Compound heterozygous variants in the multiple PDZ domain protein (MPDZ) cause a case of mild non-progressive communicating hydrocephalus

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**Abstract**

**Background:** Congenital hydrocephalus (CH) results from the accumulation of excessive amounts of cerebrospinal fluid (CSF) in the brain, often leading to severe neurological impairments. However, the adverse effects of CH can be reduced if the condition is detected and treated early. Earlier reports demonstrated that some CH cases are caused by mutations in \(L1CAM\) gene encoding the neural cell adhesion molecule L1. On the other hand, recent studies have implicated the multiple PDZ domain (MPDZ) gene in some severe forms of CH, inherited in an autosomal recessive pattern.

**Methods:** In this study, whole-exome and Sanger sequencing were performed on a 9 months old Emirati child clinically diagnosed by CH. In addition, *in silico*, cellular, and molecular assays have been conducted to confirm pathogenicity of the identified variants and to establish disease mechanism.

**Results:** Whole exome sequencing revealed two compound heterozygous novel variants (c.394G > A and c.1744C > G) in the affected child within the \(MPDZ\) gene. Segregation analysis revealed that each of the parents is heterozygous for one of the two variants and therefore passed that variant to their child. The outcome of the *in silico* and bioinformatics analyses came in line with the experimental data, suggesting that the two variants are most likely disease causing.

**Conclusions:** The compound heterozygous variants identified in this study are the most likely cause of CH in the affected child. The study further confirms \(MPDZ\) as a gene underlying some CH cases.

**Keywords:** Congenital hydrocephalus, Multiple PDZ, \(L1CAM\), Compound heterozygous variants, Autosomal recessive
Another gene associated with hydrocephalus is the AP1S2 which encodes a subunit of the adaptor protein 1 [11]. Interestingly, Al-Dosari et al., have recently reported cases from two Saudi families with congenital hydrocephalus and identified a founder homozygous mutation in MPDZ gene leading to a truncated protein. The pattern of disease in these families and segregation analysis clearly suggested a recessive mode of inheritance. The hydrocephalus in the two families was very severe leading to stillbirths in two of the affected cases, and death of several affected children in early infancy [12]. More recently, three novel null mutations were described in MPDZ gene in three unrelated hydrocephalic fetuses with multiple ependymal malformations. Histological and confocal studies performed on post-mortem sections revealed multifocal ependymal rosette formation and absence of MPDZ in ependymal lining [13].

MPDZ is an acronym from Multiple PSD95, DLG and ZO-1, they are protein modules found in many cytoplasmic proteins [14]. The MPDZ gene is located on the cytogenetic band 9p23 of chromosome 9, and encodes a 13-domain MPDZ protein that localizes at cellular junctions. In addition, numerous interactions have been documented for MPDZ suggesting that it might have multifaceted roles in the assembly and localization of the integral membrane proteins, including cell-cell junction. Hence, it regulates conductive permeability, prevents spontaneous acrosomal exocytosis and maintains GABA receptors, olfactory and melatonin receptor signaling [15]. Therefore, it is not surprising that various MPDZ domains bind to those receptors [16]. Moreover, MPDZ proteins are made of multiple domains that can bind multivalent scaffold proteins, and regulate intracellular signaling, receptor clustering and polarity of epithelial cells [17].

We report in this manuscript an Emirati family with a child affected by congenital non-progressive communicating hydrocephalus, who was found to be compound heterozygous for two novel likely pathogenic variants in the MPDZ gene. The phenotype of the affected child is milder than the previously reported cases associated with mutations in MPDZ; which indicates that the associated clinical phenotypes can be variable. This variability could be due to the nature of the variants and/or the presence of as yet unidentified disease-modifier genes.

**Methods**

The participating family was referred to Tawam Hospital in Al Ain city (UAE) for evaluation of macrocephaly and communicating hydrocephalus in their first born male child aged 9 months. DNA samples from peripheral blood were extracted for the affected child and his parents, using Flexigene DNA extraction kit (Qiagen, Gmbh, Germany) following the manufacturers’ instructions.

**Whole-exome sequencing and variant analysis and prioritization**

Whole-exome sequencing for the child was performed as a service at the Medical Genetic Laboratories of Baylor College of Medicine (www.bcmgeneticlabs.org). The output data from Illumina HiSeq were converted to FastQ files by CASAVA 1.8 software, and mapped by BWA program, the variant calls were performed using Atlas-SNP and Atlas-indel developed in house by BCM HGSC. The variants were interpreted according to ACMG guidelines [18] and patient phenotypes. The WES report included minor allele frequency (MAF) data for variants reported in the NHLBI GO Exome Sequencing Project (ESP) ESP5400 database. For example, “1/3738 1/7017” in AA/EA, respectively. That means for the variant listed, the minor allele frequency was observed one time, the major allele was observed 3738 times in (AA). The annotated SNPs were mapped to dbSNP v131, v132 and 1000 Genome. Moreover, candidate variants were cross compared with Ensembl database [19], as well as to the Exome variant server [20]. In order to determine the effect of the candidate variants at the protein level, prediction analyses using Mutation Taster, Polyphen2 and SIFT/Provean were as well performed [21–25].

