD1 splice variants and fragments fused to a 3xFLAG-tag. Upon immunoprecipitation of HA-lebercilin via anti-HA beads, all 3xFLAG-OFD1 protein variants and fragments coprecipitated (Figure 3D), although 3xFLAG-OFD1\(^{\Delta\text{ex10}}\) and 3xFLAG-OFD1\(^{\Delta\text{ex10}–1012}\) apparently coprecipitated less efficiently. The unrelated 3xFLAG-LRRK2 protein did not coprecipitate with HA-lebercilin; this finding confirmed specificity of the interaction between lebercilin and OFD1 in the communoprecipitation assay (Figure 3D).

The communoprecipitation experiment was repeated with anti-FLAG beads in the immunoprecipitation step. Again, coprecipitation of HA-lebercilin with 3xFLAG-OFD1 was observed. In this experimental set up, 3xFLAG-OFD1\(^{\Delta\text{ex10}}\) and 3xFLAG-OFD1\(^{\Delta\text{ex10}–1012}\) bound more efficiently; therefore, no conclusions on the relative interaction efficiencies of the OFD1 fragments can be drawn. To provide additional evidence for the OFD1-lebercilin interaction, we performed communoprecipitation assays in which the 3xFLAG and HA tags for OFD1 and lebercilin were swapped. These experiments corroborated our previous findings: again, coprecipitation of both proteins could be observed (Figure S3).

### Colocalization of OFD1 with Lebercilin

To provide in vivo support for the in vitro OFD1-lebercilin interaction, we performed colocalization experiments in ciliated hTERT-RPE1 cells and rat retina. GFP-OFD1 was previously reported to be targeted to the centrosome because it colocalized with the centrosomal marker \(\gamma\)-tubulin in HEK293 cells. Lebercilin localizes at the basal body and primary cilium, but association with microtubules has been observed as well. We found that upon expression in hTERT-RPE1 cells, eCFP-OFD1\(^{\Delta\text{ex10}}\) and eCFP-OFD1\(^{\Delta\text{ex10}}\) showed a characteristic pericentriolar localization concentrated around the basal body, which was sostained with the basal-body and cilium marker \(\alpha\)-glaumatylated tubulin (GT335) (Figures 4A–4F). A similar localization has also been reported for CEP290 and PCM-1. This characteristic localization did not depend on the presence of exon 10 in the OFD1 construct: both splice variants showed similar pericentriolar localization. When we coexpressed eCFP-OFD1\(^{\Delta\text{ex10}}\) and mRFP-lebercilin in hTERT-RPE1 cells, partial colocalization was observed (Figures 4G–4I). In rat photoreceptor cells, both OFD1 and lebercilin localize to the connecting cilium region, which is essentially the ciliary transition zone (Figures 4J–4O). This specific connecting cilium localization was evident from coexisting of both proteins with the cilium marker centrin.

### OFD1 Mutations Reduce Interaction with Lebercilin and Change Ciliary Localization

To explain the phenotypic spectrum observed for OFD1 mutations, we tested whether six OFD1 mutations, p.K948fs and p.E923fs (XL-JS; this report), p.I784fs (OFD1\(^{27}\)), p.E709fs (SGBS\(^{28}\)), p.N630fs (OFD1\(^{38}\)), and p.S86fs (OFD1\(^{27}\)), affect interaction with lebercilin and/or subcellular localization. In a yeast two-hybrid interaction assay, the p.I784fs, p.N630fs, and p.S86fs mutations completely disrupted interaction of OFD1 with the fragment containing the predicted coiled-coil domains 1 and 2 of lebercilin, as evidenced by the fact that none of the reporter genes for interaction were activated (Figure 5A). For the p.E923fs mutant, yeast growth was significantly reduced in comparison to wild-type OFD1 under conditions selecting for interaction, indicating a reduced binding affinity. The p.K948fs mutation only slightly

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(C) Both GST-OFD1\(^{\Delta\text{ex10}}\) and the GST-fused OFD1 fragments spanning amino acid residues 240–1012 and 356–1012 (10% input of the assay shown in panel 3) efficiently pulled down 3xFLAG-tagged full-length lebercilin (10% input shown in panel 1; pull-down assay shown in panel 2). Unfused GST failed to pull down lebercilin at all, whereas, in a positive control experiment, GST-neprhocrystin-4-I pulled down RPRGIP1\(^{C2-end}\).

