Abnormal glycosylation in Joubert syndrome type 10

Megan S. Kane1,2†, Mariska Davids1,†, Michelle R. Bond3, Christopher J. Adams1, Megan E. Grout4, Ian G. Phelps4, Diana R. O’Day4, Jennifer C. Dempsey4, Xeuli Li5, Gretchen Golas1, Gilbert Vezina6, Meral Gunay-Aygun7,8,9, John A. Hanover3, Dan Doherty4, Miao He5, May Christine V. Malicdan1,*, William A. Gahl1 and Cornelius F. Boerkoel1,10

Abstract

Background: The discovery of disease pathogenesis requires systematic agnostic screening of multiple homeostatic processes that may become deregulated. We illustrate this principle in the evaluation and diagnosis of a 5-year-old boy with Joubert syndrome type 10 (JBTS10). He carried the OFD1 mutation p.Gln886Lysfs*2 (NM_003611.2: c.2656del) and manifested features of Joubert syndrome.

Methods: We integrated exome sequencing, MALDI-TOF mass spectrometry analyses of plasma and cultured dermal fibroblasts glycomes, and full clinical evaluation of the proband. Analyses of cilia formation and lectin staining were performed by immunofluorescence. Measurement of cellular nucleotide sugar levels was performed with high-performance anion-exchange chromatography with pulsed amperometric detection. Statistical analyses utilized the Student’s and Fisher’s exact tests.

Results: Glycome analyses of plasma and cultured dermal fibroblasts identified abnormal N- and O-linked glycosylation profiles. These findings replicated in two unrelated males with OFD1 mutations. Cultured fibroblasts from affected individuals had a defect in ciliogenesis. The proband’s fibroblasts also had an abnormally elevated nuclear sialylation signature and increased total cellular levels of CMP-sialic acid. Ciliogenesis and each glycosylation anomaly were rescued by expression of wild-type OFD1.

Conclusions: The rescue of ciliogenesis and glycosylation upon reintroduction of WT OFD1 suggests that both contribute to the pathogenesis of JBTS10.

Keywords: Joubert syndrome, Ciliopathy, Glycosylation, Molar tooth sign, CMP-sialic acid

Background

Joubert syndrome (JBTS) is a rare genetic condition characterized by hypotonia, developmental delay, cerebellar dysfunction, a neuroradiologic “molar tooth sign” (MTS), and variable involvement of other systems [1, 2]. JBTS is a ciliopathy, and mutations in more than 30 genes encoding products localizing primarily to and around the cilia are implicated in its pathogenesis [3, 4]. The genetic locus associated with JBTS type 10 (OMIM 300804) is OFD1; mutations in this gene also cause the X-linked dominant disorder orofaciodigital syndrome I (OFDI, OMIM 311200), the X-linked recessive disorder Simpson–Golabi–Behmel syndrome, type 2 (SGBS2, OMIM 300209), and an X-linked form of retinitis pigmentosa (RP23, OMIM 300424) [5].

The ciliary function of the OFD1 protein has not been fully elucidated, although it does play a role in primary ciliogenesis and regulation of ciliary length [6, 7]. Loss of OFD1 in animals causes defects in left/right axis development, ciliogenesis, and neurological development [8–10]. These phenotypes are consistent with features

*Correspondence: megan.kane@nih.gov; maychristine.malicdan@nih.gov
†Megan S. Kane and Mariska Davids contributed equally to this work
1 NIH Undiagnosed Disease Program, Common Fund, Office of the Director, and National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA
Full list of author information is available at the end of the article

© The Author(s) 2017. This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
observed in other ciliopathy disorders and reflect the role of primary cilia in the Shh, Wnt, and planar cell polarity (PCP) signaling pathways [4]. In the centrosome, OFD1 interacts with lebercilin, a key protein in the pathogenesis of Leber congenital amaurosis [11, 12]. Based upon its nuclear localization, there is a hypothesis that OFD1 plays a role in chromatin remodeling [13].

The National Institutes of Health Undiagnosed Disease Program (NIH UDP) evaluates individuals with rare and undiagnosed diseases and an indication of a genetic etiology. Physicians across multiple specialties consult on each patient to obtain a full phenotypic description. In addition to targeted diagnostic tests, the NIH UDP implements broader, agnostic screens of exome, glycome, and metabolome.

Herein we report the evaluation of a 5-year-old boy with Joubert syndrome secondary to mutation in $OFD1$. His cultured dermal fibroblasts had impaired ciliogenesis, and his plasma and cultured dermal fibroblasts had abnormal $N$- and $O$-linked glycosylation profiles. Glycomic evaluation of two unrelated individuals with JBTS10 detected similar plasma and fibroblast $N$- and $O$-linked glycosylation profiles. Cultured fibroblasts from the proband also had increased CMP-sialic acid. Overexpression of wild-type $OFD1$ cDNA rescued both the glycosylation and ciliary phenotypes. We posit, therefore, that glycosylation anomalies are a previously undocumented contributor to the pathogenesis of JBTS10.

**Methods**

**Patient enrollment and consent**
The proband, UDP-3331, was evaluated in the National Institutes of Health Undiagnosed Diseases Program (NIH UDP) and was enrolled in clinical protocol 76-HG-0238, approved by the NHGRI Institutional Review Board. His mother, UDP-4786, provided informed consent. NIH-452 was enrolled in protocol “Clinical and Molecular Investigations into Ciliopathies” (Trial NCT00068224); his legal guardian provided informed consent. Both protocols conform to the 1964 Helsinki Declaration and subsequent amendments standard for patient protection.

**Variant identification and Sanger sequencing**
Genomic DNA was extracted from peripheral whole blood of UDP-3331 and UDP-4786 using the Gentra Puregene Blood Kit (Qiagen, Valencia, CA) and an AutoGen FlexStar with standard procedures. The DNA of all family members was subjected to an integrated set of genomic analyses including high-density single-nucleotide polymorphism (SNP) arrays and whole exome sequencing (WES) as described previously [14–17]. WES was performed on these individuals using the Illumina HiSeq2000 platform and the TrueSeqV2 capture kit (Illumina, San Diego, CA). Sample library preparation, sequencing, and analysis were performed using the standard NIH Intramural Sequencing Center (NISC) pipeline [18]. Sequence data were aligned to human reference genome (hg19) using Novoalign (Novocraft Technologies, Selangor, Malaysia). To test for copy number variants and to form segregation BED files for exome analysis, Omni Express 12 (hg19) SNP arrays were run on genomic DNA from all family members as described [14]. Variants listed in the Variant Call Files (VCFs) were filtered based on rarity, Mendelian segregation, and predicted deleteriousness on the Cartagenia platform (Cartagenia N.V., Leuven, Belgium).

Exome variants were filtered using ExAC release 0.3 population frequency data (MAF < 0.02, 95% confidence interval; homozygote count ≤ 25) and UDP founders cohort population frequency data (variant allele count < 8). Variants were prioritized based on coding effect (nonsynonymous, frameshift, stopgain, stoploss, startloss, inframe), proximity to splice sites (within 20 base pairs of a canonical splice site into the intron, or 5 base pairs into the exon), and CADD v1.3 Phred scores. Using the Integrative Genome Viewer (https://www.broadinstitute.org/igv/home), we assessed the quality of alignment and genotype call of variants. The single-nucleotide deletion identified by exome sequencing was confirmed using Sanger sequencing of the genomic DNA from both UDP-3331 and UDP-4786; the forward primer used was 5′-AGTAGGTCAGGTCAAGGAGTGG-3′ and the reverse primer was 5′-ATCTGTGACGCTTTCACAC-3′. DNA amplification was performed with Platinum Pfx DNA polymerase (ThermoFisher Scientific, Waltham, Massachusetts) and standard thermal cycling protocol with an annealing temperature of 55 °C for a total of 30 cycles.