**Sanger DNA sequencing**

The segregation of the candidate variants was confirmed by Sanger DNA sequencing in all available family members. PCR Primers were designed using Primer3 software [26] . Moreover, we checked the novelty of the variants from 100 ethnically matched control samples. Sanger sequencing was performed using Big Dye Terminator Kit (Applied Biosystems, USA) on 3130xl Genetic Analyzer (Applied Biosystems, USA). Clustal Omega algorithm was applied to analyze sequencing results [27].

**RNA extraction and cDNA sequencing**

Total RNA was extracted from whole blood using SV Total RNA Isolation system (Promega, USA), then reverse transcribed with random primers and GoScript reverse transcriptase enzyme according to the manufacturer’s instructions (Promega, USA). After cDNA was synthesized, primers were designed to hybridize the flanking region, which contained the variant (c.394G > A) of the cDNA using Primer3 software. That region was then amplified with GoTaq® Flexi DNA polymerase (Promega, USA). The presence of the variant was confirmed using Sanger sequencing by Big Dye Terminator Kit (Applied Biosystems, USA) on 3130xl Genetic Analyzer (Applied Biosystems, USA).
Generation of the mutant cDNAs for mammalian expression
Site-directed mutagenesis (SDM) was performed on the
MPDZ expression clone (EX-E1329-M10), Accession No.: NM_003829.4 from GeneCopoeia (http://genecopoeia.com). Both identified variants (c.394G > A; Gly132Ser) and (c.1744C > G; Leu582Val) were generated individually on the c-Myc- His-tagged ORF cDNA clones using Pfu Ultra High-Fidelity DNA polymerase (Stratagene). The following are the primers which were designed according to QuikChange Site-Directed Mutagenesis Protocol,

MPDZ-Gly132Ser-Forward: 5’CAAAAATATGGCCA
TATCAGCCATGTAGAAG3’.
MPDZ-Gly132Ser-Reverse: 5’CTTCTACATGGGCAC
TCTGGGCCCATATTTTG3’.

Both identified variants were successfully amplified by three times wash with PBS and incubation with anti-mouse IgG Fab2 Alexa Flour 555 antibody (dilution: 1:200; Cell Signaling Technology). Images were acquired using Nikon confocal Eclipse 80I microscope (Japan). The affected child was the first born son to a non-consanguineous couple from the UAE (Fig. 1a). The pregnancy was uncomplicated, but ultrasound at 20 weeks gestation revealed large head with dilated ventricles. Delivery was normal at term with a birth weight 4.1R gene [28]. To demonstrate the splice-effect of the identified variant (c.394G > A) on the mRNA level, exon 5 and parts of its flanking introns were sub-cloned into the mini-cassette vector using BStEI (GGTCACC) and NheI (GCTAGC) restriction enzymes (New England Biolabs), they were included in the following cloning primers: Forward primer: 5’-ATGGGTGTTCCACAC
TTACTAAG-3’, Reverse primer: 5’-TATTATGAGGTA
TTGCTAGCATGG-3’. After amplifying the desired genomic fragment of MPDZ gene which was around 647 bps, it was further purified using MinElute PCR Purification Kit (Qiagen. Gmbh, Germany). The respective enzymes were applied to digest the insert as well as the vector, and the latter was run along with its non-digested counterpart on gel electrophoresis in order to verify digestion. The digested vector was then gel-purified using MinElute Gel Extraction Kit (Qiagen. Gmbh, Germany). The PCR product of MPDZ gene fragment was then ligated to the vector and transformed. The clones were then sequenced to verify correct insertion.

To generate the variant in the minigene construct, site-directed mutagenesis (SDM) was performed on the vector harboring the wild-type MPDZ using the following primers: Forward primer: 5’-TCTAAACAGAGATC
GCCATG- 3’, Reverse primer: 5’-CATGGCGACTCTG
TTAGGA-3’. After amplifying the desired fragment, the mutant vector was sequenced to confirm mutagenesis.

HEK 293 T cells were maintained in HyClone™ High Glucose Modified Eagles medium (GE Healthcare Life Sciences), supplemented with 10% fetal bovine serum, 4 mM L-Glutamine, 4500 mg/L Glucose, Sodium Pyruvate, and 100 U/ml Penicillin/Streptomycin. They were then seeded in a 6-well plate. On the following day, two wells were transfected with the wild-type minigene construct, while the two other wells were transfected with the mutant, and cultured for 48 h at 37 °C, 5% CO2. Cells were, then scraped and washed with PBS, and RNA was extracted using SV Total RNA Isolation System (Promega, USA), then reverse transcribed with random primers and GoScript reverse transcriptase enzyme according to the manufacturer’s instructions (Promega, USA). Subsequently, the resulting cDNA was analyzed on agarose gel electrophoresis, and sequenced to assess RNA splicing.

