Astrocyte Subtype Vulnerability in Stem Cell Models of Vanishing White Matter

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Objective: Astrocytes have gained attention as important players in neurological disease. In line with their heterogeneous character, defects in specific astrocyte subtypes have been identified. Leukodystrophy vanishing white matter (VWM) shows selective vulnerability in white matter astrocytes, but the underlying mechanisms remain unclear. Induced pluripotent stem cell technology is being extensively explored in studies of pathophysiology and regenerative medicine. However, models for distinct astrocyte subtypes for VWM are lacking, thereby hampering identification of disease-specific pathways.

Methods: Here, we characterize human and mouse pluripotent stem cell–derived gray and white matter astrocyte subtypes to generate an in vitro VWM model. We examined morphology and functionality, and used coculture methods, high-content microscopy, and RNA sequencing to study VWM cultures.

Results: We found intrinsic vulnerability in specific astrocyte subpopulations in VWM. When comparing VWM and control cultures, white matter–like astrocytes inhibited oligodendrocyte maturation, and showed affected pathways in both human and mouse cultures, involving the immune system and extracellular matrix. Interestingly, human white matter–like astrocytes presented additional, human-specific disease mechanisms, such as neuronal and mitochondrial functioning.

Interpretation: Astrocyte subtype cultures revealed disease-specific pathways in VWM. Cross-validation of human- and mouse-derived protocols identified human-specific disease aspects. This study provides new insights into VWM disease mechanisms, which helps the development of in vivo regenerative applications, and we further present strategies to study astrocyte subtype vulnerability in neurological disease.

Ann Neurol 2019;86:780–792

Astrocytes are increasingly recognized to play a role in the pathophysiology of different neurological diseases and suggested as the target in new therapies.1,2 Astrocytes consist of functionally and morphologically heterogeneous populations of cells that develop at different times and different locations in the central nervous system.3,4 Defects in specific astrocyte subtypes are shown in different neurological disorders,1,2 including epilepsy,5 amyotrophic lateral sclerosis,6 Alzheimer disease,7 and vanishing white matter (VWM).8 VWM is one of the more prevalent leukodystrophies, caused by mutations in EIF2B1–5,9 and specific astrocyte subpopulations, like the astrocytes of the white matter, are affected.8,10,11 This vulnerability in astrocyte subtypes needs to be taken into consideration to better understand pathophysiology and to develop more targeted treatment options. Mouse studies have advanced our

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knowledge on gene function in neural cell development and provide robust and standardized protocols, but do not always recapitulate the cellular or clinical phenotypes of human patients. The discovery of human induced pluripotent stem cells (hiPSCs) greatly enhanced the possibility to study neural disorders with patient-specific cells, but we still lack models to study phenotypes in specific neural subtypes. To identify disease mechanisms for a disorder like VWM, we are in need of iPSC differentiation protocols generating specific astrocyte subtypes.

In this study, we created in vitro models for VWM using hiPSCs and mouse induced pluripotent stem cells (miPSCs; Fig 1). We generated VWM- and control iPSC-derived astrocyte subtypes, using signaling molecules stimulating gray (using fetal bovine serum [FBS]) and white (including ciliary neurotrophic factor [CNTF]) matter. Interestingly, both hiPSC- and miPSC-derived white matter–like astrocytes showed specific vulnerability to VWM mutations. RNA sequencing (RNAseq) analysis on hiPSC-derived white matter–like astrocytes indicated differential gene expression in VWM compared to control astrocytes involved in several cellular mechanisms, including neuronal, mitochondrial, and vasculature-related functioning. Of these pathways, genes involved in the immune system, cell development, and extracellular matrix also came up in miPSC-derived models, demonstrating the strength of cross-species validation in finding disease mechanisms induced by monogenic changes. These new insights can help development of more cell- or pathway-targeted therapies in VWM and other astrocyte-associated diseases.

**Materials and Methods**

**hiPSC Differentiation toward Astrocytes**

The institutional review board of Amsterdam UMC, Vrije Universiteit Amsterdam approved this study with waiver of informed consent. hiPSCs were generated from fibroblasts of 2 VWM patients with *EIF2B5* mutations (one 3-year-old male