**OFD1 cloning**
Wild-type (WT) cDNA expression constructs were made by PCR amplification of the $OFD1$ cDNA sequence from a sequence-verified TrueORF Gold clone (RC220154, Origene, Rockville, Maryland). The PCR product was cloned into pENTR/D-TOPO vector (ThermoFisher Scientific) and subcloned into pLENTI6.3/V5-DEST (ThermoFisher Scientific) using the Gateway LR Clonase II enzyme (ThermoFisher Scientific). Clones were verified by Sanger sequencing.

**Lentiviral infection and rescue**
Lentivirus was generated by transfecting 293FT cells with the pLENTI6.3-OFD1 construct and the ViralPower HiPerform Lentiviral Gateway expression kit (ThermoFisher Scientific) as per the manufacturer’s protocol.
Lentiviral culture supernatant was applied to patient and adult WT control cells in the presence of polybrene (10 µg/mL, EMD Millipore, Billerica, Massachusetts). Stable integration of the OFD1 construct was selected by addition of 5 µg/mL blasticidin (ThermoFisher Scientific) to the medium and confirmed by qPCR and immunoblotting. Subsequently, retention of the integrated active OFD1 construct was selected by growth in medium supplemented with 2.5 µg/mL blasticidin.

**Primary fibroblast isolation and culture**

Primary dermal fibroblasts from UDP-3331 were derived from a forearm skin punch biopsy. Additional fibroblast lines were obtained from ATCC (ATCC-PCS-201-012, Manassas, Virginia) and Coriell Repository [GM00409 (7 y.o. male) and GM08398 (8 y.o. male), Camden, New Jersey]. The established fibroblast lines were cultured in growth medium, consisting of Dulbecco’s Modified Eagle Medium (DMEM) with 1 g/L glucose (low glucose DMEM), 10% heat inactivated fetal bovine serum (HI FBS, ThermoFisher Scientific), and 1% antibiotic/antimycotic (ThermoFisher Scientific). Culture of primary fibroblasts for glycome analysis utilized α-MEM (Cellgro, Mediatech, Manassas, Virginia) supplemented with 15% HI FBS and 1% antibiotic/antimycotic for 24–48 h prior to cell harvest.

**RNA extraction and qRT-PCR analysis**

RNA extraction from cultured dermal fibroblasts was performed as per the manufacturer’s protocol using the Qiagen RNeasy Mini kit (Qiagen, Hilden, Germany) with on-column DNA digestion. Equal amounts of RNA were reverse-transcribed from each cell line using the OmniScript cDNA first-strand synthesis kit (Qiagen). SYBR Green reagent (Qiagen) was used to amplify OFD1 and GAPDH for a ΔΔCt relative quantification of OFD1 expression on an ABI7500 Fast RealTime PCR machine with the Melt Curve option. Normalization and quality control parameters were according to the standard manufacturer’s settings. Primers used for OFD1 were 5′-CGGAGCAGAAAGTGGGTCTTT-3′ and 5′-TGGCATGTCCCCCTGACATT-3′; primers for GAPDH were 5′-TGCACCACCAACTGCTTAGC-3′ and 5′-TGGCATGTTCCCTGCAGATT-3′; primers for GAPDH were 5′-TGCACCACCAACTGCTTAGC-3′ and 5′-TGGCATGTTCCCTGCAGATT-3′.

**Immunoblotting**

Fibroblasts were cultured to near 100% confluence, rinsed twice with PBS, and harvested by in-dish lysis with RIPA buffer (Sigma-Aldrich, St. Louis, Missouri) supplemented with 1× complete Ultra Protease Inhibitor Cocktail (Sigma-Aldrich). Total protein concentration of the lysates was determined using the DC Protein Assay (Bio-Rad, Hercules, California) and equal amounts of protein were run for each sample on SDS-PAGE gels (4–12% gradient, BioRad) in SDS–Tris–glycine buffer. Proteins were transferred using a standard wet-transfer method to Immobilon-FL PVDF membranes in Tris–glycine buffer with 20% methanol. Blocking was performed with Odyssey blocking buffer in PBS (LiCor, Lincoln, Nebraska); primary antibodies were incubated overnight at 4 °C in fresh blocking solution. Membranes were washed three times with PBS-Tween 20 (0.1%) and incubated with Odyssey near-IR conjugated secondary antibodies for 1 h in Odyssey blocking buffer with Tween-20 (0.1%) and SDS (0.2%). Prior to imaging on a LiCor Odyssey CLx, membranes were washed three times with PBS-Tween 20 and rinsed once with diH2O. Band quantification and analysis were performed using the CLx imaging software (LiCor). Antibodies used were anti-OFD1 (1:1000, TA308968, Origene), anti-α-tubulin (1:5000, ab7291, Abcam; Cambridge, Massachusetts), anti-rabbit IR800 (1:10,000, 925–32,211, LiCor), and anti-mouse IR680 (1:10,000, 925–68,020, LiCor).

**Immunofluorescence of cultured fibroblasts**

Staining of cultured fibroblasts, either with antibodies or lectins, utilized a similar protocol with minor variations detailed below. Cultured cells were fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS) at room temperature for 10 min, or in ice-cold 100% methanol at −20 °C for 15 min, or a combination of both. The fixed cells were washed three times with PBS. Paraformaldehyde-fixed cells were permeabilized with 0.5% NP-40 detergent in PBS at room temperature for 10 min then washed three times in PBS. The fixed and permeabilized cells were then incubated with a blocking buffer for 1 h. Primary staining reagent(s) were diluted in blocking buffer and applied for 1–2 h at room temperature or overnight at 4 °C. The cells were then washed with PBS three times. The cells were then incubated with secondary detection reagents with fluorophore conjugates diluted in blocking buffer for 1 h at room temperature. After washing the cells three times with PBS, the coverslips were mounted onto slides with the indicated mounting media prior to imaging.

**OFD1 localization and cell morphology analysis**

Cells were grown on 12-mm round coverslips to 50–70% confluence at the time of fixation. Cells were fixed either in paraformaldehyde or methanol, depending upon manufacturers’ recommendations for the primary antibodies. The blocking reagent used was 4% BSA (bovine serum albumin) in PBS, and the primary antibodies used were OFD1 (1:100, Origene), γ-Tubulin (1:1000; T5326, Sigma-Aldrich), α-tubulin (1:2000, ab7291, Abcam), PCM1 (1:100, sc-50164, SantaCruz Biotechnology, Dallas, Texas).
Texas), GM-130 (1:250, BD Biosciences, San Jose, California), TGN-46 (1:200, GTX74290, Genetex, Irvine, California), or KDEL (1:100, ab12223, Abcam). Alexa Fluor conjugated secondary antibodies used were Alexa488 Donkey anti-Rabbit, Alexa555 Donkey anti-Mouse, and Alexa647 Donkey anti-Rabbit (ThermoFisher Scientific). Where indicated, DNA was counterstained with Hoechst 33342 (ThermoFisher Scientific) and coverslips were rinsed with PBS. The mounting medium was ProLong Gold antifade mounting medium (ThermoFisher Scientific) and the slides were imaged using a 20× or 40× objective on a Zeiss LSM 700 confocal laser-scanning microscope (Carl Zeiss Microscopy, GmbH, Jena, Germany).

Cilia measurements
Fibroblasts were grown to 80% confluence and then serum starved for 48 h. Cells were fixed with 4% paraformaldehyde for 5 min at room temperature followed by ice-cold methanol for 4 min at −20 °C. The blocking buffer used was PBS containing 10% normal donkey serum, 1% BSA, and 0.1% triton X-100 for 1 h. Fixed cells were incubated overnight with primary antibodies (1:1000–1:2000, mouse anti-acetylated Tubulin, T6793, SigmaAldrich; 1:200, goat anti-γ tubulin, sc-7396, Santa Cruz). Secondary antibodies used were AlexaFluor conjugated secondary antibodies (Life Technologies). Coverslips were mounted using Fluormount G with 4',6-diamidino-2-phenylindole (DAPI) (Southern Bio- tech, 0100-20, Birmingham, Alabama).

Fourteen image z-stacks with 0.3 mm spacing were taken with a CoolSNAP HQ2 digital monochrome camera (Photometrics, Tucson, Arizona, USA) through a Marians live cell imaging system (Intelligent Imaging Innovations, Denver, Colorado, USA) using a Plan Apochromat 63X, 1.4 NA oil objective.