Results
Clinical report
The affected child was the first born son to a non-consanguineous couple from the UAE (Fig. 1a). The pregnancy was uncomplicated, but ultrasound at 20 weeks gestation revealed large head with dilated ventricles. Delivery was normal at term with a birth weight 17 with their downstream and upstream flanking intron of 4.1R gene [28]. To demonstrate the splice-effect of the identified variant (c.394G > A) on the mRNA level, exon 5 and parts of its flanking introns were sub-cloned into the mini-cassette vector using BStEI (GGTCACC) and NheI (GCTAGC) restriction enzymes (New England Biolabs), they were included in the following cloning primers: Forward primer: 5’-ATGGGTGTTCCACAC
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Results
Clinical report
The affected child was the first born son to a non-consanguineous couple from the UAE (Fig. 1a). The pregnancy was uncomplicated, but ultrasound at 20 weeks gestation revealed large head with dilated ventricles. Delivery was normal at term with a birth weight
of 3400 g, length of 56 cm, and head circumference of 39 cm (>97th centile). There were no other reported neonatal problems. Family history revealed that the brother of the mother has a daughter with hydrocephalus and ventriculoperitoneal (VP) shunt. Unfortunately, no further clinical or radiological information were available for this child because the family refused to participate in the research; and hence, no clinical or molecular studies were performed on this branch of the family.

MRI brain at 3 months revealed widened CSF spaces in the frontoparietal region bilaterally, as well as mild dilatation of lateral and third ventricles (Fig. 1b). The cerebral volume was preserved and the white matter appeared normal. Findings were thought to be consistent with external hydrocephalus [29].

The patient was evaluated in our Genetics Clinic at the age of 9 months. His weight was 10 kg (75th centile), length 77 cm (90th–95th centile), and head circumference 50 cm (>97th centile; Z score 3.34). He had large head with frontal bossing and high arched palate, but no other dysmorphic features. He had normal developmental milestones. At the age of 14 months he was walking and saying a few words, and continued to have a large head. At 15 months his weight was 11.5 kg (>50th centile), height 84 cm (>90th centile) and head circumference 51 cm (>97th centile). Detailed ophthalmological examination was normal. Progress brain MRI at age 17 months showed less dilatation of ventricles and less widening of CSF spaces, in keeping with the natural progress of external hydrocephalus. Negative investigations included: microarray, urine organic acid, urine mucopolysaccharides (MPS), creatine phosphokinase (CPK) and skeletal survey.

Whole-exome sequencing revealed that the affected child has two compound heterozygous variants in the MPZD gene

Clinical whole-exome sequencing analysis at Baylor College Genetics Laboratory revealed two compound heterozygous variants (c.394G > A and c.1744C > G) in the MPDZ gene. Sanger sequencing confirmed the segregation of these compound heterozygous variants in this family with the child being compound heterozygous for both variants (Fig. 2a). The mother was heterozygous for the c.394G > A variant (Fig. 2a, left panels), while the father was heterozygous for the c.1744C > G variant (Fig. 2a, right panels). The first variant (c.394G > A) has been reported as a SNP (rs201101621); which leads to the substitution of glycine at 132 with serine (p.Gly132Ser). However, this SNP has a minor allele frequency of T = 0.0004 based on 1000 genome, and no homozygote were reported for this variation. In addition, computational analysis predicted that this variant is disease causing/damaging using Mutation Taster and SIFT/Provean software. More importantly, this change was also predicted to cause acceptor splice site change by Neural Network software (www.fruitfly.org) [30]. It predicted a score of 0.78 for the acceptor splice-site present at c.394-1G, which is one nucleotide upstream to the variant. Furthermore, the software predicted that the variant might lead to the use of the cryptic acceptor-splice sites within intron 5–6 with a score of 0.74, leading to skipping of exon 5 and the generation of a stop codon.