(D) The immunoblot (panel 4) shows that 3xFLAG-tagged OFD1 fragments communoprecipitated with HA-lebercilin, whereas the unrelated 3xFLAG-LRRK2 protein did not. Five percent of the cell lysate inputs used in the assays are shown in panels 1 and 2, and the HA immunoprecipitates are shown in panel 3. As a positive control, 3xFLAG-RPRGIP1\(^{C2-end}\) communoprecipitated with HA-fused nephrocystin-4-I.

(E) When anti-FLAG immunoprecipitation was performed, HA-lebercilin communoprecipitated with all 3xFLAG-tagged OFD1 fragments, as shown in the immunoblot in panel 4. Panels 1 and 2 show five percent of the cell lysate inputs used in the assays, and panel 3 depicts FLAG immunoprecipitates.
reduced yeast growth, whereas the p.E709fs mutation did not apparently affect yeast growth. For the mutants that retained interaction in the yeast two-hybrid assay, we also assessed the effects on lebercilin binding in a semi-quantitative liquid β-galactosidase assay, demonstrating activation of the LacZ reporter gene (Figure 5B). This assay showed that the p.E709fs mutation moderately decreased OFD1 binding to lebercilin (60% residual reporter gene activity compared to that of wild-type OFD1). The p.K948fs and p.E923fs mutations more strongly decreased OFD1 binding to lebercilin (24% and 18% residual reporter gene activity, respectively).

We also investigated to what extent OFD1 mutations influenced cellular localization of the encoded eCFP-fusion protein. In hTERT-RPE1 cells, the JS-associated p.K948fs and p.E923fs mutants (Figures 6A–6F) and the SGBS2-associated p.E709fs mutant (Figures 6J–6L) retained pericentriolar localization, comparable to that of wild-type
In contrast, the OFD1 syndrome-associated p.N630fs and p.S586fs mutant proteins localized diffusely throughout the cytoplasm and showed only a limited, severely reduced concentration around the basal body (Figures 6M–6R). The p.I784fs mutant protein, which is also associated with OFD1 syndrome, showed comparable scattering throughout the cytoplasm but showed clear concentration at both centrioles (Figures 6G–6I). This localization obviously differed from the pericentriolar localization observed for wild-type OFD1 and for the XL-JS and SGBS2-associated mutant proteins.

Discussion

Until now, Joubert syndrome has been recognized as an autosomal-recessive disorder. In this study, we report that in two unrelated families with X-linked recessive JS...
as defined by the presence of the molar tooth sign), exon 21 of OFD1 contains two separate protein-truncating mutations. The mutations cosegregated with the disease and were not found in more than 250 controls. Affected males primarily presented with mental retardation accompanied by postaxial polydactyly. Central apnea was noted in at least one family member of the XL-JS family and in isolated patient UW87. In the XL-JS family, early-onset retinal degeneration was reported, showing that OFD1 mutations can cause classic JS with retinal involvement. The facial dysmorphism, oral anomalies, and overgrowth in patient UW87 reveal phenotypic overlap between OFD1-related JS, SGBS2, and OFD1 syndrome.

The OFD1 gene has previously been associated with OFD1 syndrome, which is a male-lethal X-linked dominant condition primarily involving malformations of the face, oral cavity, and digits (Table 2). Cystic kidney disease is also commonly associated with this disorder, as are central nervous system abnormalities. However, a molar tooth sign on a brain MRI has never been reported. OFD1 syndrome belongs to the broad spectrum of cilia-related disorders or ciliopathies; the OFD1 protein was found to localize to centrosomes and basal bodies in renal epithelial cells. Mice lacking Ofd1 reproduced the main features of the human disease and showed several hallmarks of ciliary dysfunction, such as failure of left-right axis specification, polydactyly, and renal cyst development. Mutations in OFD1 have also been found in a single family presenting with X-linked recessive mental retardation, macrocephaly, and respiratory ciliary dyskinesia (SGBS2; Table 2). The index patient in this study, who was the only living affected individual in the family, showed no molar tooth sign on a brain MRI. However, no brain-scan data were available for the other, more severely affected patients, who died soon after birth, so it can not be completely excluded that SGBS2 belongs to the spectrum of JSRDs.