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**FIGURE 1:** Experimental overview. (A) Fibroblasts from human and mouse, both vanishing white matter (VWM) genotype and control/wild type, were reprogrammed to induced pluripotent stem cells (iPSCs), and differentiated to glial cells for various assays. (B) The iPSCs were further differentiated to mouse and human gray matter and white matter astrocyte subtypes. The different astrocyte subtypes were used as an in vitro model for VWM, in which oligodendrocyte maturation and apoptosis, morphology, proliferation rate, and mRNA expression were studied. qPCR = quantitative polymerase chain reaction.
with 1484A>G mutation; one 9-year-old female with 806G>A mutation.) and 3 controls (males of 44, 46, and 74 days old) using a polycistrionic lentiviral transduction of the Yamanaka factors.14 Cells were maintained and characterized as previously described,15 by alkaline phosphatase staining, immunocytochemistry, real-time polymerase chain reaction (RT-PCR) analysis, RNAseq PluriTest, and germ layer embryoid body (EB) differentiation assay, and karyotyped or tested on the Infinium Global Screening Array for DNA abnormalities. hiPSCs were differentiated toward human astrocytes as described previously.15 To generate human astrocyte subtypes, glial precursors were generated as described previously,15 after which medium was switched to N2B27-vitA medium supplemented with either epidermal growth factor (20ng/ml), fibroblast growth factor 2 (FGF2; 20ng/ml), and CNTF (20ng/ml) for human white matter–like astrocytes (human WM astrocytes), or with 10% FBS for human gray matter–like astrocytes (human GM astrocytes) for another 2 passages in 15 days. All differentiations are performed at least 3 times per iPSC clone, with at least 2 iPSC clones per donor.

miPSC Differentiation toward Glial Cells and Astrocytes

miPSCs from wild-type (wt), Eif2b4Arg191His/Arg191His (2b4), and Eif2b4Arg191His/Arg191His (2b4; hu) mice were made from fibroblasts with viral transduction of the Yamanaka factors.14 Cells were maintained in 2i medium on gelatin-coated plates. miPSCs were characterized by using immunocytochemistry, PCR analysis, EB differentiation assay, and teratoma formation assay. miPSCs were differentiated via EB formation in basal medium followed by neural induction in N2-based medium supplemented with retinoic acid (0.2μM) and purmorphamine (1μM). After 8 days, EBs were plated in neural maintenance medium (N2-based) supplemented with 20ng/ml FGF2 for 12 days.

For glial differentiation, the glial progenitor cells were subsequently cultured in mouse neural maintenance medium supplemented with 30ng/ml T3 and 10ng/ml NT3 from day 12 on. At day 18, cells were used for analysis. For mouse astrocyte subtypes cells were cultured in mouse neural maintenance medium supplemented with 10% FBS for mouse gray matter–like astrocytes (mouse GM astrocytes) or 10ng/ml CNTF for mouse white matter–like astrocytes (mouse WM astrocytes) from day 12 onward. To obtain purer astrocyte cultures, cells were passed every 7 days for an additional 2 passages. At day 32, cells were used for analysis. All immunostaining and PCR analysis of miPSC-derived cells are an average of at least 3 independent differentiations and based on 3 to 4 iPSC lines per genotype.

Isolation and Culture of Primary Cells

Primary astrocytes and oligodendrocyte precursor cells were isolated from forebrain of embryonic day 18 mice as described previously.10 Oligodendrocyte precursor cells were sorted on expression of platelet-derived growth factor receptor α with magnetically activated cell sorting according to the manufacturer’s protocol (Miltenyi Biotec, Bergisch Gladbach, Germany).10

Analysis

Cells were analyzed by immunocytochemistry as previously described.15 Primary antibodies targeted glial fibrillary acidic protein (GFAP; Sigma, St Louis, MO; G3893, 1:1,000), GFAP (Dako, Carpinteria, CA; Z0334, 1:1,000), nestin (BD Biosciences, Franklin Lakes, NJ; 611658, 1:500), S100β (Proteintech Group, Rosemont, IL; 15146-1-AP, 1:1,000), Olig2 (for mouse, gift of J. H. Alberts, 1:500), Olig2 (for human, Millipore, Billerica, MA; AB9610, 1:500), Sox9 (Cell Signaling Technology, Danvers, MA; 82630, 1:500), myelin basic protein (MBP; Covance, Princeton, NJ; SMI-99P, 1:2,000), CD44 (Developmental Studies Hybridoma Bank, Iowa City, IA; H4C4, 1:250), ID3 (Cell Signaling Technology, 9837, 1:250), ezrin (Santa Cruz Biotechnology, Santa Cruz, CA; sc-32759, 1:450), myelin oligodendrocyte glycoprotein (MOG; Millipore, MAB5680, 1:500), cleaved caspase 3 (Cell Signaling Technology, 9661, 1:400), OCT3/4 (Santa Cruz Biotechnology, sc-5279, 1:1,000), nanog (Abcam, Cambridge, MA; AB80892, 1:1,000), Lin28a (Cell Signaling Technology, 3978s, 1:1,000), β-tubulin-III (R&D Systems, Minneapolis, MN; MAB1195, 1:1,000), α-smooth muscle actin (Progen, Heidelberg, Germany, 61001, 1:1,000), α-fetoprotein (R&D Systems, MAB1368, 1:1,000), Tra-1-60 (Santa Cruz Biotechnology, sc-21705, 1:200), and SSEA4 (Developmental Studies Hybridoma Bank. [SSEA-4]-s, 1:50).