The second variant (c.1744C > G) was predicted to cause a missense mutation at amino acid position leucine 582 (p.Leu582Val). At the same locus, a different SNP (rs201275925) has been reported with G/A alleles. This SNP had a minor allele frequency of A = 0.0002. However, the change was synonymous. In silico analysis using Mutation Taster predicted that the p.Leu582Val variant was disease causing. However, SIFT/Provean predicted that the change was tolerable/neutral. Both variants were absent in Exome Variant Server for all the
tested population. Furthermore, there were no damaging variants in \textit{MPDZ} gene in a homozygous state in normal individuals in this database. Likewise, there were no reports of any homozygous damaging variations at the same locus in \textit{MPDZ} gene in NCBI. Additionally, WES analysis data identified two variants in heterozygous states in genes known to be implicated in hydrocephalus. One of which was in \textit{CCDC88C} gene (c.4327G > A) on chromosome 14; a previously reported reference SNP (rs189215037). This variant leads to an amino acid substitution (p.Ala1443Thr) which was predicted to be tolerated/benign by SIFT/PolyPhen-2 tools with ESP5400 AA/EA frequency of 0/3086 16/6636. The other variant was found in \textit{KIF7} gene (c.2777G > A) on chromosome 15 which was predicted to be a tolerable amino acid substitution (p.Arg926Gln), with a frequency was not reported in ESP5400 AA/EA. In addition, heterozygous variants in these two genes are not predicted to be disease-causing and therefore were excluded as relevant to the phenotype in the child.

cDNA sequencing revealed a likely unstable copy of the c.394G > A variant mRNA
It was predicted that c.394G > A variant would result in a splice site change. \textit{MPDZ} cDNA from the mother, the affected child and from a healthy control were all prepared and sequenced. In all three cases, a single sequencing peak (G) at position c.394 was observed (Fig. 2b). This was different from what we observed from the genomic DNA sequencing of the affected child and mother (Fig. 2a (left panels)). This discrepancy can be explained by the activation of nonsense-mediated decay (NMD) of the mutated copy of the mRNA, but not the wild type. This activation could be due to the introduction of the predicted premature stop codons.

The c.394G > A variant caused exon 5 skipping
We wanted to further investigate the effect of c.394G > A on mRNA splicing, therefore exon 5 and its flanking regions of \textit{MPDZ} gene were sub-cloned into a mini-cassette vector used for the analysis of splicing (Fig. 3). RNAs extracted from the transfected cells harboring wild type and mutant mini-cassette vectors were converted into cDNAs and then analyzed by agarose gel electrophoresis which revealed an aberrant band with a molecular weight of less than 200 bps appeared compared to the wild type band of 300 bps (Fig. 3). Upon sequencing both bands, the mutant band revealed skipping of exon 5 of \textit{MPDZ}, while conserving both of the vector’s exons, just as expected by the neural network prediction tool. On the other hand, the wild-type cDNA retained exon 5 along with the vector’s exons (Fig. 3).

Protein modeling of the p.Leu582Val variant indicated possible disruption of the protein structure
The second variant (c.1744C > G) affected a highly conserved residue (p.Leu582Val) (Fig. 4a). The images in fig. 4b and c were generated using Schrödinger Prime and rendered using PyMOL. The variant which was located within the MPDZ protein was predicted to cause functional disruption of the domain. The mutant amino acid valine lies between βC and αA of PDZ4, interfering with the conserved GLG repeat which is essential for hydrogen bond coordination of the c-terminal carboxylate group. Moreover, as already mentioned, the site of leucine at this position displays evolutionary conservation consistent with the damaging nature of the change at that position.

Confocal analysis revealed that the pathogenicity of the compound heterozygous variants is not due to subcellular localization defects
To establish whether the pathogenicity of the amino acid substitution (p.Leu582Val) is due to the subcellular mislocalization of the mutant protein, HeLa cells were
**Fig. 3** Illustrative model of the mini-gene analysis protocol. Patterned box depicts exon 5 and its flanking introns of the subcloned MPDZ gene, black ones depict exons 13 and 17 of 4.1R gene which was subcloned into pRc/CMV vector. Splicing of the wild type cDNA shows the authentic splicing and the inclusion of the three exons and that is noticed from the size of the amplified band on the gel (WT). Whereas, the mutant cDNA (Mu) shows a smaller band due to cryptic splicing and exon skipping. On the right side, the chromatograms represent both splice variants, dashed lines separate the joined exons.

**Fig. 4 a** Multiple sequence alignments for MPDZ protein among different Eukaryotic species by Clustal Omega show the conservation of amino acid leucine among selected Eukaryotic species. **b** The rat PDZ4 (wild type) was used to generate the image in PyMOL. The conserved GLG loop is shown in red with ball and stick representation. The secondary structures (βA, βB, βC, αA, βD, βE, αB, βF) are shown with the wild-type Leucine located between βC and αA. **c** PDZ4 (mutant type). The rat PDZ4 structure (PDB ID: 4XH7) was used to generate a model of the human PDZ4 domain using Schrodinger Prime. The image was generated in PyMOL. The mutant valine which lies between βC and αA could disrupt or interfere with the GLG loop.
transfected with plasmids harboring the wild type and the mutant MPDZ cDNAs. Although we have shown that the c.394G > A variant is altering RNA splicing, we wanted to test the alternative possible implication of this variant (i.e. generating of a missense mutation at position p.Gly132Ser), we therefore generated this change as well, using the MPDZ cDNA and then expressed the mutants (p.Leu582Val, p.Gly132Ser) and wild type in HeLa cells. In all cases, cells were co-transfected with GFP-H-Ras as a plasma membrane marker. MPDZ protein is known to co-localize with synaptic markers and with tight junctions such as claudin-5 at cell membranes [31], as well as the cytosol [32]. Confocal imaging showed that the localization of the two variants is similar to that of the wild type (Fig. 5), thus, excluding the possibility that the pathogenicity of both variants is linked to mis-localization of the MPDZ protein.