Percentage of ciliated cells and length measurements were made in FIJI on 16-bit sum projection images of z-stacks. Cilia were defined as linear acetylated tubulin-positive structures >1-μm long with a punctate gamma-tubulin signal at one end. The percentage of ciliated cells was determined by dividing the number of ciliated cells by the number of DAPI-positive nuclei in the same images. To measure cilia length, we manually painted a mask over each cilium (defined by acetylated tubulin) with a 3 pixel-wide brush, skeletonized the cilium mask, and applied the maximum branch length function.

Lectin staining of cultured fibroblasts
Biotinylated Maackia amurensis lectin II (MAL II), fluorescein Sambucus nigra (elderberry) bark lectin (SNA), and rhodamine Peanut Agglutinin (PNA) were obtained from Vector Laboratories (Burlingame, California). Cells were seeded at 0.5 × 10^5 per 12-mm round coverslip and cultured overnight. The cells were fixed with paraformaldehyde or methanol as indicated in the text. Blocking was performed with 4% BSA in PBS, followed by primary incubation with MAL II, SNA, and/or PNA lectins (20 μg/mL). Detection of biotinylated MAL II required secondary incubation with Streptavidin AlexaFluor 350 (S-11,249, ThermoFisher Scientific). In cells without MAL II staining, DNA was stained with Hoechst 33342 (ThermoFisher Scientific). Coverslips were mounted on slides using ProLong Gold anti-fade reagent then imaged using a 20× objective on a Zeiss LSM 700 confocal laser-scanning microscope. Image analysis was performed using ImageJ software [19].

Neuraminidase treatment and cilia measurement
Adult WT fibroblasts were plated directly on coverslips at 0.5 × 10^5 cells per 12-mm coverslip density, allowed to adhere overnight, and then grown for additional 48 h in the absence of serum. Half of the coverslips were washed with Hank’s Balanced Salt Solution (HBSS) and then incubated for 1 h at 37 °C in 1 mU/mL neuraminidase (Sigma, N2133) dissolved in HBSS. Untreated cells were left in serum-free media. Cells were subsequently washed three times in PBS and then fixed using ice-cold methanol. Staining for cilia utilized anti-ARL13B (1:250, ProteinTech, Rosemont, Illinois) and anti-gamma tubulin (1:250, Sigma-Alrdich) with Hoechst 33342 DNA counterstaining. DNA counterstaining in conjunction with lectin staining, which was performed as stated above, utilized the far-red dye DRAQ5 (Abcam). Cilia were defined as above and measured with FIJI software; the percentage of ciliated cells was defined as above.

Glycome analysis
Primary dermal fibroblasts were cultured and processed to release N- and O-linked glycans as previously reported [20]. Fibroblasts were cultured in DMEM with 10% HI FBS and 1% antibiotic/antimycotic to 80% confluence in a 15-cm culture dish. After one rinse with PBS, the medium was changed to α-MEM containing 1000 mg/L d-glucose, 15% HI FBS, and 1% antibiotic–antimycotic. Upon reaching 100% confluence, the cells were washed twice with PBS then harvested using a cell scraper. The cells were then pelleted in PBS by centrifugation. Fibroblast pellets were lysed in 300 μL PBS with sonication and 400 μg of total glycoprotein was collected from the cell lysate. Free glycans were removed by Ultrace1-10k Centrifugal filter (EMD Millipore, Billerica, Massachusetts) and the enriched glycoproteins were divided into 200-μg aliquots for N-glycan and O-glycan preparation. N-linked glycans were released from 200 μg of total glycoproteins or 10-μL plasma samples using PNGase F digestion (New
England Biolabs, Ipswich, Massachusetts, USA). The released N-linked glycans from plasma or cells were purified and desalted by solid-phase extraction using a Sep-Pak C18 and carbograph column. O-linked glycans were released using β-elimination with sodium borohydride. O-linked glycans from plasma or cells were purified and desalted using an AG 50 W-X8 resin cation exchange column. N-linked glycans or O-linked glycans were permethylated with sodium hydroxide and iodomethane in dimethyl sulfoxide (DMSO) as described before [21]. Following permethylation, glycans were extracted with water/chloroform (2:1, vol/vol) for four times and dried. Samples were then dissolved in 50% methanol, spotted with 11% 2,5-dihydroxybenzoic acid matrix (1:1 vol/vol), and measured by MALDI-TOF using the positive mode on Ultraflex MALDI-TOF/TOF system (BrukerDaltonics, Billerica, Massachusetts).

Nucleotide sugar analysis
Fibroblasts were cultured in growth medium in 15-cm culture disks until 90–100% confluent, rinsed with PBS and harvested by scraping into PBS. Cells were counted using an automated cell counter (BioRad TC20) and pelleted at 500 RCF for 5 min. Nucleotide sugars were extracted based on previously published methods [22, 23]. Briefly, cell pellets were lysed in 70% ethanol, 300 µL of ethanol per million cells, and subjected to three rounds of sonication for 5 s at 15% amplitude. An aliquot of lysate was taken for total protein quantification. Lysates were pelleted at 16,000×g for 10 min at 4 °C and the supernatant was collected and lyophilized. The lyophilized supernatant was resuspended in 40 mM sodium phosphate buffer (pH 9.2) at 60 µL per million cells and further diluted based on total protein from the initial lysate relative to adult WT. The supernatant was applied to a Millipore spin column (10kD MW cut off) and spun for 30 min at 16,000×g. The extracted nucleotide sugars were analyzed by high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) on a Dionex instrument as previously reported [22] and quantified by comparison to a standard curve of dilutions of CMP-sialic acid (C8271, Sigma-Aldrich) in 40 mM sodium phosphate.

Results
Clinical report
The proband, UDP-3331, is a 5-year-old male, born to unaffected parents; he had an unaffected sister. At 14 months, he presented with severe feeding problems requiring placement of a G-tube. At 18 months, he was evaluated for hypotonia and dysmorphia. Over the next 4 years, his documented features included oral motor dysfunction, sleep apnea, myopathy, developmental delay, easy bruising, hypertrichosis, rocker bottom feet, and dysmorphic facial features, including a notched upper lip and submucous cleft palate. Following 5 years of non-diagnostic evaluations, the proband was evaluated at the NIH UDP. His brain MRI revealed a molar tooth sign (Fig. 1a) that, combined with his clinical features (Table 1), suggested a diagnosis of Joubert syndrome.

Exome sequencing identifies an OFD1 mutation
Analysis of exome sequence data for variants in genes associated with Joubert syndrome revealed NM_003611.2: c.2656del, a maternally inherited hemizygous mutation in OFD1. Sanger sequencing confirmed the variant in the proband and mother (UDP-3331 and UDP-4786, respectively; Fig. 1b) and refined the diagnosis to JBTS10. Considering the apparently normal phenotype of the mother, no further testing or analysis of samples from individual UDP-4786 was performed.

The OFD1 mutant protein is stable and appropriately localized
NM_003611.2: c.2656del, which occurred in the 20th exon, is predicted to encode a frameshift leading to a premature stop codon, p.Gln886Lysfs*2. To test if this mutation caused nonsense-mediated mRNA decay and thereby loss of OFD1 function, we measured OFD1 mRNA levels in cultured dermal fibroblasts by qRT-PCR. Following normalization to GAPDH mRNA abundance, the UDP-3331 OFD1 mRNA levels were approximately 30% of those in control dermal fibroblasts (Fig. 1c), a finding consistent with loss of function and previous reports [11]. OFD1 protein levels, as measured by immunoblotting, were approximately 50% of those in control dermal fibroblasts following normalization to α-tubulin (Fig. 1d). Lentiviral transduction of a WT copy of OFD1 cDNA into the dermal fibroblasts of UDP-3331 partially rescued OFD1 mRNA and protein expression (Fig. 1c, d).