Bromodeoxyuridine (BrdU) assay was performed (2 independent differentiations of 4 human astrocyte lines per genotype) by 2-hour incubation of BrdU at 37°C and stained with anti-BrdU antibody (Abcam) according to the manufacturer’s protocol. RNA was collected with TRIzol (Invitrogen, Carlsbad, CA). RNA isolation, cDNA synthesis, and PCR analysis were performed as described previously (see Supplementary Table 1 for primer sequences).15 Calcium imaging of glutamate uptake in human astrocytes was performed and analyzed as described previously.15 For reactivity assay, human astrocytes were treated with either 10μg/ml polyinosinic:polycytidylic acid (Tocris Bioscience, Bristol, UK) or dH2O for 21 hours, after which RNA was isolated and analyzed as described previously.15

For RNAseq, an Illumina (San Diego, CA) TrueSeq stranded mRNA kit was used according to the manufacturer’s protocol. Sequence fragments were aligned to the reference genome (mouse: GRCh38, annotated genes GENCODE vM12; human: GRCh38, annotated genes GENCODE v25). Only genes on autosomal chromosomes were extracted, and all genes with a transcripts per kilobase million of <1 in 50% of samples were excluded. Differential expression analysis was performed using R package DESeq2. Genes with Bonferroni corrected p < 0.05 were considered significantly differentially expressed genes (DEGs) and used for gene-set enrichment analysis using Gene Ontology (GO) terms. For human samples, analysis was not done with covariate factors, which included individual, the number of clones per individual, and the number of differentiation repetitions per clone.

Statistical Analysis

For all experiments, results were considered significant at α = 0.05 after multiple test correction. Apart from the RNAseq data, all other data are analyzed with IBM (Armonk, NY) SPSS
and the severely affected 2b4 astrocytes (~15% were differentiated toward glia cultures containing both
was used.
data, 1-way analysis of variance with Dunnett post hoc tests
markers GFAP, nestin, Id3, CD44, and SOX9 by immuno
were smaller, and had higher expression of S100b, ALDOC, AQP4, GLAST, GFAP, and CD44 by RT-PCR
(data not shown). Functionality of human astrocytes was
demonstrated using calcium imaging, showing glutamate
uptake in both control and VWM lines (data not shown).
To confirm increased proliferation in VWM astrocytes as
was described earlier for primary human astrocytes,8 we
performed a BrdU incorporation assay. The VWM astrocytes
showed a significantly higher percentage of proliferating
cells compared to control human astrocytes (independent
t3 = 2.56, p = 0.042). To show that human astrocytes
can mediate an oligodendrocyte maturation defect via secre-
tion of factors in media as suggested by mouse studies,10
primary mouse oligodendrocyte precursor cells were
cultured in medium conditioned by control and patient
iPSC-derived astrocytes. Media conditioned by VWM
astrocytes significantly impaired oligodendrocyte maturation,
as measured by the percentage of MBP-/Olig2-positive
cells (independent t3 = 2.34, p = 0.049). As hyaluronic acid
has been previously described to inhibit oligodendrocyte
maturation16 and is increased in brains of VWM patients,17
media conditioned by human astrocytes was treated with
hyaluronidase. Hyaluronidase-treated conditioned medium
significantly increased oligodendrocyte maturation in VWM
cultures (independent t3 = −8.62, p = 0.002), which recov-
ered toward control levels. Altogether, our results demon-
strate that hiPSC models recapitulate findings in both
primary human cells and in miPSC cell models. Furthermore,
we confirmed an astrocyte-intrinsic defect in VWM using
hiPSC models.

miPSC-Based Models Show Selective Involvement of Astrocytic Subtypes in VWM
An increasing number of studies confirm heterogeneity
among astrocytes, such as white and gray matter astro-
cytes. However, we lack iPSC-based differentiation proto-
cols generating specific subtypes. CNTF and FBS are both
used in astrocyte differentiation protocols.18,19 Where
CNTF-reactive astrocytes are predominantly found in the
white matter,20 FBS administration is standardly used to
maintain cortical astrocytes,21 leading to a flat morphology
that is common for gray matter astrocytes in culture. Here
we used CNTF- and FBS-based media to generate mouse
WM astrocytes and mouse GM astrocytes, respectively.
The mouse WM astrocytes had many thin protrusions,
were smaller, and had higher expression of Glast, S100b,
and Gfap (data not shown) compared to the larger and
rounder mouse GM astrocytes (Fig 4). To assess functional
defects in the 2 astrocyte subtypes in VWM, wt mouse pri-
mary oligodendrocyte precursor cells were cultured in
media conditioned by mouse WM and GM astrocytes,
derived from wt, 2b4, and 2b4 miPSCs. Oligoden-
drocyte maturation was not affected in conditioned