**Discussion**

The reported child has congenital communicating hydrocephalus with no associated abnormalities and near normal development. The hydrocephalus was non-progressive and did not require shunting. The affected child was found to have compound heterozygous variants in the MPDZ gene. A single homozygous CH-causing mutation in this gene has been recently reported in two individuals from two unrelated Saudi Arabian families. The phenotype in these previously reported families is much more severe compared to our patient. In addition to a severe progressive hydrocephalus requiring shunting, other abnormalities were reported in the affected individuals including simplified gyration pattern in all of them, and coloboma with congenital heart disease in one of them. In addition, there were two still births presumably due to CH in one family and six CH cases in the second extended family. The cognitive outcome of the two long term survivors in Al-Dosari et al. (2013) correlated with the speed of neurosurgical intervention initiation after birth [12]. Moreover, five hydrocephalic stillbirths from three unrelated families were identified. Three probands were genetically tested to find null mutations in MPDZ gene. Medical termination was achieved to all fetuses except for one in whom the child died from severe hydrocephalus [13]. Table 1 lists all the
previously reported mutations in \textit{MPDZ} gene in patients with congenital hydrocephalus.

The mutation reported by Al-Dosari et al. (2013), led to a 12-domain truncated protein out of, originally 13 domains, abolishing most of the interactions with other proteins and rendering the affected members severely hydrocephalic. This mutation likely leads to a complete loss-of-function of the \textit{MPDZ} protein. Compared to our study, we detected a splice site and missense variants that might, provisionally affect domains (between L27 and PDZ-1, as well as PDZ4, respectively). \textit{MPDZ} protein has many possibilities of interactions due to its multiple-domain structure. Each domain is highly conserved among species and each is made up of six beta sheets and two alpha helices (Fig. 4b and c). Furthermore, the 13 PDZ domains are > 50% homologous in rat MUPP1/MPDZ protein. [33] We present in fig. 6 the alignment of the 13 domains in human \textit{MPDZ}, showing homology in the preserved GLG loop, as well as other locations. This loop lies between $\beta$A and $\beta$B sheets. It functions as a coordinator for hydrogen bond of the C-terminal carboxylate group of the binder protein. Another important domain in \textit{MPDZ} is the hydrophobic pocket, which lies between $\beta$B and $\alpha$B, constituting the canonical binding sites for \textit{MPDZ}. Opposite to this binding groove, at about 30 residues C-terminal extension of the PDZ domain lie other non-canonical

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|l|l|}
\hline
DNA Change & Protein Change & No. of Patients & Origin & Phenotypes & Ref. \\
\hline
\textit{c.628C > T} & p.(Gln210*) & 1 & Saudi & Severe congenital supratentorial hydrocephalus, mild hypotonia, coloboma, atrial septal defect & [12] \\
\textit{c.628C > T} & p.(Gln210*) & 1 & Saudi & Severe congenital supratentorial hydrocephalus & [12] \\
\textit{c.1291_1294del} & Frame-shift & 2 & Senegalese & Massive hydrocephalus & [13] \\
\textit{c.533 + 1G > T} & Frame-shift & 2 & – & Foetal macrosomia with severe isolated ventriculomegaly & [13] \\
\textit{c.2248C > T} & p.(Arg750*) & 3 & sibs & Recurrent hydrocephalus & [13] \\
\textit{(Compound heterozygous)} & Predicted: p.Gly132Ser & 1 & Emirati & Congenital non-progressive communicating hydrocephalus with normal developmental milestones & Current study \\
\textit{c.394G > A} & Experimentally shown to cause exon 5 skipping & & & & \\
\textit{c.1744C > G} & & & & & \\
\end{tabular}
\caption{Reported \textit{MPDZ} mutations in patients with congenital hydrocephalus}
\end{table}