We next determined the intracellular localization of the residual OFD1 protein by immunofluorescent studies of dermal fibroblasts. In both UDP-3331 and control, residual OFD1 localized with the centrosome as judged by co-localization with centrosome markers γ-tubulin and PCMI (Fig. 2a). Notably, in both affected and control fibroblasts, we observed variable co-localization of all three centrosomal proteins; the variability is likely due to the presence of non-centrosomal aggregates of PCMI [24] and cell-cycle differences among the population of cells resulting in variable γ-tubulin localization to the centrosome [25, 26]. Microtubule morphology (α-tubulin) appeared to be unaffected in the JBTS10 cell line as compared to control (Fig. 2b).
also examined the structure of these compartments in the affected cells [27]. As judged by immunofluorescence, there was no change in the morphology of the endoplasmic reticulum and the cis- and trans-Golgi of UDP-3331 compared to control adult dermal fibroblasts (Fig. 2c, d).

Decreased ciliation number in OFD1 p.Gln886Lysfs*2 mutant cells
The above observations raised the question as to whether the remaining OFD1 protein that localized to the centrosome in UDP-3331 dermal fibroblasts was sufficient to support ciliogenesis. Comparable to a fetal-affected JBTS10 cell line (UW172-4, NM_003611.2: c.277G > T, p.Val93Phe [28]), UDP-3331 dermal fibroblasts showed a decreased ability to form cilia that was rescued to near WT levels by expression of WT OFD1 (Fig. 2c, d). Thus, consistent with previous studies [8], the reduction in OFD1 mRNA and protein levels in UDP-3331 dermal fibroblasts is sufficient to impede cilia formation.

JBTS10 tissues have OFD1-dependent anomalies of protein glycosylation
Evaluation of UDP-3331 included a screen for anomalies of N- and O-linked glycans in plasma and of free oligosaccharides in urine. This detected a decrease in mature, fully sialylated and galactosylated N-linked glycans at m/z 2973 (Table 2), whereas monosialo glycan at m/z 2431 and monosialo, monogalacto glycan at m/z 2227 were increased. In plasma O-linked glycans sialylated T antigen (see Additional file 1: Table S1) was decreased, and T antigen was at the low end of the normal range. Both the plasma N- and O- glycosylation profiles indicated an abnormality in sialic acid utilization in the trans-Golgi or abnormal metabolism of sialic acids (see Additional file 2: Figure S1). Urinary oligosaccharides were within the normal range. Supporting the association of glycosylation anomalies with JBTS10, similar plasma N-linked findings were detected in another individual with JBTS10 (NIH-452, NM_003611.2: c.149A>G [29, 30], Table 2); O-linked
Table 1  Phenotype of JBTS10 patients

| Joubert syndrome phenotypes (JBST1; OMIM: 213,300) | UDP-3331 | NIH-452 |
|---------------------------------------------------|----------|---------|
| **Head and neck**                                  |          |         |
| Macrocephaly                                       | +        | +       |
| **Face**                                           |          |         |
| Prominent forehead                                 | −        | +       |
| High, rounded eyebrows                             | −        | +       |
| Hemifacial spasms                                  | −        | −       |
| Low-set ears                                       | +        | +       |
| ‘Tilted’ ears                                      | −        | −       |
| **Eyes**                                           |          |         |
| Abnormal, jerky eye movements                      | −        | +       |
| Impaired smooth pursuit                            | +        | +       |
| Impaired saccades                                  | +        | +       |
| Oculomotor apraxia                                 | +        | +       |
| Coloboma of optic nerve                            | −        | −       |
| Chorioretinal coloboma                             | −        | −       |
| Epicanthal folds                                   | +        | −       |
| Ptosis                                             | −        | +       |
| **Nose**                                           |          |         |
| Upturned nose                                      | +        | +       |
| Anteverted nostrils                                | −        | +       |
| **Mouth**                                          |          |         |
| Triangular-shaped open mouth                       | +        | +       |
| Protruding tongue                                  | −        | −       |
| Rhythmic tongue movements                          | −        | −       |
| **Respiratory**                                    |          |         |
| Neonatal breathing dysregulation                   | +        | +       |
| Hyperpnea, episodic                                | −        | +       |
| Tachypnea, episodic                                | −        | +       |
| Central apnea                                      | +        | +       |
| **Liver**                                          |          |         |
| Hepatic fibrosis (less common)                     | −        | +       |
| **Central nervous system**                         |          |         |
| Delayed psychomotor development                    | +        | +       |
| Mental retardation/intellectual disability         | +        | +       |
| Ataxia                                             | −        | +       |
| Hypotonia                                          | +        | +       |
| Occipital meningocele (less common)                | +        | −       |
| Hypoplasia of the brainstem                        | −        | −       |
| Malformation of brainstem structures               | +        | +       |
| ‘Molar tooth sign’ on MRI                          | +        | +       |
| Cerebellar vermis hypoplasia                        | +        | +       |
| Dysgenesis or agenesis of the cerebellar vermis    | +        | +       |
| Deep posterior interpeduncular fossa               | +        | +       |
| Thick and elongated superior cerebellar peduncles  | +        | +       |
| **Behavioral psychiatric manifestations**           |          |         |
| Hyperactivity                                      | −        | −       |
| Aggressiveness                                     | −        | −       |
| Self-mutilation                                    | +        | +       |
| **Additional phenotypes**                          |          |         |
| Submucosal cleft palate                            | +        | −       |
| Notched upper lip or tongue                        | +        | +       |
| Oral motor dysfunction                             | +        | +       |
| Recurrent aspiration pneumonia                     | +        | −       |
| Easy bruising                                      | −        | −       |
| Rockerbottom feet                                  | −        | −       |
| Hypertricosis                                      | +        | −       |
| Moderate hearing loss                              | +        | −       |
| Double hair whorl                                  | +        | −       |
| GI tube placed in infancy                          | +        | +       |
glycosylation was not tested in this individual due to limited sample amount.

To exclude the possibility that these findings were non-specific, we defined the N- and O-linked glycome profile in cultured dermal fibroblasts from JBTS10 patients UDP-3331 and UW172-4. Both cell lines had increased levels of mono-sialylated N-glycan species compared to controls (Table 3). Both cell lines also had abnormal formation of core 2 mature, sialylated O-linked glycans (see Additional file 3: Table S2) and thus a relative high abundance of the core 1 species. Core 1 sialylated T antigen from cellular protein was not reduced. Each N-glycosylation abnormality in the cultured JBTS10 dermal fibroblasts was partially rescued by expression of WT OFD1 (Table 3, UDP-3331 + OFD1). As was observed for the plasma results in Table 2, we observed a defect in sialylated glycans in JBTS10 patient fibroblast samples.

Cultured JBTS10 dermal fibroblasts have increased nuclear CMP-sialic acid

To characterize further the glycomic profile abnormalities, we performed lectin immunofluorescence using SNA, which preferentially binds to α-2,6-linked sialic acids, and MALII, which primarily binds α-2,3-linked sialic acids. We observed increased nuclear localization of the SNA fluorescently labeled lectin signal in both of the JBTS10 dermal fibroblast lines, and this was reduced upon rescue with WT OFD1 (Fig. 4a, b, \( p < 0.05 \)).

Given that the increased nuclear SNA signal was only observed in JBTS10 cells fixed with paraformaldehyde but not with methanol (Fig. 4c), we hypothesized that the SNA-reactive epitope was a small molecule, lipid, or protein not preserved by methanol fixation. Unlike other nucleotide sugars, which are synthesized in the cytosol, CMP-sialic acid (CMP-SA) is synthesized in the nucleus and, under aldehyde fixation, could constitute a small SNA-reactive epitope [31]. Using HPSEC-PAD, comparison of nucleotide sugars levels between UDP-3331 and adult WT cultured dermal fibroblasts showed a 39% increase in CMP-SA in UDP-3331 (Fig. 5a, b); introduction of WT OFD1 decreased the CMP-SA levels by 50% (\( p < 0.005 \), unpaired \( t \) test).