Results

miPSC-Based Models Recapitulate Oligodendrocyte Precursor Cell Maturation Inhibition by VWM Astrocytes
Previous studies showed that astrocytes isolated from
VWM mouse models impair primary oligodendrocyte pre-
cursor cell maturation via secreted factors, suggesting that
astrocytes are the primary affected cell type in VWM.10
To show that iPSC-derived cell models recapitulate findings
in primary cell models and to confirm that astrocytes cause cellular defects in VWM, we developed an in vitro
miPSC model based on previous experiments with primary
mouse cells. We generated miPSCs from wt, 2b4, and the severely affected 2b4 mouse. The miPSCs
were differentiated toward glia cultures containing both
astrocytes (-15–25% GFAP-positive cells) and oligoden-
drocytes (-30–40% Olig2-positive cells; Fig 2). Similar to previous primary mouse studies, oligodendrocyte maturation
was significantly impaired in VWM cultures, as was demonstrated by decreased ratios of MBP-/Olig2-positive cells
in 2b4 mice (Figure 2, p = 0.005; post hoc Dunnett
t wt vs 2b4, p = 0.039) and MOG-4,6-diamidino-
dendrocyte maturation was not affected in conditioned

VWM Patient iPSC-Based Models Confirm Intrinsic Defects in VWM Astrocytes
To recapitulate findings with hiPSC models, we generated
hiPSCs from VWM patients and healthy control donor
fibroblasts. To generate astrocytes, we differentiated control
and VWM hiPSCs toward astrocytes according to protocols described earlier.15 We characterized the human
astrocytes for expression of the astrocyte-associated
markers GFAP, nestin, Id3, CD44, and SOX9 by immuno-
cytochemistry (Fig 3), and NESTIN, BLBP, S100b,
**FIGURE 2:** Wild-type (wt) astrocytes rescue the maturation defect of vanishing white matter (VWM) mouse induced pluripotent stem cell (miPSC)-derived oligodendrocytes. (A–C) Glial differentiation contained glial fibrillary acidic protein (GFAP)-positive astrocytes, as shown by immunocytochemistry (A, B) and quantification (C). (D–I) Oligodendrocyte maturation was addressed in wt, 2b5^ho, and 2b4^ho2b5^ho glia by immunostaining for myelin basic protein (MBP), myelin oligodendrocyte glycoprotein (MOG), and Olig2. Cell counts confirmed the decrease in the number of MBP-positive (G) and MOG-positive cells (H) in VWM without a decrease in the number of Olig2^+ cells (I). Percentages of positive cells compared to the number of Olig2-positive cells are shown for MBP; H and I show the percentage of total number of 4,6-diamidino-2-phenylindole (DAPI)-positive cells (wt, n = 3; 2b5^ho, n = 3; 2b4^ho2b5^ho, n = 3). (J–O) To investigate the effect of healthy astrocytes on the oligodendrocyte precursor cell maturation defect, wt, 2b5^ho, and 2b4^ho2b5^ho glia were grown on a monolayer of wt primary mouse astrocytes, and oligodendrocyte maturation was assessed using an immunostaining for MBP, MOG and Olig2. The oligodendrocyte maturation was no longer reduced in VWM cultures, as demonstrated by the absence of significant differences in the MBP/Olig2 (M) and MOG/DAPI (N) ratios between wt, 2b5^ho, and 2b4^ho2b5^ho cultures. The Olig2/DAPI ratio was slightly, but not significantly, increased in 2b5^ho cultures (O; wt, n = 3; 2b5^ho, n = 3; 2b4^ho2b5^ho, n = 4). Scale bars = 50μM. *Significant at p < 0.05. Bars in C, G–I, M–O represent mean ± standard error of the mean. NS = not significant.
Astrocyte differentiation of hiPSC