Fig. 6 The alignment of the 13 PDZ domains in \textit{MPDZ} protein of human, indicating the homologous regions in colored boxes showing a graphical view of those conserved regions by Jalview [40]
binding sites [34]. In the present study, the change from leucine (non-polar and bulkier side chain) to valine (non-polar with smaller side chain) at position 582 is predicted to be disturbing the GLG loop (Fig. 4c). Despite the fact that glycine is neither at the GLG loop nor within the hydrophobic pocket, it is predicted that the reported variant c.394G > A is in an acceptor splice site region, leading to a stop codon and truncation of the majority of the protein domains, rendering the protein with only L27 domain. PDZ-1 domain enhances binding of other PDZ domains. For example, PDZ-2 domain of Syntaxin (a PDZ domain that functions as an adaptor) binds neurexin, ephrin-1 and syndecan only when paired with PDZ-1 [35, 36]. Moreover, Grootjans et al. (2000) showed that PATJ, a protein involved in cell polarity and a mammalian paralogue of MPDZ, has a PDZ-1 domain linked to L27 domain. When they decreased both domains in mammalian cell culture, it resulted in disassembly in tight junctions. Strikingly, they had to be linked together for PATJ to function properly [36].

Moreover, the absence of the mutated copy of mRNA that was revealed by sequencing the cDNA for the mother and child suggested an NMD mechanism which is consistent with the prediction of the variant (c.394G > A) affecting an acceptor splice-site according to Neural Network prediction tool. Following that, we had to investigate the effect of that variant on MPDZ expression by quantifying mRNA using real time PCR. Unfortunately, MPDZ is expressed at very low levels in the hematopoietic lineages (https://www.ncbi.nlm.nih.gov/unigene). However, transcripts could be detected in patient blood samples by RT-PCR, which enabled further characterization of the mutant. Besides, higher tissue sources expressing MPDZ such as skin fibroblasts could not be obtained from the patient.

Furthermore, to experimentally verify the effect of the variant (c.394G > A) on mRNA splicing and to reveal the possible disease mechanism; mutant and wild mini-gene constructs were designed to analyze splicing variants [37]. Sequencing both cDNAs revealed the actual adjoining of the exons in the wild-type, while skipping of exon 5 of MPDZ gene within the mutant. This is explained by the fact that the variant (c.394G > A) is located at the first nucleotide of exon 5 which was predicted to cause a splice-site aberration (Fig. 3).

The second variant (c.1744C > G), leucine is also conserved at that position among multiple species (Fig. 4a). It lies between βC sheet and αA of PDZ-4 domain. This change could be disease causing/probably damaging according to Mutation Taster and PolyPhen, respectively. It has been shown that binding of PDZ-5 domain of GRIP (Glutamate receptor interacting protein) to its receptor GluK2 failed to occur in the absence of PDZ-4 domain, indicating its role in the proper folding of the other domain [38]. Although our variant did not take place in any of the canonical binding sites of MPDZ, function of PDZ-4 was not only restricted to such binding, rather PDZ-4 was shown to be involved in the direct binding of PDZK1 (a MPDZ domain-containing protein in hepatocytes which controls the expression and localization of HDL receptors) to the inner leaflet of the cell membrane. And that this binding was not due to any of the four PDZ domains in that protein, but to PDZ-4 solely [39]. It was assumed that since MPDZ localizes at cell-cell adhesion sites and is involved in regulating permeability, mutations in this protein would disrupt this localization. Hence, alteration in CSF secretion would result in hydrocephalus. After performing confocal microscopy on the wild type and mutant plasmids, we observed no difference in the subcellular localization of either mutant compared to the wild type. Therefore, we assume that the pathogenesis of these missense changes is not due to trafficking defects; rather it could be due to disruption or complete loss of the protein function or an essential protein-protein interaction.

Conclusions
The identified compound heterozygous variants detected by whole-exome sequencing and confirmed by Sanger sequencing in the affected child are the most likely cause of his CH. The deleterious effects of the two variants have been illustrated experimentally and by bioinformatics tools. The clinical presentation is consistent with CH which we strongly argue to be caused by the compound heterozygous variants in MPDZ gene.

Abbreviations
AA: African American; CH: Congenital hydrocephalus; CPK: Creatine phosphokinase; CSF: Cerebrospinal fluid; DAPLE: Dvl-Associating Protein With A High Frequency Of Leucine; EU: European; GRIP: Glutamate receptor interacting protein; L1CAM: L1 Cell Adhesion Molecule; MAF: Minor Allele Frequency; MPDZ: Multiple PDZ; MPS: Mucopolysaccharides; NMD: Nonesense-mediated decay; SDM: Site-directed mutagenesis; VP: Ventriculoperitoneal

Acknowledgments
We thank the family for taking part in this study. We also thank the United Arab Emirates University for funding (Grant 31 M135) and supporting NKA PhD studies (Grant 31 M144). We would also like to thank Dr. Mireille Deguillien and her co-workers for providing the (p(13,17)/CMV) splicing cassette.

Funding
United Arab Emirates University grants 31M135 and 31M144.

Availability of data and materials
The datasets supporting the conclusions of this article are included within the article.

