REPORT

Bi-allelic variants in HOPS complex subunit VPS41 cause cerebellar ataxia and abnormal membrane trafficking

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Membrane trafficking is a complex, essential process in eukaryotic cells responsible for protein transport and processing. Deficiencies in vacuolar protein sorting (VPS) proteins, key regulators of trafficking, cause abnormal intracellular segregation of macromolecules and organelles and are linked to human disease. VPS proteins function as part of complexes such as the homotypic fusion and vacuole protein sorting (HOPS) tethering complex, composed of VPS11, VPS16, VPS18, VPS33A, VPS39 and VPS41. The HOPS-specific subunit VPS41 has been reported to promote viability of dopaminergic neurons in Parkinson’s disease but to date has not been linked to human disease. Here, we describe five unrelated families with nine affected individuals, all carrying homozygous variants in VPS41 that we show impact protein function. All affected individuals presented with a progressive neurodevelopmental disorder consisting of cognitive impairment, cerebellar atrophy/hypoplasia, motor dysfunction with ataxia and dystonia, and nystagmus. Zebrafish disease modelling supports the involvement of VPS41 dysfunction in the disorder, indicating lysosomal dysregulation throughout the brain and providing support for cerebellar and microglial abnormalities when vps41 was mutated. This provides the first example of human disease linked to the HOPS-specific subunit VPS41 and suggests the importance of HOPS complex activity for cerebellar function.

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**Keywords:** cerebellar ataxia; membrane trafficking; neurodevelopmental disorder; VPS41; zebrafish disease modelling
Abbreviations: CRISPR = clustered regularly interspaced short palindromic repeats; ESC = embryonic stem cell; GFP = green fluorescent protein; HOPS = homotypic fusion and vacuole protein sorting; LRO = lysosome-related organelle; VPS = vacuolar protein sorting

Introduction

Membrane trafficking in eukaryotic cells is critical for protein transport and processing of endocytic cargo for degradation. It is regulated by highly conserved vacuolar protein sorting (VPS) proteins, which were first discovered through genetic screening for yeast mutants exhibiting several classes of distinct trafficking phenotypes (Conibear and Stevens, 1995; Bonangelino et al., 2002). Analogous phenotypes have since been observed in mammalian cells (Huizing et al., 2008) and trafficking defects and VPS protein deficiencies have been linked to numerous human neurological diseases (Supplementary Table 1). For example, a homozygous founder mutation in VPS11 was found in eight affected individuals from four unrelated Ashkenazi Jewish families who presented with hypomyelination, developmental delay, hypotonia, microcephaly, and seizures (Zhang et al., 2016), while VPS33A loss-of-function has been linked to psychomotor retardation, delayed myelination, cerebral calcification, and lysosomal dysfunction (Kondo et al., 2017). However, whether other VPS proteins also have a role in disease remains to be clarified.

VPS proteins frequently function within multi-subunit tethering complexes, the best-documented being the hexameric HOPS (homotypic fusion and vacuole protein sorting) and CORVET (class C core vacuole/endosome tethering) complexes. The HOPS complex is thought to regulate fusion of lysosomes with endo- or autophagosomes and consists of six VPS subunits: VPS39 and VPS41, which are HOPS-specific, as well as VPS11, VPS16, VPS18 and VPS33A, which are also present in the CORVET complex (Bowers and Stevens, 2005). VPS39 and VPS41 interact with RAB7 to specifically target the HOPS complex to late endosomes (Balderhaar and Ungermann, 2013).

Yeast strains with disrupted VPS41 function exhibit deformities in vacuolar morphology and trans-Golgi trafficking of vacuolar components that can be rescued upon complementation with the wild-type gene (Radisky et al., 1997). As lysosomes in higher eukaryotes are analogous organelles to yeast vacuoles, abnormalities in lysosome-related organelles (LROs)—including lysosomes, melanosomes and phagosomes, among others—are likely to occur in instances of VPS41 deficiency, for example in vertebrate cells (Delevoye et al., 2019). However, proof of this theory could not be established to date because of early embryonic lethality of the only currently described Vps41 mouse knockout model preventing further analysis (Aoyama et al., 2012).