The two XL-JSRD mutations reported here lead to a frameshift that results in a premature termination codon.
in the OFD1 mRNA; this in turn renders the mutant mRNA subject to nonsense-mediated decay. Consequently, only about 30%–60% of OFD1 transcript persists. This is also the case for the SGBS2 mutation (Figure S1). Most OFD1-syndrome-associated mutations also result in premature termination codons and are thus predicted to reduce transcript levels as well. Therefore, we postulate that the phenotypic variability observed for OFD1 mutations is primarily caused by changes in activity of the remaining truncated OFD1 protein (Figure 7). Overall, the severity of the phenotype correlates with a reduction in OFD1 protein length. All mutations before amino acid residue 631 are lethal for males and cause OFD1 syndrome in females. The SGBS2-associated p.E709fs mutation causes macroencephaly and early death in males because of severe respiratory-tract infections, but it leaves females unaffected. Male patients with the p.K948fs and p.E923fs mutations, which are located in the coiled-coil domain nearest to the C-terminal end of the protein, have JS and may live beyond the age of 30 years, whereas carrier females

Table 2. Clinical Features of the XL-JS Family and Isolated JS Patient UW87 as Compared to SGBS2 and Females with OFD1 Syndrome

<table>
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<tr>
<th>Clinical Symptom</th>
<th>XL-JS</th>
<th>XL-JS (UW87)</th>
<th>SGBS2&lt;sup&gt;28&lt;/sup&gt;</th>
<th>OFD1&lt;sup&gt;28&lt;/sup&gt;</th>
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<td>recessive</td>
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<td>recessive</td>
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<td>Recurrent infections</td>
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<td>-</td>
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<tr>
<td>Juvenile RP</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Cystic kidneys</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Obesity</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Typical age of male death</td>
<td>&gt;20 yrs</td>
<td>&gt;12 yrs</td>
<td>&lt;3 yrs</td>
<td>Prenatal</td>
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Mental retardation severe/profound severe severe mild
| Hypotonia | - | + | + | - |
| Central apnea | + | + | - | - |
| Molar tooth sign | + | + | - | - |
| Corpus callosum agenesis | - | - | - | + |

Dysmorphisms
| Macrocephaly | - | + | + | - |
| High-arched palate | - | - | + | - |
| Low-set ears | + | - | + | - |
| Broad nasal bridge | + | - | - | + |
| Hypotelorism | - | - | - | + |
| Hypoplasia of the nasal alae | - | +/- | - | + |
| Downslanting palpebral fissures | - | - | - | + |

Oral Abnormalities
| Cleft lip/palate | - | +/-<sup>**</sup> | - | + |
| Tongue abnormalities | - | + | - | + |
| Abnormal dentition | - | - | - | + |

Digital Abnormalities
| (Postaxial) polydactyly | + | + | + | +<sup>*</sup> |
| Brachydactyly | - | + | + | + |
| Clinodactyly | - | - | - | + |
| Broad thumbs | - | +/- | - | - |
| Long digits | + | - | - | - |

<sup>*</sup> Postaxial polydactyly has been reported in OFD1 syndrome but occurs rarely (<5%).
<sup>**</sup> Notched lip.
Figure 7. Genotype-Phenotype Correlation of OFD1 Mutations with OFD1, SGBS2, and XL-JS

Mutations thus far reported as occurring before amino acid residue 631 are all embryonic lethal for males. In females, these cause OFD1 syndrome. The SGBS2-associated p.E709fs mutation causes early death in males, but females are not affected. Male patients with the p.E923fs and p.K948fs mutations that cause XL-JS may live beyond the age of 30 years. We have also indicated the position of the exon 17 p.I784fs mutation that results in OFD1 syndrome in females and is lethal to males to show that the phenotype boundaries are variable (dotted lines). However, the overall severity of the phenotype inversely correlates with the size of the remaining OFD1 protein.

are not affected. Interestingly, two exon 17 mutations that are positioned between the SGBS2 and the XL-JS mutations have been described\(^6\): one is a frameshift (p.I784fs), and one is a deletion of the complete exon. These exon 17 mutations are lethal to males and cause OFD1 syndrome in females. This shows that the phenotypic boundaries are variable and that other factors, such as variable degrees of RNA degradation, contribute to the differences in phenotype as well.