A–F) Immunostaining showed expression of astrocyte markers glial fibrillary acidic protein (GFAP; A, D), nestin and Id3 (B, E), and CD44 and SOX9 (C, F) in both control (A–C) and VWM (D–F) human astrocyte cultures. To quantify proliferation of the astrocytes, bromodeoxyuridine (BrdU) labeling was performed. (G, H) Representative control (G) and VWM (H) astrocyte lines are shown. Quantification of proliferation was assessed using the percentage of BrdU-labeled cells of the total population of cells (4,6-diamidino-2-phenylindole [DAPI]) in control human iPSC (hiPSC)-derived astrocytes. (I) VWM astrocytes showed increased proliferation compared to control astrocytes. (J, K) Immunostaining for Olig2 and myelin basic protein (MBP) of primary wild-type embryonic day 18 mouse oligodendrocyte precursor cells cultured for 7 days in media conditioned by hiPSC-derived astrocytes in representative examples of VWM and control cultures is shown. Maturation of the oligodendrocyte precursor cells was assessed using a ratio of MBP-positive cells of the total Olig2-positive population, in the presence of media conditioned by control and VWM iPSC-derived astrocytes. Astrocyte conditioned media were treated with either vehicle or hyaluronidase (HAse). (L) Oligodendrocyte maturation was impaired in VWM compared to control cultures, and HAse rescued oligodendrocyte maturation comparable to control level. Scale bars = 100μm. *p < 0.05, **p < 0.01.
FIGURE 4: Mouse white matter (WM) and gray matter (GM) astrocytes are differentially affected by vanishing white matter (VWM) mutation. (A, B) Immunostaining for cytoplasmic cell surface marker ezrin of mouse induced pluripotent stem cell (miPSC)-derived WM (A) and GM astrocytes (B) showed morphological differences between the cells. (C–H) To assess functional defects of VWM mouse WM and GM astrocytes, wild-type (wt) mouse primary oligodendrocyte precursor cells were cultured in conditioned medium collected from wt (n = 3), 2b5<sup>+</sup> (n = 4), and 2b4<sup>+</sup>2b5<sup>+</sup> (n = 4) iPSC-derived GM or WM astrocytes (respectively GM and WM in I–K). Oligodendrocyte maturation was quantified as the percentage of myelin basic protein (MBP)-positive oligodendrocyte precursor cells were cultured in conditioned medium collected from wt (n = 3), 2b5<sup>+</sup> (n = 4), and 2b4<sup>+</sup>2b5<sup>+</sup> (n = 4) iPSC-derived GM or WM astrocytes (respectively GM and WM in I–K). Oligodendrocyte maturation was quantified as the percentage of myelin basic protein (MBP)-positive oligodendrocytes of the number of Olig2-positive cells. (I, J) This percentage was significantly decreased in 2b4<sup>+</sup>2b5<sup>+</sup> GM medium compared to wt WM medium and 2b4<sup>+</sup>2b5<sup>+</sup> GM medium (I), whereas the percentage of Olig2-positive cells was unchanged between the conditions (J). To assess apoptosis in the oligodendrocyte cultures, a cleaved caspase 3 (CC3) assay was conducted in oligodendrocyte cultures with conditioned medium of GM and WM astrocytes of wt (n = 4), 2b5<sup>+</sup> (n = 4), and 2b4<sup>+</sup>2b5<sup>+</sup> (n = 4). (K) The percentage of CC3-positive cells of 4,6-diamidino-2-phenylindole (DAPI)-positive cells was significantly decreased in VWM compared to control WM cultures. *p < 0.05, **p < 0.01. Scale bars = 50 μm. Bars in I–K represent mean ± standard error of the mean. (L) Volcano plot of differential expression analysis between wt (n = 4) and 2b4<sup>+</sup>2b5<sup>+</sup> (n = 4) mouse WM astrocytes with significant differentially expressed genes (DEGs). Significant DEGs are colored in red and are labeled. (M) Significantly enriched Gene Ontology terms for DEGs in different categories.
medium of VWM GM astrocytes. However, the mouse WM astrocyte conditioned medium from the 2b4ho2b5ho mutants significantly decreased the percentage of MBP-/Olig2-positive cells compared to mouse WM astrocyte conditioned medium from controls (F2, 8 = 6.23, p = 0.023; post hoc Dunnett wt vs 2b4ho2b5ho, p = 0.014) and mouse GM astrocyte conditioned medium from 2b4ho2b5ho mutants (t5 = −6.111, p = 0.009). The percentage of Olig2-positive cells was unchanged between conditions. In the cultures containing conditioned media of mouse WM astrocytes, the percentage of oligodendrocyte precursor cells positive for apoptotic marker cleaved caspase 3 was significantly reduced in the 2b5ho and 2b4ho2b5ho mutants compared to controls (F2, 9 = 9.231, p = 0.007, post hoc Dunnett wt vs 2b5ho, p = .014; wt vs 2b4ho2b5ho, p = 0.006). No significant differences in the percentage of cleaved caspase 3–positive cells were observed in the GM condition. Altogether, these findings demonstrate that mouse WM astrocytes show a higher vulnerability to VWM mutations.