Here, we report five unrelated families with nine affected individuals with homozygous missense variants in VPS41, presenting with an autosomal recessive, progressive developmental disorder consisting of cerebellar atrophy/hypoplasia, cognitive impairment, motor dysfunction with ataxia and dystonia, and abnormal membrane-bound vesicles, as assessed by electron microscopy in lymphocytes and lymphoblastoid cells.

Materials and methods

Detailed methodology is provided in the Supplementary material.

Human subjects

All probands were investigated by their referring physicians. Most genetic analyses were performed in a diagnostic setting. Legal guardians of affected individuals gave informed consent for genomic investigations and publication of anonymized data. This study was approved by institutional review board of KFSHRC (RAC# 2120022). Exome sequencing was performed as previously described (AlMuhaizea et al., 2020; Perenthaler et al., 2020).

In silico analyses

Functional pathway gene ontology enrichment analyses were performed using Database for Annotation, Visualization and Integrated Discovery (DAVID) (Dennis et al., 2003) and Ingenuity Pathways Analysis (IPA) (QIAGEN Inc.). A right-tailed Fisher’s exact test was used to calculate a P-value determining the probability that the biological function (or pathway) assigned to that dataset is explained by chance alone. Gene interaction network analysis was performed using IPA. Genes were mapped to their corresponding gene objects in the Ingenuity® Knowledge Base. SwissModel, RaptorX and iTASSER were used to produce homology models. Models were manually inspected and mutations evaluated using PyMOL.

Human embryonic stem cell experiments

VPS41 knockout embryonic stem cells (ESCs) were generated by clustered regularly interspaced short palindromic repeats (CRISPR)-Cas9 mutagenesis, as previously described (Barakat et al., 2018; Perenthaler et al., 2020). ESCs were used in rescue experiments in which wild-type or mutant VPS41 was expressed transiently. Full experimental details are provided in the Supplementary material.

CRISPR/Cas9 zebrafish mutagenesis

CRISPR/Cas9 mutagenesis was performed largely as previously described using guide RNA (gRNA) targeting exon 4 of vps41. Cas9 nuclease was synthesized (Kuil et al., 2019) and co-injected with gRNAs into fertilized zebrafish oocytes. Indel efficiency and frequency were determined using Sanger sequencing.
LysoTracker™ staining

For LysoTracker™ staining, carried out as previously described (Kuil et al., 2019), zebrafish larvae were incubated in the dark at 2°C for 40 min in 1.5 ml tubes with 100 μl LysoTracker™ Red DND-99 (ThermoFisher) diluted to a final concentration of 10 μM in E3 media containing 200 μM PTU. Medium was replaced with fresh E3-PTU and incubated 20 min before imaging.

In vivo imaging and image analysis

Larvae were anaesthetized using tricaine (0.016%) prior to being mounted in 1.8% low melting point agarose (Kuil et al., 2019). In vivo fluorescence imaging used a Leica SP5 intravital confocal microscope with a 20× water-dipping objective (Leica Plan-Apochromat, NA = 1.0). Images were analysed using ImageJ software (Schindelin et al., 2012).

Optokinetic response assay

Zebrafish embryos [5 days post-fertilization (dpf)] were positioned in 6% methylcellulose/1× E3 within clear plastic wells and placed on a platform surrounded by a motorized drum (20 rpm) lined with black and white stripes of defined spatial frequencies on interchangeable inserts. Embryos were assessed for 1 min per trial, with trials consisting of assessment in both clockwise and counterclockwise directions with inserts achieving either 0.06 or 0.10 cycles per degree. Infrared monitoring and a custom program written by Dr Frank Schaeffel (University of Tubingen) were used to monitor and measure eye movement/position.

Statistical analyses

For continuous variables, data are reported either as individual values with the median and standard error of the mean (SEM) indicated or as box and whisker plots indicating the interquartile range (IQR) and median, minimum, and maximum values of the range. Categorical variables are summarized as percentages. Continuous variables were compared by Student’s t-test (or Mann-Whitney U-test) or one-way ANOVA as applicable, while the categorical variables are compared by Fisher’s exact test. Statistical analyses were conducted using GraphPad Prism 6.00 (La Jolla, California, USA) and PARTEK Genomics Suite (Partek Inc., St. Lois, MO, USA). The statistical level of significance was set at P < 0.05.