Authors’ contributions
NKA had undertaken the research work, prepared the manuscript and analyzed the data. BA and LG are joint corresponding authors who have provided the intellectual input, supervision and designs and approved the protocols to be followed in the study, in addition to correcting and proofreading the manuscript. AJ has done the sequencing, RV conducted
the protein modelling, JS and AMA had provided the MRI image and the clinical presentation of the patient and revised the manuscript. SB had revised the manuscript and helped with the initial data analysis. All authors read and approved the manuscript.

Ethics approval and consent to participate
The study on elucidating the genes and mutations underlying rare genetic disorders in the UAE has been approved by Al-Ain Medical Human Research Ethics Committee (protocol number 10/09 and ERH-2015-3241 15–115). Written informed consent was obtained from the participating family.

Consent for publication
Consent for publication was obtained from the child’s parents.

Competing interests
The authors declare that there were no competing interests.

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Received: 23 February 2017 Accepted: 12 February 2018
Published online: 02 March 2018

References
1. Tully HM, Dobyns WB. Infantile hydrocephalus: a review of epidemiology, classification and causes. Eur J Med Genet. 2014;57(8):359–68.
2. Vintzileos AM, Ingardia CJ, Nochimson DJ. Congenital hydrocephalus: a review and protocol for perinatal management. Obstet Gynecol. 1983;62(5):539–49.
3. Munch TN, Rostgaard K, Rasmussen MLH, Wohlfahrt J, Juhler M, Melbye M. Familial aggregation of congenital hydrocephalus in a nationwide cohort. Brain. 2012;135(Pt 2):409–15.
4. Schrander-Stumpel C, Fryns JP. Congenital hydrocephalus: nosology and phenotype correlations in L1 syndrome: a guide for genetic counselling and prediction of sequence variations revisions 2007. Genet Med. 2008;10(4):294–300.
5. Abdul-Karim R, Iliya F, Iskandar G. Consecutive hydrocephalus: report of two cases. Obstet Gynecol. 1964;24:376–8.
6. Lapunzina P, Delicaco A, De Torres ML, Mori MA, Perez-Pacheco RF, Lopez-Palacios I. Autosomal recessive hydrocephalus due to aqueduct stenosis: report of a further family and implications for genetic counselling. J Matern Fetal Med. 2002;12(1):64–6.
7. Barros-Nunes P, Rivas F. Autosomal recessive congenital stenosis of the aqueduct of Sylvius. Genet Couns. 1993;4(1):19–23.
8. Petrini M, Dulau G, Rochiccioli P. Autosomal recessive inheritance of hydrocephalus with Sylvius aqueductal stenosis. J Genet Hum. 1981;29(2):155–60.
9. Hamada H, Watanabe T, Sugimoto M, Yasuoka M, Yamada N, Kubo T. Autosomal recessive hydrocephalus due to congenital stenosis of the aqueduct of Sylvius. Prenat Diagn. 1999;19(11):1067–9.
10. Vos YJ, De Walle HEK, Bos NK, Stegeman JA, Ten BERGE HEK, Bos KK, Stegeman JA, Ten Berge AM, Bruining M, Van Maarel KC, Elting MW, Den Hollandier NS, Hamel B, et al. Genotype–phenotype correlations in L1 syndrome: a guide for genetic counselling and mutation analysis. J Med Genet. 2010;47(3):169–75.
11. Saillour Y, Zanni G, Des Portes V, Heron D, Guibaud L, Iba-Zizen MT, Pedespan JL, Poirier K, Castelnaud L, Julien C, et al. Mutations in the AP152 gene encoding the sigma 2 subunit of the adaptor protein 1 complex are associated with syndromic X-linked mental retardation with hydrocephalus and calcifications in basal ganglia. J Med Genet. 2007;44(11):739–44.
12. Al-Dosari MS, Al-Owain M, TuBhaish M, Kurdi W, Adly N, Al-Hemmian A, Masoodi TA, Albash B, Alkurya FS. Mutation in MPDZ causes severe congenital hydrocephalus. J Med Genet. 2013;50(1):54–8.
13. Saugier-Veber P, Marguet F, Lecoquiere F, Adle-Biassette H, Guimiot F, Cipriani S, Patier S, Brasseur-Daudruy M, Goldenberg A, Layet V, et al. Hydrocephalus due to multiple ependymal malformations is caused by mutations in the MPDZ gene. Acta Neuropathol Commun. 2015;7(1):36.