We show that OFD1 physically interacts with lebercilin, the protein encoded by the LCA-associated gene LCA5. The functional significance of this interaction is stressed by the fact that the X-linked JS family we report here also suffers from retinal degeneration. For the family members IV-5 and IV-6, as well as for isolated patient UW87, no retinal phenotype could be diagnosed. However, monitoring of these patients is advisable because a retinal pathology could develop in the future. Variability in the onset and severity of the retinal phenotype among and within families is a common feature with isolated RP and even more so in (retinal) ciliopathies, which often express a variable combination of phenotypes. Furthermore, we show that OFD1 is present in retinal photoreceptor cells, where it colocalizes with lebercilin in the transition zone of the photoreceptor sensory cilium. In other cell types, both OFD1 and lebercilin localize to the basal body and/or to primary cilia. Our interaction data show that the p.K948fs, p.E923fs, and p.E709fs mutations in OFD1 weakened the interaction with lebercilin but did not totally disrupt it. OFD1-eCFP fusion proteins containing the p.K948fs, p.E923fs, and p.E709fs mutations showed normal pericentriolar localization in hTERT-RPE1 cells. In contrast, the male-lethal dominant OFD1 p.I784fs, p.N630fs, and p.S586fs mutations completely abolished interaction with lebercilin, and upon expression of the corresponding mutated recombinant proteins in hTERT-RPE1 cells, pericentriolar localization was lost completely; however, apart from diffuse cytoplasmic localization, the p.I784fs mutant protein showed concentration at both centrioles, and such an observation was not made for the p.N630fs and p.S586fs mutant proteins. The p.I784fs mutation lies in exon 17, which is further downstream than the p.N630fs and p.S586fs mutations and results in a longer remaining mutant protein. This mutant protein might retain the ability to concentrate at the centrioles, but it has lost pericentriolar localization as well as lebercilin binding affinity. Taken together, these data support the hypothesis that differences in binding affinity of the OFD1 mutant protein to functionally interacting proteins, such as lebercilin, underlie the phenotypic variability observed for OFD1 syndrome, SGBS2, and XL-JS.

The exact cellular function of OFD1 is not known yet, but evidence from studies with knock-out mice shows that OFD1 is required for primary cilia formation and left-right axis specification.\(^54\) In limb buds of these mice, which consistently showed polydactyly, altered expression of Hoxa and Hoxd genes was observed. These genes regulate limb patterning downstream of Gli3, which is a transcription factor in the Shh pathway.\(^36\) Although Gli3 itself and other players in the Shh pathway showed no altered gene expression in the limb buds of Ofd1-null mice, OFD1 disruption could affect Gli3 function, which would explain the upregulation of Hox genes. In this line of thinking, it is interesting to note that Ofd1 knock-out mice show overall phenotypic overlap with RPGRIP1L-null mice.\(^20,26\) Studies in RPGRIP1L/Ftm-null mice identified RPGRIP1L as a necessary component for cilia-mediated Shh signaling.\(^26\) It is tempting to speculate that OFD1 might also be involved in this pathway; such involvement would explain the occurrence of polydactyly in patients with OFD1 mutations. Furthermore, mutations in the JS gene Arh3b were identified as underlying the lethal hemin mouse mutant that shows disruption of the ciliary axoneme and displays a Shh-like phenotype, including defective neural-tube patterning and polydactyly.\(^13\) These studies suggest that cilia-mediated regulation of Shh signaling might be a commonly disrupted pathway in JS.

In conclusion, we have identified OFD1 as a JSRD gene and have expanded the recognized inheritance pattern of JSRDs to include X-linked recessive. Mutations in OFD1 are now known to be associated with three disorders: OFD1 syndrome, SGBS2, and XL-JS. We propose that the inverse correlation between OFD1 mutant protein length and phenotypic severity could be explained by differences in binding to functionally interacting proteins and disruption of ciliary localization.
Supplemental Data

Supplemental Data include three figures and one table and can be found with this article online at http://www.cell.com/AJHG/.

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Web Resources

The URLs for data presented herein are as follows:

Cilia Proteome, http://www.ciliaproteome.org
Human Genome Browser, http://www.genome.ucsc.edu/
SMART, http://smart.embl-heidelberg.de/smart/set_mode.cgi?NORMAL=1

References

polyporphosphate-5-phosphatase E, link phosphatidyl inositol signaling to the ciliopathies. Nat. Genet. 41, 1032–1036.