Data availability

The data that support the findings of this study are available from the corresponding authors, upon reasonable request, with the exception of primary patient sequencing data, which cannot be made available due to patient consent.

Results

Clinical phenotype and genetic investigations

Affected individuals were born after uneventful pregnancies and presented in most cases early in life with developmental delay (Fig. 1, Table 1, Supplementary Figs 1–5 and Supplementary Table 2). Various degrees of ataxia, hypotonia, and dystonia were present in all affected individuals, preventing independent ambulation. Likewise, nystagmus was commonly described. In addition, all affected individuals displayed intellectual disability and speech delay. Two siblings further presented with therapy-resistant epilepsy. No major dysmorphic features were found. In two individuals, retinal pigment alterations were noticed. Brain MRI revealed mild cerebellar atrophy and vermian atrophy without other major structural abnormalities in most affected individuals while in one case (Subject 9) bilateral hyperintensities at the nucleus caudatus area were noted (Fig. 1C and Supplementary Fig. 5). No hearing or vision problems were noted and in cases where nerve conduction studies were performed, these were normal. Transmission electron microscopy (TEM) on peripheral blood lymphocytes from Subject 2 and lymphoblastoid cells from Subject 3 revealed more multilayered vesicles compared to control cells (Supplementary Fig. 6).

Exome sequencing did not identify variants in known disease genes but revealed homozygous missense variants in VPS41 (GenBank: NM_014396) that fully segregated with the phenotype in families where segregation analysis was performed (Fig. 1A and Supplementary Figs 2–5). No other likely pathogenic variants were identified. Other than c.853T>C (p.Ser285Pro), which was found twice in a heterozygous state in gnomAD (minor allele frequency: 0.000007), the encountered variants [c.1898G>C (p.Arg633Pro), c.2372G>T (p.Cys791Phe), and c.38A>G (p.Glu13Gly)] have not previously been reported in publicly available genome databases (Fig. 1D). All variants were family-specific except for c.853T>C (p.Ser285Pro), which was encountered in two different families from Saudi Arabia. All missense variants occur in evolutionarily conserved amino acid positions that in silico modelling suggests are likely to disturb protein structure (Fig. 1D and E and Supplementary Fig. 7A and B).

In silico analysis predicts reduced VPS41 function in affected individuals

Protein modelling showed that VPS41 contains an acidic N-terminal extension (residues 1–29), a WD40 β-propeller domain (residues 30–361), a central superhelical structure composed of clathrin heavy-chain repeats (CHCR; residues 362–81) and a C-terminal RING (really interesting new gene) zinc finger (787–852) (Fig. 1D and E). Glu13Gly
attenuates the acidic charge of the N-terminal extension. Glu-rich acidic regions are frequently used for protein localization and ligand recognition (Uversky, 2013) and the mutation might abrogate such functions. Ser285Pro affects the WD40 domain. Ser285 is part of a buried \( \beta \)-strand and a proline in this position would disrupt this secondary structure and decrease shape complementarity within the densely packed protein core. Because of this disruptive effect on the WD40 domain, Ser285Pro might affect ARL8B binding (Khattar et al., 2015). The interaction of VPS41 with ARL8B through its effector SKIP (SifA and kinesin-interacting protein) contributes to lysosomal localization and trafficking and degradation of EGFR (Khattar et al., 2015). Interestingly, when Arl8b is silenced, lysosomal cargo delivery is impaired (Khattar et al., 2015). The Arg633Pro mutation found within the CHCR distorts the \( \alpha \)-helix because the rigidity of proline and its inability to donate a hydrogen bond lead to strong kinks in helices. Finally, residue 791 represents the first cysteine of the zinc finger/RING-type domain and its mutation to Phe is expected to abolish zinc binding and would introduce steric clashes (Matthews and Sunde, 2002). Thus, each of the identified variants is
<table>
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<th>Family 1</th>
<th>Family 2</th>
<th>Family 3</th>
<th>Family 4</th>
<th>Family 5</th>
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<td>c.853T &gt; C</td>
<td>c.1898G &gt; C</td>
<td>c.2372G &gt; T</td>
<td>c.38A &gt; G</td>
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<td>NR</td>
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<tr>
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<td>Hypotonic</td>
<td>Hypotonic</td>
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<tr>
<td>MRI</td>
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<td>Midline vermian atrophy, no iron deposition</td>
<td>Midline vermian atrophy, no iron deposition</td>
<td>Mild vermis hypoplasia or atrophy</td>
<td>Mild vermis hypoplasia or atrophy</td>
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</tbody>
</table>

DTR = deep tendon reflex; NA = not applicable; NR = not reported.
predicted to affect protein domains implicated in mediating ligand interactions, possibly leading to an impact on VPS41 function and stability. The level of weakening or disruption of these interactions possibly explains the diverse patient specific differences (Fig. 1E, Table 1 and Supplementary Table 2).