Neuraminidase treatment destabilizes cellular primary cilia

Given the association between abnormal sialylation and ciliogenesis in the JBTS10 patient lines, we sought to examine the effects of disrupting sialylation on cilia stability in control cells. Following growth in ciliogenic conditions, WT cells were either left untreated or treated with neuraminidase to remove sialic acids from the cellular glycoproteome. The neuraminidase used is isolated from *Clostridium perfringens* and can cleave α-2,3-, α-2,6-, or α-2,8-linked sialic acids, although it has highest affinity for α-2,3-linkages. Cells were then fixed and stained for cilia markers and cilia were measured. We observed a significant decrease (\( p < 0.001 \)) in average cilia length in WT cells following neuraminidase treatment and a modest (\( p < 0.05 \)) decrease in the number of ciliated cells in the population (Fig. 6a–d). Neuraminidase treatment successfully disrupted glycoprotein sialylation as indicated by increased lectin staining with PNA, which strongly binds galactose epitopes normally masked by terminal sialic acids (Fig. 6e). Additionally, in the absence of neuraminidase treatment, we observed co-localization of SNA-reactive epitopes with the cilium marker ARL13B, indicating that some proteins localized to the primary cilia are sialylated (Fig. 6f).

Discussion

Herein we have detailed the clinical and molecular phenotypes of a boy, UDP-3331, with Joubert syndrome type 10 (JBTS10). His clinical, molecular, and cellular findings are consistent with other reports of JBTS10. Interestingly, we observed a global alteration in the N- and O-linked glycosylation profiles of three JBTS10 patients. The most pronounced and consistent glycosylation change was a decrease of mature, fully sialylated N-glycans and an increase in total cellular CMP-sialic acid. Within cultured skin fibroblasts, lentiviral rescue with WT *OFD1* cDNA normalized the abnormal glycosylation profiles in...
Fig. 2 Cellular morphology and analysis of JBTS10 skin fibroblasts. a Co-localization analysis of OFD1 (magenta) signals in proband, proband rescued, and adult WT cells with γ-tubulin (red) and PCM1 (green). Inset higher magnification of the region outlined with dotted lines. b Immunofluorescence staining for OFD1 (green) and α-tubulin (red) in fibroblasts of the proband (UDP-3331), proband rescued (UDP-3331 + OFD1), and adult WT control. c Comparison of endoplasmic reticulum structure (KDEL, green) in UDP-3331 and WT cells. d Comparison of the cis- (GM-130, green) and trans- (TGN-46, red) Golgi in UDP-3331 and WT cells. DNA was stained with Hoechst 33,342 (blue). All scale bars represent 20 µm.
JBTS10 patient cells and the ciliogenesis phenotype previously associated with OFD1 mutations.

The observed decrease in primary cilia stability following neuraminidase removal of sialic acids from the glycoproteome and the co-localization of sialylated glycoproteins with the primary cilium suggest that protein sialylation plays a role in ciliary homeostasis. We hypothesize, therefore, that the abnormal glycosylation might contribute to the ciliogenesis defect. Two factors, however, remain to be addressed. First, our glycomic studies on primary dermal fibroblasts were performed under growth conditions that do not promote ciliogenesis, and therefore, it is unclear how much of the abnormal glycosylation is attributable to ciliary proteins. Second, because of technical limitations to performing glycome profiling on the cilia, we are unable to assess biochemically whether the ciliary proteome is aberrantly glycosylated in patient cells. These two limitations provide testable hypotheses for future investigations.

Regardless of the impact directly on the ciliary proteins, the accumulation of the sialic acid epitope that binds to the lectin SNA in the nuclei of JBTS10 patient cells and the increased CMP-SA and the accumulation of incompletely sialylated N-glycans are consistent with a defect in CMP-SA metabolism or transport. Unlike other nucleotide sugars, CMP-SA is synthesized in the nucleus, rather than the cytosol; thus, CMP-SA must be transported through two distinct membrane-bound organelles for proper sialylation to occur [32, 33]. There are many possible ways in which alterations in levels of functional OFD1 could lead to changes in sialylated glycoproteins. First, nuclear OFD1 [13] might contribute to CMP-SA metabolism. Second, defects of ciliogenesis might alter cell signaling [34–36] or the role of the basal body in regulating microtubules and intracellular transport to impair CMP-SA transport from the nucleus and subsequent utilization in the Golgi. Disruption of the centrosome structure and tethering of the daughter centrioles are associated with abnormal Golgi structure [37], and alterations in the structure of the Golgi cisternae would undoubtedly influence the glycosylation enzymes that are targeted specifically to cis- or trans-Golgi compartments [38]. However, we did not observe any obvious changes in centriole separation or Golgi structure. Third, the increased sialylation of O-linked core 1 T antigen on cellular proteins and the decreased sialylation of O-linked core 1 T antigen on secreted glycoproteins in plasma might suggest altered exocytosis and endocytosis of glycoproteins. Finally, elevated processing of sialoglycoproteins or enhanced turnover of surface sialoglycoproteins might contribute to the abnormal glycosylation signature observed [39]. Each of these potential mechanisms highlights the important role of ciliary and basal body proteins in cell physiology and the delicate homeostatic balance that can be indirectly disrupted by single-gene mutations.

Prior associations between primary ciliopathies and altered glycosylation have been limited to observations of the mutated ciliary proteins. Polycystin-2, polycystin-1, and aquaporin-11 mutations associate with abnormal glycosylation of ciliary proteins [40–42]. Changes in protein trafficking, which result in altered protein glycosylation, have been linked to polycystic liver disease [43].

There exists corroborating evidence that altered glycosylation might contribute to the pathogenesis of some ciliopathies. Primary defects in glycosylation, collectively referred to as Congenital Disorders of Glycosylation (CDGs), cause phenotypes similar to those of primary ciliopathies and disrupt cilia function. Defects in the N-linked
glycosylation genes ALG3, ALG8, and ALG9, as well as the O-linked glycosylation gene B3GALT1, variably result in polycystic kidneys, cleft lip or palate, polycystic ovaries, and abnormal development of the brain including the corpus callosum [44–47]. Mutations in GALNT11, encoding polypeptide N-acetylgalactosaminyltransferase, cause ciliary dysfunction and heterotaxy [48].

The importance of glycosylation in neural development is well illustrated by animal models of disrupted glycosylation. A knock-out of the only sialyltransferase gene in D. melanogaster caused significant neurological symptoms, implicating sialylation as a key factor in neurodevelopment and function [49]. Additionally, disruption of proteoglycan formation in C. elegans by alteration of microRNA levels causes aberrant migration of neurons [50]. Alterations of polysialic acid epitope formation in mice also cause abnormal neuronal development and disrupted neuron migration, likely due to changes in sialylation of NCAM [51]. The cerebellum and midbrain anomalies that result in the molar tooth sign (MTS) are thought to arise due to aberrant migration of neurons in the cerebellum. Given that both N- and O-linked CDGs present with abnormal development of other brain structures, and animal models with altered sialylation show distinct neurodevelopmental defects, it is possible that the altered glycosylation we observe in JBTS10 patients may contribute to the formation of MTS. Recent work by Feng and colleagues to identify the N-glycoproteome in the mouse brain revealed that several key proteins for brain development are N-glycosylated [52].