Because mouse WM astrocytes are selectively affected by VWM mutations, we performed transcriptome analysis on wt and 2b4ho2b5ho cultures to identify DEGs in the affected astrocytes. Interestingly, only a selected number of genes were significantly affected by VWM mutations; in total, 13 genes were significantly differentially expressed between the wt and 2b4ho2b5ho WM astrocytes (Supplementary Table 2), as labeled in Figure 4L. Based on enrichment analysis for DEGs with GO terms and literature,22–25 DEGs were overrepresented in the categories “Immune System,” “Development and Proliferation,” “Extracellular Matrix,” “RNA Polymerase II Transcription,” and “P53 Mediated Signaling” (see Fig 4L), suggesting that these processes are differentially regulated between VWM and wt mouse WM astrocytes.  

**hiPSCs Differentiate into Distinctive Astrocytic Subpopulations**

To show astrocyte subtype-specific abnormalities in human VWM cells, we used CNTF- and FBS-based media to generate human WM astrocytes and human GM astrocytes, respectively. Similar to mouse cultures, human WM astrocytes presented as smaller cells with many and thin protrusions compared to the rounder and larger human GM astrocytes (Fig 5A, B) and showed significantly increased expression in astrocyte markers including SOX9, NESTIN, and BLBP (data not shown). Both astrocyte subtypes showed appropriate upregulation of reactivity-related genes in response to cellular stressor polyinosinic-polycytidylic acid (data not shown), but no differences in reactive response were observed. As, to our knowledge, there are no transcriptome profiles of cultured, or purified, human gray and white matter astrocytes, we performed whole-genome transcriptome analysis on our human WM and GM astrocyte subtypes. Figure 5C shows a volcano plot of DEGs in human WM compared to human GM astrocytes. In total, 346 genes were significantly differentially expressed between the subtypes (Supplementary Table 3). Enrichment analysis with GO terms (Molecular Functions) on the DEGs between human WM and GM astrocytes could be classified in the categories “Receptor/Ligand Activity,” “Extracellular Space,” “Enzymatic Activity,” “Immune System,” and “Cytoskeleton” (see Fig 5D). These findings show that we generated 2 human astrocyte subpopulations in vitro: human WM and GM astrocytes, with specific morphological characteristics and differential expression profiles.

**VWM Mutations Predominantly Affect White over Gray Matter Astrocyte Subtype in hiPSC-Based Cultures**

To perform unbiased differential expression analysis, we performed transcriptome analysis of VWM and control human WM astrocytes. In total, 63 genes were significantly differentially expressed between the VWM and controls (Supplementary Table 4). A volcano plot of covariate-corrected DEGs in VWM compared to controls is shown in Figure 6A. When comparing VWM and control human GM astrocytes, 37 DEGs were detected (Supplementary Table 5), which is almost half the number of DEGs found in the white matter subtype. A volcano plot presents the covariate-corrected DEGs in VWM compared to control human GM astrocytes in Figure 6B. Enrichment analysis of GO terms (Biological Processes) on the DEGs between control and VWM WM astrocytes could be categorized as “Immune System,” “Extracellular Space,” “Cell Development,” “Neuronal Functioning,” and “Vasculature-Related,” whereas the DEGs between control and VWM GM astrocytes did not show significant enrichment in the latter 2 categories (see Fig 6). Because these results demonstrate that the human WM astrocytes presented a more broadly modulated transcript profile by a VWM genotype than the human GM astrocytes, the differences in the human WM astrocyte cultures were further investigated. The VWM human WM astrocytes showed different morphologies compared to controls, although these did not reach statistical significance when quantified using automated Columbus software (PerkinElmer, Waltham, MA). The VWM cells were rounder (2-sample t4 = 0.62, p = 0.58), showed a larger surface area (2-sample t9 = 1.78, p = 0.15), and had a reduced perimeter corrected for the surface area (2-sample t4 = −1.00, p = 0.37) compared to control cells. Furthermore, expression analysis for astrocyte-associated markers by quantitative PCR analysis showed that VWM
Human WM astrocytes had a significantly higher expression of AQP4 (2-sample $t_S = 2.94, p = 0.032$) and NESTIN (2-sample $t_S = 3.61, p = 0.015$). There was no difference in the expression of cleaved caspase 3 in oligodendrocyte precursor cells that were cultured in medium conditioned by VWM or control human WM astrocytes, or for human GM astrocyte conditioned media. Together, these findings demonstrate that human WM astrocytes were more profoundly affected by the VWM genotype than human GM astrocytes.
FIGURE 6: Vanishing white matter (VWM) human astrocyte subtypes differ in morphology and mRNA expression level from control human astrocyte subtypes. (A, B) Differential expression analysis revealed significant differentially expressed genes (DEGs) between VWM and control in human white matter (WM) astrocytes (A; control, n = 3; VWM, n = 4) and human gray matter (GM) astrocytes (B; control, n = 4; VWM, n = 4). The volcano plots indicate significant DEGs in red, with the 15 most significantly up- or downregulated DEGs labeled. (C) Significantly enriched Gene Ontology (GO) terms classified the DEGs in different categories of biological processes. (D, E) Morphology of control (D) and VWM (E) human WM astrocytes was visualized using CD44 and SOX9 immunostaining. (F–H) Morphological analysis showed differences in roundness (F), area (G), and perimeter corrected for area (H) between VWM and control human WM astrocytes based on the CD44 immunostaining. (I) A quantitative polymerase chain reaction on VWM human WM astrocytes (n = 4) showed differential expression of S100B, SLC25A18, AGP4, MLC1, SOX9, CD44, NESTIN, ALDH1L1, ALDOC, SLCT1A2, AQP4, GFAP, and BLBP relative to control human WM astrocytes (n = 4). *p < 0.05. (J) The cleaved caspase 3 (CC3)-positive percentage of the 4,6-diamidino-2-phenylindole (DAPI)-positive cells was determined in oligodendrocyte cultures containing media conditioned by GM or WM astrocytes derived from control (n = 4) and VWM (n = 4) lines. Bars represent mean ± standard error of the mean.
Discussion