VPS41 is moderately expressed in the cerebral cortex and cerebellum (Supplementary Fig. 8A), but is expressed highly in Purkinje cells (Supplementary Fig. 8B). Since co-expressed genes are considered more likely to be functionally related and may provide clues about gene function and regulation (Stuart et al., 2003; van Dam et al., 2015), we searched for genes that are co-regulated with VPS41 in over 4000 human RNASeq data (van Dam et al., 2015; Al-Harazi et al., 2019). We identified 105 genes (correlation coefficient ≥ 0.4), of which almost 80%, like VPS41, are expressed in the nervous system (Supplementary Table 3). Functional enrichment and pathway analyses of VPS41 co-expressed genes revealed significant enrichment for terms associated with nervous system development and function, vesicle-mediated transport, cellular component organization, phagosome maturation and autophagy pathways (VPS39, VPS41 and PIK3C3) (all P-values < 0.01) (Supplementary Fig. 9 and Supplementary Tables 4–6).

**VPS41 variants show functional defects in human embryonic stem cells**

To functionally test VPS41 variants in a cellular model, we generated human ESCs knock-out (KO) for VPS41 by CRISPR-Cas9 engineering (Barakat et al., 2018; Perenthaler et al., 2020). A clone was obtained harbouring a homozygous 1 bp insertion (chr7:38908803_38908804insT), leading to a frameshift and premature stop codon (c.110_111insA, N37Kfs*5), resulting in the full absence of all VPS41 as assessed by immunoblotting (Supplementary Fig. 10A–C). VPS41 KO did not alter ESC growth characteristics but resulted in minor flattened morphology compared to parental wild-type ESCs (Supplementary Fig. 10D). Previous work showed that the transcription factor TFE3 can be cytoplasmic or nuclear, depending on its phosphorylation state, and that this is regulated by mTORC1 signalling and lysosomes (Yang et al., 2018; Shin and Zoncu, 2020). In agreement with lysosomal dysfunction (Villegas et al., 2019), the absence of VPS41 resulted in solely nuclear TFE3 localization (Fig. 2A). VPS41 KO cells showed upregulation of the lysosomal protein LAMP2 (Fig. 2C, D and Supplementary Fig. 10E), likely due to nuclear TFE3 localization (Fan et al., 2018). Thus, the absence of VPS41 provided two clear phenotypical differences between wild-type and VPS41 KO ESCs and we used the cellular phenotype of this VPS41 KO background to assess the functionality of wild-type and mutant VPS41 proteins in rescue experiments. We used the expression of V5-VPS41-T2A-GFP fusions, allowing the tracking of transfected cells at the population and single cell level. Fluorescence-activated cell sorting (FACS) analysis 48 h after transient transfection showed a significant reduction in the percentage (Fig. 2E) and intensity (Supplementary Fig. 10F) of GFP+ cells for the Arg633Pro and Cys791Phe variants, whereas transfection efficiency for a spiked-in mCherry control was similar amongst all transfections (Supplementary Fig. 10G). In agreement, immunoblotting confirmed reduced protein expression of these both mutants, as well as a slightly reduced expression of Ser285Pro (Fig. 2C and D). This indicates that the variants encountered result in reduced RNA or protein expression and possibly stability. Whereas transfection of wild-type VPS41 resulted in downregulation of LAMP2 in transfected KO ESCs, expression of Ser285Pro, Cys791Phe, and to a lesser extent Arg633Pro, failed to downregulate LAMP2 at a population level (Fig. 2D). We then moved to single cells and evaluated the cellular distribution of TFE3. Upon transient transfection of wild-type VPS41, cytoplasmic localization of TFE3 was restored in GFP+ cells (Fig. 2A and B). In contrast, transient transfection of Glu13Gly resulted in a significantly reduced rescue of the TFE3 cytoplasmic localization in GFP+ cells, indicating a reduced functionality of this mutant despite normal protein expression (Fig. 2A and B). TFE3 relocalization was similar for Ser285Pro, Arg633Pro, and Cys791Phe in GFP+ cells, indicating that if sufficient overexpression can be achieved, these mutants can promote TFE3 relocalization. Together, these data further support our *in silico* analysis, providing functional evidence that the encountered variants affect VPS41 protein expression levels and result in measurably reduced VPS41 function in human cells.