### Table 2 Summary of N-glycan profiles of JBTS10 patient plasma

| Centroid mass (m/z) | Predicted glycan species | UDP-3331 (% total glycans) | NIH-452 (% total glycans) | Ref low (% total glycans) | Ref high (% total glycans) |
|---------------------|--------------------------|-----------------------------|---------------------------|--------------------------|---------------------------|
| 1580.2              | Hex5 HexNAc2             | 0.69                        | 0.91                      | 0.23                     | 1.88                      |
| 1662.3              | Hex3 HexNAc4             | 0.03                        | 0.00                      | 0.00                     | 0.22                      |
| 1784.4              | Hex6 HexNAc2             | 0.51                        | 0.48                      | 0.00                     | 0.87                      |
| 1866.5              | Hex4 HexNAc4             | 0.19                        | 0.00                      | 0.06                     | 0.53                      |
| 1982.5              | Hex4 NeuAc1 HexNAc3      | 0.48                        | 0.43                      | 0.08                     | 0.80                      |
| 1988.5              | Hex7 HexNAc2             | 0.19                        | 0.00                      | 0.00                     | 0.39                      |
| 2070.6              | Hex5 HexNAc4             | –                           | 0.15                      | 0.00                     | 0.30                      |
| 2156.7              | Hex4 dHex1 NeuAc1        | 0.05                        | 0.00                      | 0.06                     | 0.17                      |
| 2186.7              | Hex4 dHex1 NeuAc1 HexNAc3| –                           | 0.39                      | 0.00                     | 0.89                      |
| 2192.7              | Hex8 HexNAc2             | 0.32                        | 0.00                      | 0.06                     | 0.61                      |
| 2227.7              | Hex4 NeuAc4 NeuAc1       | 1.38                        | 1.54                      | 0.40                     | 1.28                      |
| 2285.2              | Hex4 dHex1 HexNAc5       | 0.17                        | –                         | 0.00                     | 0.16                      |
| 2431.9              | Hex5 HexNAc4 NeuAc1      | 5.87                        | 9.88                      | 2.38                     | 6.74                      |
| 2606.0              | Hex5 dHex1 HexNAc4 NeuAc1| 0.54                        | 1.87                      | 0.79                     | 2.87                      |
| 2793.3              | Hex5 HexNAc4 NeuAc2      | 20.29                       | 40.58                     | 28.38                    | 58.23                     |
| 2966.5              | Hex5 dHex1 HexNAc4 NeuAc2| 1.49                        | 2.89                      | 2.49                     | 3.36                      |
| 3241.6              | Hex6 HexNAc5 NeuAc2      | 1.13                        | 1.52                      | 0.08                     | 1.70                      |
| 3415.6              | Hex6 dHex1 HexNAc5 NeuAc2| 0.25                        | 0.00                      | 0.00                     | 0.87                      |
| 3602.9              | Hex6 HexNAc5 NeuAc3      | 3.31                        | 6.72                      | 1.64                     | 6.10                      |
| 3776.6              | Hex6 dHex1 HexNAc5 NeuAc3| 0.94                        | 0.38                      | 0.00                     | 0.72                      |

The percentages of total glycans for the indicated N-glycan species in the plasma of UDP-3331 and NIH-452 are shown along with reference (Ref) range derived from controls. Species with values in the low-normal range in the JBTS10 samples are flagged with a pound sign (#). Species below the normal range in JBTS10 sample(s) are flagged with a dagger (†). Species that are elevated above the normal range are flagged with a caret (^).

*a*Low normal in JBTS10

* Above normal in JBTS10

* Below normal in JBTS10
**Table 3  Summary of N-glycan profiles of JBTS10 patient fibroblasts**

| Centroid mass (m/z) | Glycan structure | UDP-3331 (% total glycans) | UDP-3331 +OFD1 (% total glycans) | UW172-4 (% total glycans) | Adult control (% total glycans) | Ref low (% total glycans) | Ref high (% total glycans) |
|---------------------|------------------|-----------------------------|-----------------------------------|---------------------------|-------------------------------|--------------------------|---------------------------|
| 1171.9              | Hex3 HexNAc2     | 4.09                        | 7.07                              | 1.63                      | 5.00                          | 0.23                     | 12.38                     |
| 1346.0              | Hex3 dHex1       | 3.86                        | 5.28                              | 1.06                      | 5.09                          | 0.00                     | 6.11                      |
| 1417.1              | Hex3 HexNAc3     | 0.00                        | 0.79                              | 0.00                      | 0.78                          | 0.00                     | 0.78                      |
| 1580.2              | Hex5 HexNAc2     | 8.19                        | 11.68                             | 4.55                      | 8.46                          | 1.83                     | 14.29                     |
| 1621.3              | Hex4 HexNAc3     | 0.56                        | 0.76                              | 0.00                      | 0.98                          | 0.00                     | 0.98                      |
| 1662.3              | Hex3 HexNAc4     | 0.00                        | 0.00                              | 0.00                      | 0.00                          | 0.00                     | 0.29                      |
| 1784.4              | Hex6 HexNAc2     | 19.88                       | 17.97                             | 15.17                     | 18.31                         | 6.54                     | 20.74                     |
| 1825.4              | Hex5 HexNAc3     | 0.57                        | 0.62                              | 0.00                      | 0.90                          | 0.00                     | 0.90                      |
| 1866.5              | Hex4 HexNAc4     | 0.74                        | 0.81                              | 0.00                      | 0.66                          | 0.00                     | 0.70                      |
| 1982.5              | Hex4 NeuAc1      | 0.00                        | 0.00                              | 0.00                      | 0.00                          | 0.00                     | 0.30                      |
| 1988.5              | Hex7 HexNAc2     | 9.81                        | 9.62                              | 12.04                     | 9.67                          | 4.87                     | 14.86                     |
| 2029.6              | Hex6 HexNAc3     | 0.54                        | 0.52                              | 0.00                      | 0.93                          | 0.00                     | 2.19                      |
| 2070.6              | Hex5 HexNAc4     | 4.39                        | 4.03                              | 2.03                      | 3.20                          | 0.00                     | 3.88                      |
| 2156.7              | Hex4 dHex1       | 0.00                        | 0.34                              | 0.00                      | 0.52                          | 0.00                     | 0.75                      |
|                    | Hex3 HexNAc3     |                            |                                   |                           |                               |                          |                           |
| 2192.7              | Hex8 HexNAc2     | 16.18                       | 13.96                             | 27.66                     | 16.04                         | 0.00                     | 32.90                     |
| 2227.7              | Hex4 HexNAc4     | 0.00                        | 0.00                              | 0.00                      | 0.00                          | 0.00                     | 0.20                      |
| 2396.9              | Hex9 HexNAc2     | 9.28                        | 7.49                              | 18.11                     | 9.09                          | 0.24                     | 32.34                     |
| 2431.9              | Hex5 HexNAc4     | 2.18                        | 1.47                              | 2.92                      | 1.83                          | 0.00                     | 1.83                      |
| 2606.0              | Hex5 dHex1       | 6.85                        | 4.19                              | 2.23                      | 4.70                          | 0.00                     | 4.70                      |
|                    | Hex3 HexNAc4     |                            |                                   |                           |                               |                          |                           |
| 2793.3              | Hex5 HexNAc4     | 0.86                        | 0.45                              | 1.53                      | 1.00                          | 1.21                     | 6.80                      |
| 2967.3              | Hex5 dHex1       | 2.69                        | 1.45                              | 4.74                      | 2.02                          | 3.38                     | 10.79                     |
| 3241.6              | Hex6 HexNAc5     | 0.00                        | 0.00                              | 0.00                      | 0.17                          | 0.00                     | 1.96                      |
| 3415.6              | Hex6 dHex1       | 0.45                        | 0.22                              | 0.84                      | 0.39                          | 0.43                     | 5.84                      |
| 3602.9              | Hex6 HexNAc5     | 0.33                        | 0.00                              | 0.00                      | 0.28                          | 0.64                     | 23.23                     |
| 3776.6              | Hex6 dHex1       | 0.21                        | 0.00                              | 0.79                      | 0.11                          | 0.00                     | 5.46                      |

The percentages of total glycans of the indicated N-glycan species for UDP-3331, UDP-3331 + OFD1 rescue, UW172-4, and adult WT control fibroblasts samples are shown along with reference (Ref) low and high values derived from control samples. Species with values in the low-normal range in the JBTS10 sample(s) are flagged with a pound sign (#). Species that are elevated above the normal range in JBTS10 sample(s) are flagged with a caret (^). Species below the normal range in JBTS10 sample(s) are flagged with a dagger (†).

* Low normal in JBTS10
^ Above normal in JBTS10
† Below normal in JBTS10
Fig. 4 Lectin staining for sialic acid epitopes in JBTS10 and control skin fibroblasts. 

**a** Lectin staining of α-2,6-sialic acid (SNA, green) and α-2,3-sialic acid (MALII, blue) linkages in affected, rescued and control cells, as indicated. Arrows indicate atypical nuclear staining by SNA in UDP-3331. Scale bars 20 μm. 