In this study, we examined astrocyte subtype vulnerability in VWM. We generated functional astrocytes through differentiation protocols for control and VWM white and gray matter–like astrocyte subtypes from hiPSCs and miPSCs, using respectively CNTF or FBS supplementation. VWM astrocytes showed intrinsic defects. The WM astrocytes in both mouse and human models confirmed that specific astrocytic subtypes are more vulnerable for the VWM genotype. Pathways that were differentially regulated in VWM WM astrocytes involved immune system and extracellular matrix. Only in VWM patient models were changes in pathways related to neuronal functioning and vasculature identified. Cross-validation in different species models provided new insights into disease pathways in specific astrocyte subtypes in VWM, thereby helping development of new treatment strategies.

Astrocytes play a central role in pathology in VWM. The presented astrocytes derived from mouse and human iPSCs recapitulate earlier findings in mouse and human, such as increased proliferation, morphological abnormalities, and induction of decreased oligodendrocyte maturation. Interestingly, WM astrocytes are more vulnerable to VWM mutations than GM astrocytes, as indicated before in patient tissue. To our knowledge, we are the first to present VWM iPSC models involving subtype-specific astrocytes. Using mouse cultures, we demonstrated that WM astrocytes, but not GM astrocytes, inhibited oligodendrocyte maturation. This effect was not mediated via increased apoptosis of oligodendrocytes, as medium conditioned by VWM mouse WM astrocytes decreased the percentage of apoptotic cells compared to control astrocytes. This effect was not present in cultures containing medium conditioned by human WM astrocytes. This finding as well as other discrepancies between the mouse and human iPSC-based models may represent species-specific features. For example, earlier studies indicated that hyaluronic acid was immensely increased in brains of VWM patients, but only enhanced to some extent in severely affected VWM mice. Moreover, hyaluronic acid was not a clear determinant in disease phenotypes in primary cultures of VWM mouse cells. In the brain, hyaluronic acid is mainly produced by astrocytes and is known to inhibit oligodendrocyte maturation, although a contribution of a neuronal and an oligodendrocyte component to elevated hyaluronic acid in the VWM patient brain cannot be excluded. Together, our findings show that iPSCs can be used to study intrinsic differences between astrocyte subtypes and to identify shared and unique disease mechanisms between species.

Transcriptome analyses on mouse and human iPSC cultures both suggest involvement of immune system and extracellular matrix in VWM pathology. In mouse cultures, only 13 DEGs between VWM and control were found, of which some genes have been related to VWM disease mechanisms previously. Cdi4 encodes a transmembrane receptor that regulates cellular responses to hyaluronic acid, and was significantly upregulated in VWM compared to control mouse WM astrocytes. Previous studies showed an increase in CD44 expression in VWM patient white matter astrocytes in postmortem tissue. Furthermore, VWM mouse cultures showed decreased Gli1 expression, which codes for a transcription factor that is a direct target of sonic hedgehog (SHH) signaling. SHH regulates proliferation of neural progenitor cells together with the intracellular receptor smoothened (SMO) and has neuroprotective effects. Although not significant, Shh and Smo levels were also decreased (2–2.5-fold) in VWM mouse WM astrocytes. A recent study showed an impaired SHH pathway in primary astrocytes from the Eif2b7R132H/R132H mouse, confirming the involvement of the SHH pathway in VWM. Interestingly, VWM mouse cultures also showed increased Ets1 expression. Ets1 is a transcription factor that is involved in the T-cell immune response and was shown to be upregulated in astrocytes surrounding white matter lesions in a mouse model for multiple sclerosis. These findings suggest that the SHH pathway and immune response are interesting targets for VWM.