14. Fehr C, Shirley RL, Metten P, Kosobud AEk, Belknap JK, Crabbe JC, Buck KJ. Potential pleiotropic effects of MpDz on vulnerability to seizures. Genes Brain Behav. 2004;3(1):8–19.
15. Ackermann F, Zitranski N, Borth H, Buech T, Gu德ermann T, Boekhoff I. CamKIIalpha interacts with multi-PDZ domain protein MUPP1 in spermatozoa and prevents spontaneous acrosomal exocytosis. J Cell Sci. 2009;122(Pt 24):4547–57.
16. Ali M, Hocking PM, Mcklbbin M, Finnegian S, Shires M, Poulter JA, Prescott K, Booth A, Rashid Y, Jafri H, et al. MPDZ null allele in an avian model of retinal degeneration and mutations in human leber congenital amaurosis and retinitis pigmentosa. Invest Ophthalmol Vis Sci. 2011;52(8):4742–40.
17. Hamazaki Y, Itoh M, Sasaki H, Furuse M, Tsukita S. Multi-PDZ domain protein 1 (MUPP1) is concentrated at tight junctions through its possible interaction with claudin-1 and junctional adhesion molecule. J Biol Chem. 2002;277(1):455–61.
18. Richards CS, Bale S, Bellisimo DB, Das S, Grody WW, Hegde MR, Lyon E, Ward BEACMG. Recommendations for standards for interpretation and reporting of sequence variations revisions 2007. Genet Med. 2008;10(4):136–42.
19. Aken BL, Ayling S, Barrell D, Clarke L, Curwen V, Fairley S, Fernandez-Barquet J, Bills K, Garcia-Giron C, Houriier T, et al. The Ensembl gene annotation system. Database (Oxford). 2016; https://doi.org/10.1093/database/baw093.
20. Exome Variant Server. NHLBI exome sequencing project (ESP). Seattle, WA; 2015; http://evs.gs.washington.edu/EVS/. Accessed Jan 2015.
21. Choi Y, Sims GE, Murphy S, Miller JR, Chan AP. Predicting the functional effect of amino acid substitutions and Indels. PLoS One. 2012;7(10):e46688.
22. Adzhubei IA, Schmidt S, Peshkin L, Ramensky VE, Gerasimova A, Bork P, Kondrashov AS, Sunyaev SR. A method and server for predicting damaging missense mutations. Nat Methods. 2010;7(4):248–9.
23. Kumar P, Henikoff S, Ng PC. Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. Nat Protoc. 2009;4(7):1073–82.
24. Schwarz JM, Rodeleperger C, Schuelke M, Seelow D. MutationTaster evaluates disease-causing potential of sequence alterations. Nat Methods. 2010;7(5):575–6.
25. Sim NL, Kumar P, Hu J, Henikoff S, Schneider G, Ng PC. SIFT web server: predicting effects of amino acid substitutions on proteins. Nucleic Acids Res. 2012;40(W1):W452–7.
26. Untergasser A, Cutucache I, Koressaar T, Ye J, Faircloth BC, Remm M, Rozen SG. Primer3–new capabilities and interfaces. Nucleic Acids Res. 2012;40(15):e115.
27. Severs F, Wilms A, Dineen D, Gibson TJ, Karplus KI, Li W, Lopez R, McWilliam H, Remmert M, Soding J, et al. Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal omega. Mol Syst Biol. 2011;7:539.
28. Deguillen M, Huang SC, Moriniere M, Dreumont N, Benze JR, Baklouti F. Multiple cis elements regulate an alternative splicing event at 4.1 R pre-mRNA during erythroid differentiation. Blood. 2001;98(13):3809–16.
29. Zahl SM, Egge A, Helseth E, Wester K. Benign external hydrocephalus: a review, with emphasis on management. Neurosurg Rev. 2011;34(4):417–22.
30. Reese MG, Eeckman FH, Kulp D, Haussler D. Improved splice site detection in genie. J Comput Biol. 1997;4(3):311–23.
31. Poliak S, Matlis S, Ullmer C, Scherer S, Peles E. Distinct Claudins and Retinitis Pigmentosa proteins in human retinal photoreceptor cells. J Cell Biol. 2002;158(8):1049–55.
32. Al-Jezawi et al. BMC Medical Genetics (2018) 19:34
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37. Desviat LR, Perez B, Ugarte M. Minigenes to confirm exon skipping mutations. Methods Mol Biol. 2012;867:37–47.
38. Hung AY, Sheng MPDZ. Domains: structural modules for protein complex assembly. J Biol Chem. 2002;277(8):5699–702.
39. Tsukamoto K, Wales TE, Daniels K, Pal R, Sheng R, Cho W, Stafford W, Engen JR, Krieger M, Kocher O. Noncanonical role of the PDZ4 domain of the adaptor protein PDZK1 in the regulation of the hepatic high density lipoprotein receptor scavenger receptor class B, type I (SR-BI). J Biol Chem. 2013;288(27):19845–60.
40. Clamp M, Cuff J, Searle SM, Barton GJ. The Jalview java alignment. Bioinformatics. 2004;20(3):426–7.