**b** Quantification of the fluorescence signal for SNA in nucleus relative to the entire cell for affected, rescued, and control cells under paraformaldehyde fixation conditions. Error bars represent standard error. Single asterisks indicate \( p < 0.05 \); double asterisks indicate \( p < 0.001 \) relative to both adult and pediatric male controls (unpaired, heteroscedastic t test). 

**c** Comparison of methanol and paraformaldehyde fixation on the relative SNA staining intensity of the nucleus vs. the whole cell. Double asterisks indicate \( p < 0.001 \) (unpaired, heteroscedastic t test). Error bars represent standard error.
**Fig. 5** Nucleotide sugar analysis in JBTS10 and control skin fibroblasts. 

**a** Representative HPAEC trace for UDP-3331, UDP-3331 + OFD1 rescue, adult WT, adult WT + OFD1 overexpression, pediatric male control cells, and 500 nM standard CMP-sialic acid. 

**b** Quantification of CMP-sialic acid levels in affected and control cells (in μM) (*single asterisks* indicate p < 0.005 in an unpaired, heteroscedastic t test). Error bars represent standard error. 

**Fig. 6** Neuraminidase treatment of adult WT primary cells. 

**a** Average cilia length of WT cells with and without neuraminidase (±Neuase) treatment to remove sialic acids from glycan chains; neuraminidase treatment significantly reduced average cilia length by ~1 μm. Error bars represent standard error; p < 0.001 as determined in an unpaired, heteroscedastic t test. 

**b** The percentage of ciliated cells is decreased after neuraminidase treatment; p < 0.05 as determined by a two-tailed Fisher’s Exact test; error bars represent the 95% confidence interval for the binomial distribution represent by the sample population. 

**c, d** Representative images of WT fibroblasts without (c) and with (d) neuraminidase treatment. Cilia are stained with ARL13B antibody (red) and γ-tubulin (green). Scale bars 20 μm. 

**e** Lectin staining for sialic acid with SNA (green) and MAL II (white) and galactose with PNA (red) in WT fibroblasts with DRAQ5 DNA counterstaining (blue). 

**f** Representative image of SNA (green) signal showing alpha 2,6-sialylation co-localizing with ARL13B-positive cilia (red) in WT cells without neuraminidase treatment. Scale bars 5 μm.
Conclusions
Our findings highlight a potential role for glycosylation in contributing to pathogenesis of JBTS10 and possibly other ciliary disorders. The mechanism for the altered glycosylation remains to be determined. Our observations raise the question as to whether glycosylation is altered in other types of Joubert syndrome and more severe ciliopathies. The impact of altered glycosylation in contributing to pathophysiology in ciliopathies bears further study.

Additional files

Additional file 1: Table S1. O-glycan analysis results of UDP-3331 plasma.

Additional file 2: Figure S1. Overview of N- and O-linked glycosylation pathways. N-linked glycosylation begins with the synthesis of the dolichol-phosphate-linked glycan precursor on the cytosolic face of the endoplasmic reticulum (ER). Once the precursor molecule is flipped into the ER lumen, it is further built up before being transferred to the nascent polypeptide. After the glycosylated polypeptide is properly folded, it is transferred to the cis Golgi where the high-37 mannose species are trimmed and further matured by the addition of N-acetylglucosamine (GlcNAc), followed by galactose, fucose and, finally, sialic acid to form the most prevalent, Complex N-linked glycans. An alternative pathway to generate hybrid glycans also exists. The enzymes that perform each of these glycan-building steps are targeted to either cis, medial, or trans Golgi stacks to facilitate proper assembly of the sugars. Two representative O-linked glycosylation pathways are shown: O-GalNAc and O-mannose. Much of the O-linked glycosylation reactions occur in the Golgi apparatus and, similar to N-linked glycosylation, the complexity and length of the sugars extends as the protein transits through the Golgi cisternae. In JBTS10 patient samples, we observe a decrease in formation of the mature, fully sialylated N-glycans and alterations in levels of the O-linked sialylated species. These species are shown in trans Golgi cisternae, which has been highlighted in blue.

Additional file 3: Table S2. O-glycan analysis results of UDP-3331 and UW172-4 primary dermal fibroblasts.

Abbreviations
JBTS10. Joubert syndrome type 10, JBTS. Joubert syndrome; ODFI. orofaciiodental syndrome I; SGBS2. Simpson–Golabi–Behmel syndrome, type 2; PCP. planar cell polarity; NIH UDP. National Institutes of Health Undiagnosed Diseases Program; DMEM. Dulbecco’s Modified Eagle Medium; Hi FBS. Heat inactivated fetal bovine serum; PBS. phosphate-buffered saline; WT. wild-type; HPAEC-PAD. high-performance anion-exchange chromatography with pulsed amperometric detection; CMA-SA. CMP-sialic acid; CDG. congenital disorder of glycosylation; NHGRI. National Human Genome Research Institute.

Authors’ contributions
MSK designed and performed multiple experiments and was a major contributor in writing the manuscript. MD contributed to experimental design, performed multiple experiments, and was a participant in writing and revision of the manuscript. MRB and JAH contributed to CMP-sialic acid measurements and key discussions in experimental design. CIA contributed to immunofluorescence studies, performed exome variant analysis, and contributed to writing “Methods” section of the manuscript. MEG, IGP, DRO, JCD, and DD performed citogeness experiments and data analysis as well as drafting of a portion of “Methods” section. XL performed mass spectrometry measurements of N- and O-glycans and contributed in writing “Methods” section. CG, GV, MGA, and WAG were involved in clinical evaluation and diagnosis of the individuals seen at NIH (UDP-3331 and NIH-452). MH performed analysis of mass spectrometry data for all glycome testing and contributed in writing the manuscript. MCVM and CFB were instrumental in overall experimental design, project supervision, and manuscript drafting. All authors read and approved the final manuscript.

Author details
1 NIH Undiagnosed Disease Program, Common Fund, Office of the Director, and National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA. 2 Inova Translational Medicine Institute, Inova Health System, Falls Church, VA, USA. 3 National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, MD, USA. 4 Department of Pediatrics, University of Washington, Seattle, WA, USA. 5 The Michael J Palmieri Metabolic Laboratory, Children’s Hospital of Philadelphia, Philadelphia, PA, USA. 6 Children’s National Medical Center, Washington, DC, USA. 7 Medical Genetics Branch, National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA. 8 Office of the Clinical Director, National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA. 9 Johns Hopkins University School of Medicine, Department of Pediatrics and McKusick-Nathans Institute of Genetic Medicine, Baltimore, MD, USA. 10 Department of Medical Genetics, University of British Columbia, Vancouver, BC, Canada.

Acknowledgements
The authors would like to acknowledge the patients and their families for participation in their respective research protocols.

Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Consent for publication
All patient data presented within this work is presented with full consent of the legal guardian for each patient.

Ethics approval and consent to participate
Research protocols involving patients, NIH clinical protocol 76-HG-0238 and NCT00068224, were subject to approval by the National Human Genome Research Institute (NHGRI) Institutional Review Board. Informed consent was provided by the patients’ legal guardians. Additional review was provided by the NHGRI Scientific Review Committee.

Funding
This work was supported in part by the Intramural Research Program of the National Human Genome Research Institute and the NIH Common Fund Program. Additional funding for the laboratory of D. Doherty came from the Eunice Kennedy Shriver National Institute of Child Health and Human Development Grant U54HD083091, the National Institute for Neurological Defects and Stroke Grant RO1NS064077 and private donations.