**Zebrafish disease modelling recapitulates key clinical findings**

To investigate the neurological effects of VPS41 loss *in vivo*, we mutated the single *vps41* homologue in zebrafish, which shares 82% amino acid identity with the human protein, by injection of CRISPR/Cas9 and gRNA ribonucleoprotein complexes into oocytes (Supplementary Fig. 11). CRISPR-mediated mutants (crispants) generated in this way have been shown to exhibit highly efficient mutagenesis, often achieving complete loss of the wild-type target sequence, and can reliably phenocopy existing loss-of-function mutants and thus facilitate rapid and reliable investigations in F0 embryos (Shah et al., 2015; Kuil et al., 2019). Zebrafish *vps41* crispants, having a *vps41* disruption efficiency between 60 and 90%, had normal general morphology in the first 3 days but displayed a severe lack of pigmented melanocytes when compared to control gRNA-injected oocytes (Fig. 3A). Pigmentation defects were partially corrected by injection of wild-type *vps41* mRNA at the 1–2-cell stage (Supplementary Fig. 11). After 5 days, 100% of *vps41* crispants showed a lack of swim bladder inflation—ultimately resulting in a loss of viability—as well as decreased body length and reduced pigmentation both throughout the body and in the retina (Fig. 3A–C). These specific pigmentation phenotypes are
consistent with observations of loss-of-function of other HOPS components (Vps33a/VTI1A in mice and vps11, vps18 and vps39 in zebrafish (Thomas et al., 2011; Zhen and Li, 2015; Berg et al., 2016), while swim bladder defects were also noted in all three zebrafish mutants. Deficiencies in surfactant production or distribution during swim bladder development are believed to contribute to an inability to properly inflate (Chen et al., 2018). Together, these data

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**Figure 2 VPS41 variants contribute to impaired lysosomal functions in vitro.** (A) Immunocytochemistry assessing TFE3 in rescue experiments using transient expression of wild-type (WT) and mutant VPS41-t2a-GFP in VPS41 KO ESC. Transfected cells are marked by GFP and GFP + cells showing no TFE3 relocalization to the cytoplasm are marked with white asterisks. Nuclei are counterstained with DAPI. Scale bars = 50 μm. (B) Quantification of A counting a minimum of 80 GFP + cells from three replicates. ***p = 0.0001, binomial test (expected distribution based on relocalization upon transfection with the wild-type construct). (C) Western blotting detecting VPS41 (endogenous VPS41 99 kDa, V5-tag-VPS41 105 kDa, input 30 μg) and LAMP2 (100–120 kDa, input 20 μg), in wild-type and VPS41 KO ESCs and VPS41 KO ESCs transiently rescued with wild-type or mutant VPS41. (D) Quantification of C. *p < 0.05; **p < 0.01; ***p < 0.001 one-way ANOVA, multiple comparison test of mutant constructs to the wild-type. (E) Percentage of GFP + cells upon transient transfections of VPS41-KO cells with VPS41-GFP plasmid spiked with mCherry, expressing wild-type or mutant VPS41. ***p = 0.0006, ****p < 0.0001, one-way ANOVA, multiple comparison test of mutant constructs to the wild-type; >17,000 cells were analysed per sample.
Figure 3 VPS41 disease modelling in zebrafish recapitulates lysosomal abnormalities and indicates microglial and cerebellar dysfunction. (A) Gross morphology of vps41 crisprant versus control embryos at 5 dpf. Magnified views of the skin and eyes show abnormal melanocyte morphology and reduced pigmentation in both the skin melanocytes and retinal pigment epithelium. (B) Measurements of body length at 3 and 5 dpf, suggesting a reduced growth rate in vps41 crisprants versus controls (n = 15 each). (C) At 5 dpf, swim bladders were not visible in any of the vps41 crisprant embryos, while being fully developed by this time in the majority of controls (n = 50). (D) Schematic of the OKR experiment, indicating the eye angles measured. (E) Representative OKR traces for vps41 crisprant and control embryos under four different conditions. (F) Quantification of saccades produced in response to visual stimuli (n = 6 for each condition). (G) Average saccade amplitude per trial, taken from left eye traces (n = 6 for each condition). (H) LysoTrackerTM staining of 3 dpf embryos, revealing increased acidification of vps41 crisprant brains that is largely concentrated within Mpeg1 + cell (microglia) compartments. Scale bars = 20 μm. (I) Quantification of acidic compartments within microglia in the brain at 3 dpf (6–8 cells counted per embryo, n = 4 embryos each). (J) Compared to control injected embryos at 5 dpf, vps41 crisprants show a remarkably increased level of acidity across the entire brain, detected using LysoTrackerTM, that is significantly reduced when embryos are additionally injected with wild-type vps41 mRNA at the 1–2 cell stage. (K) Quantification of the LysoTrackerTM signal intensity across the whole brain of 5 dpf control, wild-type vps41 and vps41 + mRNA-injected embryos (n = 5 controls, n = 9 vps41, and n = 7 vps41 + mRNA). Data-points in B, F, G and I are individual measurements (circles) and mean (bars) and error bars indicate SEM. CW = clockwise; CCW = counterclockwise; cpd = cycles per degree; dpf = days post-fertilization. *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001.
further support a role for Vps41 and the HOPS complex in the development and function of LROs, including both melanosomes (the pigment-producing organelles within melanocytes and the retinal pigment epithelium) and the surfactant-producing organelles involved in the teleost surfactant production system (Chen et al., 2018; Delevoye et al., 2019).