In addition to the pathways described for the mouse cultures, human WM astrocyte transcripts showed differences between VWM and controls that were associated with GO terms that we categorized as “Neuronal Functioning” and “Vasculature-Related.” The DEGs related to neuronal functioning include upregulation of GRID1, GRIA2, and GRIN2A in VWM, which encode subunits of membrane ionotropic N-methyl-D-aspartate and α-amino-3-hydroxy-5-methyl-4-isoxazolpropionic acid receptors. A downstream effect of activation of these receptors is stimulation of the adenosine triphosphate (ATP)-driven Na⁺ pump. This induces glycolytic upregulation of ATP and lactate in astrocytes, as well as regulation of the extracellular potassium concentration, because the ATP-driven Na⁺ pump actively pumps potassium into the cells. The channel protein AQP4 is regulated by extracellular potassium levels and involved in water permeability of astrocytes, which can lead to neurotoxicity and myelin defects when dysregulated. Interestingly, VWM human WM astrocytes showed increased AQP4 levels. DEG analysis also included increased expression of SLC6A1, SLC6A11, and GABBR2 in VWM, which encode major transmembrane γ-aminobutyric acid (GABA) transporters GAT-1, GAT-3, and GABA receptor subunit 2, respectively. Increased extracellular levels of GABA, released by interneurons for example, lead to astrocytic Ca²⁺ elevations mediated by GABAb receptor activation. This may further lead to astrocytic glutamate release, thereby potentiating inhibitory synaptic transmission, and to astrocytic ATP/adenosine-mediated heterosynaptic suppression affecting excitatory transmission. This confirms earlier findings that neuronal dysfunction in VWM should not be overlooked.
Our human transcriptome data furthermore showed a number of DEGs involved in mitochondrial functioning, including CTGF, CYP1B1, NTRK2, CNRI, TBX2, and MT2A. CNRI, driven by PAX3 expression, both of which are upregulated in VWM human WM astrocytes, is known to regulate mitochondrial functioning and protects against neuroinflammation resulting from oxidative stress.\textsuperscript{35–37} TBX2 was also upregulated in VWM human WM astrocytes. In astroglioma cell lines, TBX2 has been shown to promote proliferation, to inhibit cleaved caspase 3–mediated apoptosis by stimulating mitochondrial fission, and to increase mitochondrial DNA content.\textsuperscript{38} In contrast to the protective effect of TBX2, our VWM human WM astrocytes also showed a decreased expression of MT2A, an antioxidative, anti-inflammatory, and anti-apoptotic metallothionein secreted by astrocytes that is well known for its neuroprotective properties.\textsuperscript{39} Mitochondrial mechanisms involved in VWM pathophysiology corroborate recent findings in primary astrocytes of Eif2b5\textsuperscript{R132H/R132H} mice indicating that an impaired oxidative phosphorylation in these cells is compensated by increased mitochondrial content and glycolysis.\textsuperscript{39} Thus, these pathways deserve further study to determine whether these can target for therapeutic strategies.

In conclusion, the transcript analysis confirmed pathways previously implicated in VWM, such as SHH and extracellular matrix–related proteins. Additionally, the immune system presented itself as a potential target. Interestingly, human specific signaling pathways that emerged suggested interaction with neuronal functioning and with the vasculature. Of note is the increased expression of various glutamate receptors as well as transmembrane GABA transporters. Finally, the finding of altered expression of a number of genes associated with mitochondrial functioning adds to recent insights in mitochondrial dysfunction in VWM pathology. Altogether, we have created new disease models for VWM using iPSC-derived white and gray matter–like astrocytes, and have demonstrated intrinsic vulnerability of the white matter–like subtypes in VWM. The use of the presented in vitro models may greatly aid further exploration of these new pathways possibly involved in VWM pathophysiology, and can be used in compound screening studies for drug development.

We thank J. Broeke for his help with the calcium imaging of the astrocytes and A. Badia for her help with the morphological analysis using Columbus software.

**Author Contributions**

P.S.L., S.D., A.E.J.H., and V.M.H. contributed to conception and design of the study; P.S.L., S.D., A.E.J.H., K.W., G.J., P.C.-S. and L.G. contributed to the acquisition and analysis of data; P.S.L., S.D., A.E.J.H., M.S.v.d.K., and V.M.H. contributed to drafting the text and preparing the figures.

**Potential Conflicts of Interest**

Nothing to report.

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