Received: 7 October 2016 Accepted: 17 February 2017
Published online: 23 March 2017

References
1. Parisi M, Glass I. Joubert syndrome and related disorders. In: Pagon RA, Adam MP, Ardinger HH, Wallace SE, Amemiya A, Bean L, Bird TD, Ledbetter N, Melford HC, Smith RJH, Stephens K, editors. GeneReviews(R). Seattle: University of Washington, 1993.
2. Valente EM, Dallapiccola B, Bertini E. Joubert syndrome and related disorders. Handb Clin Neurol. 2013. doi:10.1016/B978-0-444-59565-2.00058-7.
3. Szymanska K, Hartill VL, Johnson CA. Unraveling the genetics of Joubert and Meckel–Gruber syndromes. J Pediatr Genet. 2014. doi:10.3233/ PGE-14009.
4. Romani M, Micalizzi A, Valente EM. Joubert syndrome: congenital cerebellar ataxia with the molar tooth. Lancet Neurol. 2013. doi:10.1016/ S1474-4422(13)70136-4.
5. Webb TR, Parfitt DA, Gardner JC, Martinez A, Bevilacqua D, Davidson AE, Zito I, Thielson DL, Ressa JH, Aperi M, et al. Deep intronic mutation
in OFD1, identified by targeted genomic next-generation sequencing, causes a severe form of X-linked retinitis pigmentosa (RP23). Hum Mol Genet. 2012. doi:10.1093/hmg/ddr119.

6. Tang Z, Lin MG, Stowe TR, Chen S, Zhu M, Stearns T, Franco B, Zhong Q. Autophagy promotes primary cilia formation by removing OFD1 from centriolar satellites. Nature. 2013. doi:10.1038/nature12600.

7. Singla V, Romaguera-Ros M, Garcia-Verdujo JM, Reiter JF. Ofd1, a human disease gene, regulates the length and distal structure of ciliae. Dev Cell. 2010. doi:10.1016/j.devcel.2009.12.022.

8. Ferrante MI, Zullo A, Barra A, Bimonte S, Messaddeq N, Studer M, Dolle P, Franco B. Oral-facial-digital type I protein is required for primary cilia formation and left–right axis specification. Nat Genet. 2006. doi:10.1038/ng1694.

9. D'Angelo A, De Angelis A, Vallone E, Biscopo I, Tammaro R, Studer M, Franco B. Ofd1 controls dorso-ventral patterning and axoneme elongation during embryonic brain development. PLoS ONE. 2012. doi:10.1371/journal.pone.0052937.

10. Ferrante MI, Romio L, Castro S, Collins JE, Goulding DA, Stemple DL, Woolf DM, Rue TC, Ishikawa GE, Isabella CR, Gorden N, Adkins J, et al. Joubert syndrome: a model for untangling recessive disorders with extreme genetic heterogeneity. J Med Genet. 2015. doi:10.1136/jmedgenet-2015-103087.

11. Vilboux T, Doherty D, Glass IA, Parisi MA, Malicdan MC, Phelps IG, Cullinan AR, Zein W, Heller T, Soldatos A, et al. Molecular genetic findings and clinical correlations in 101 patients with Joubert syndrome and related disorders prospectively evaluated at a single center. Genetics Med. 2016.

12. Hoffman EA, Frey BL, Smith LM, Auble DT. Formaldehyde crosslinking: a tool for the study of chromatin complexes. J Biol Chem. 2015. doi:10.1016/j.ribo.2015.05.027.

13. Geng L, Okuhara D, Yu Z, Tian X, Cai Y, Shibazaki S, Somlo S. Polycystin-2 traffics to cilia independently of polycystin-1 by using an N-terminal RVxP motif. J Cell Sci. 2006. doi:10.1242/jcs.028188.

14. Quinn S, Taglieni M, Nauli SM, Contrino L, Takakura A, Zhou J, Kreidberg JA. Failure to ubiquitinate c-Met leads to hyperactivation of mTOR signaling in a mouse model of autosomal dominant polycystic kidney disease. J Clin Invest. 2010. doi:10.1172/JCI41531.

15. Davila S, Furru L, Gharavi AG, Tian X, Oneto N, Qian Q, Li A, Cai Y, Kamath PS, Maynard JD, King BF, et al. Mutations in SEC63 cause autosomal dominant polycystic kidney disease. Nat Genet. 2010. doi:10.1038/ng.1057.

16. Yu SH, Bond MR, Whitman CM, Kohler JJ. Metabolic labeling of glycoprotein-dependent adhesion mechanisms in cultured insect cells and mammalian cells. Anal Biochem. 2001. doi:10.1006/abio.2001.5091.

17. Schneider CA, Rasband WS, Eliceiri KW. NIH Image to ImageJ: 25 years of image analysis. Nat Methods. 2012. doi:10.1038/nmeth.2013.07.037.

18. Yu SH, Bond MR, Whitman CM, Kohler JJ. Metabolic labeling of glycoprotein-dependent adhesion mechanisms in cultured insect cells and mammalian cells. Anal Biochem. 2001. doi:10.1006/abio.2001.5091.

19. Kubo A, Tsukita S. Non-membranous granular organelle consisting of POM-1: subcellular distribution and cell-cycle-dependent assembly/disassembly. J Cell Sci. 2003. doi:10.1242/jcs.002822.

20. Muroyama A, Seldin L, Lechler T. Divergent regulation of functionally distinct gamma-tubulin complexes during differentiation. J Cell Biol. 2016. doi:10.1083/jcb.201601099.

21. Woodruff JB, Wueseke O, Hyman AA. Pericentriolar material structure and dynamics. Philos Trans R Soc Lond B Biol Sci. 2014. doi:10.1098/rstb.2013.0459.

22. Gurel PS, Hatch AL, Higgs HN. Connecting the cytoskeleton to the endoplasmic reticulum and Golgi. Curr Biol. 2014. doi:10.1016/j.cub.2014.05.033.

23. Tomiya N, Ailor E, Lawrence SM, Betenbaugh MJ, Lee YC. Determination of nucleotides and sugar nucleotides involved in protein glycosylation by high-performance anion-exchange chromatography: sugar nucleotide contents in cultured insect cells and mammalian cells. Anal Biochem. 2001. doi:10.1006/abio.2001.5091.

24. Kubo A, Tsukita S. Non-membranous granular organelle consisting of POM-1: subcellular distribution and cell-cycle-dependent assembly/disassembly. J Cell Sci. 2003. doi:10.1242/jcs.002822.
46. Tham E, Eklund EA, Hammarsjo A, Bengtson P, Geiberger S, Lagerstedt-Robinson K, Malmgren H, Nilsson D, Grigelionis G, Conner P, et al. A novel phenotype in N-glycosylation disorders: Gillessen–Kaesbach–Nishimura skeletal dysplasia due to pathogenic variants in ALG9. Eur J Hum Genet. 2016. doi:10.1038/ejhg.2015.91.

47. Heinonen TY, Maki M. Peters’-plus syndrome is a congenital disorder of glycosylation caused by a defect in the 1,3-glucosyltransferase that modifies thrombospondin type 1 repeats. Ann Med. 2009. doi:10.1080/07853890802301975.

48. Boskovski MT, Yuan S, Pedersen NB, Goth CK, Makova S, Clausen H, Brueckner M, Khokha MK. The heterotaxy gene GALNT11 glycosylates Notch to orchestrate cilia type and laterality. Nature. 2013. doi:10.1038/nature12723.

49. Repnikova E, Koles K, Nakamura M, Pitts J, Li H, Ambavane A, Zoran MJ, Panin VM. Sialyltransferase regulates nervous system function in Drosophila. J Neurosci. 2010. doi:10.1523/JNEUROSCI.15253-09.2010.

50. Pedersen ME, Snieckute G, Kajias K, Nehammer C, Multhaupt HAB, Couchman JR, Pocock R. An epidermal MicroRNA regulates neuronal migration through control of the cellular glycosylation state. Science. 2013. doi:10.1126/science.1242528.

51. Krocher T, Rockle I, Diederichs U, Weinhold B, Burkhardt H, Yanagawa Y, Gerardy-Schahn R, Hildebrandt H. A crucial role for polysialic acid in developmental interneuron migration and the establishment of interneuron densities in the mouse prefrontal cortex. Development. 2014. doi:10.1242/dev.111773.

52. Fang P, Wang XJ, Xue Y, Liu MQ, Zeng WF, Zhang Y, Zhang L, Gao X, Yan GQ, Yao J, et al. In-depth mapping of the mouse brain N-glycoproteome reveals widespread N-glycosylation of diverse brain proteins. Oncotarget. 2016. doi:10.18632/oncotarget.9737.