Abnormal eye movements are associated with cerebellar injury and can be diagnostic of ataxia (Cogan et al., 1982). Thus, we analyzed the optokinetic response (OKR) of vps41-mutated zebrafish larvae, using a well-established zebrafish assay for cerebellar dysfunction (Namikawa et al., 2019). In contrast to control injected larvae, which demonstrated normal responses to visual stimuli, vps41 crispants showed a dramatically reduced response and, when present, saccades had both reduced amplitude and frequency (Fig. 3D–G), indicating defects in eye movement control and suggesting cerebellar dysfunction. Spontaneous eye movements in the absence of stimulation were observed, removing the possibility that crispants were simply unable to move their eyes. The underlying mechanisms of reduced visual performance seen in vps41 crispants remain unclear. Nevertheless, data from both vps11 and vps39 mutant zebrafish indicate that ineffective digestion of material phagocytosed by the retinal pigment epithelium could contribute by hindering proper development and function of photoreceptor cells, further pointing to a defect in LRO function (Schonthaler et al., 2007; Thomas et al., 2011).

While no differences were seen in the gross morphology or size of vps41 crispant brains (Supplementary Fig. 11C), we sought to further investigate potential underlying neurologic abnormalities that could help understand the progressive neurodevelopmental phenotypes seen in human disease. During early neurodevelopment microglia are highly endocytic and phagocytic, functions that rely heavily on efficient membrane trafficking and LRO function, the dysfunction of which is readily observed in zebrafish larvae (Berg et al., 2016; Kuil et al., 2019). Early defects in microglial function have been linked to a wide range of neurological diseases, including lysosomal storage diseases, where lysosomal dysfunction in microglia has been implicated as a primary driver (Kuil et al., 2019). Thus, to assess whether microglial dysfunction may play a role in VPS41 variant-driven pathology, we used neutral red staining to observe microglia during development of vps41 crispants and control larvae (Supplementary Fig. 11B). This revealed that vps41-deficient microglia have distinct morphological abnormalities suggestive of larger, or possibly more numerous, lysosomal compartments compared to microglia from control larvae. To more carefully dissect this phenotype, we sought to visualize lysosomes in vivo in the developing brain using LysoTracker™ in zebrafish where microglia were transgenically tagged with green fluorescent protein (GFP). Confocal imaging revealed that vps41 deficiency resulted in a dramatic increase in LysoTracker™ signal throughout the entire brain, in addition to highly amoeboid, swollen microglia with large, supernumerary acidic compartments in contrast to those in control injected zebrafish (Fig. 3H and I).

Importantly, LysoTracker™ signal in the brain was significantly abrogated by the injection of wild-type zebrafish vps41 mRNA into crispant oocytes at the 1–2-cell stage (Fig. 3J and K), again supporting the specificity of our observations and utility of this model for the study of vps41 function. Together, these data further support a role for lysosomal dysfunction upon vps41 disruption and, in parallel with observations of macrophage defects seen in vps39 mutant zebrafish (Schonthaler et al., 2007), suggest disruption of microglial function. However, while the largest LysoTracker+ compartments both in crispants and controls appeared within mpeg+ cells, it is important to note that a substantial increase in LysoTracker+ compartments inside cells other than microglia was also observed only in vps41 crispants. This supports a more global LRO defect that likely impacts neural progenitors as well, contributing to developmentally seeded neurological defects that are likely further compounded by microglial dysfunction.

**Discussion**

Here we establish that VPS41 bi-allelic mutations cause cerebellar ataxia. As the identified variants fail to completely rescue the lysosomal dysfunction seen in VPS41 KO ESCs, and VPS41 deficiency causes lysosomal abnormalities in multiple brain cell types in the developing zebrafish brain, it is possible that membrane trafficking defects in neurons underlie cerebellar atrophy. Additionally, gene network analysis suggests that VPS41 dysfunction may disrupt a number of signalling pathways linked to neuronal development and maintenance of brain function (Supplementary Fig. 9 and Supplementary Tables 4–6). Nearly all spinocerebellar ataxias have been characterized by Purkinje cell deterioration and cerebellar atrophy (Huang and Verbeek, 2018). A scenario of reduced viability of Purkinje cells is also plausible for VPS41-related ataxia and may help explain cerebellar atrophy. Although we did not detect clear differences directly in zebrafish Purkinje cells during the early development of vps41 crispants (Supplementary Fig. 11D), cellular dysfunction is supported by our OKR results and may precede visible deterioration. However, we did observe distinct lysosomal accumulation and abnormalities at a very early stage in microglia. As lysosomal accumulation in microglia inhibits their function and thus is detrimental to brain development and function, combined with the understanding that a loss of microglia has been shown to cause a spectrum of brain abnormalities, including cerebellar hypoplasia (Oosterhof et al., 2019; Burns et al., 2020), further investigation of the role of microglia in the disease pathogenesis is warranted.

It is interesting to note the overlap of some phenotypes observed in models of VPS loss-of-function, yet ours is the first report to explicitly link variants in a HOPS-specific VPS gene to human brain disease. Intriguingly, many of the existing Vps/vps mutant animal models were identified in genetic screens for pigment defects and have been proposed to model Hermansky-Pudlak syndrome (HPS), a disorder
Bi-allelic variants in VPS41 cause ataxia

typed by albinism and bleeding, but typically no neurological abnormalities. Mutations associated with HPS include mutations in genes involved in LRO biogenesis, though to date no VPS mutations have been linked to this disease in humans (Merideth et al., 2020). This suggests that while LRO dysfunction contributes to familiar phenotypes across a spectrum of human diseases, VPS (and specifically HOPS) mutations diverge from other LRO-associated diseases as they appear to more prominently affect the nervous system. VPS mutant animal models may therefore require reassessment and possible reclassification as models of cerebellar dysfunction.

Together, our work provides evidence that genetic defects in VPS41, and thus the HOPS complex, result in dysregulated neurodevelopment in vertebrates, with a predominant ataxia phenotype, driven at least partially through LRO abnormalities. We suggest that screening for VPS41 variants, and perhaps other HOPS members, should therefore be considered among cases of unsolved autosomal recessive ataxia. Our work adds to the increasing knowledge on autosomal recessive cerebellar ataxias by providing evidence that genetic defects in the HOPS-specific subunit VPS41 lead to abnormalities in lysosome-related organelles in multiple cell types, including microglia. This contributes to disturbed neurodevelopment and ataxia, though further studies are needed to clearly dissect the underlying mechanisms of these connections.

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Competing interests


Supplementary material

Supplementary material is available at Brain online.

References
