Proteomic and Metabolomic Analyses of Vanishing White Matter Mouse Astrocytes Reveal Deregulation of ER Functions

Lisanne E. Wisse‡, Renske Penning‡, Esther A. Zaal‡, Carola G. M. van Berkel‡, Timo J. ter Braak‡, Emiel Polder‡, Justin W. Kenney†, Christopher G. Proud†, Celia R. Berkers‡, Maarten A. F. Altelaar‡, Dave Speijer‡, Marjo S. van der Knaap‡ and Truus E. M. Abbink*

1 Pediatrics, VU University Medical Center, Amsterdam, Netherlands, 2 Biomolecular Mass Spectrometry and Proteomics Group, Utrecht Institute for Pharmaceutical Sciences, Bijvoet Center for Biomolecular Research, Utrecht University, Utrecht, Netherlands, 3 Centre for Biological Sciences, University of Southampton, Southampton, United Kingdom, 4 Medical Biochemistry, Academic Medical Center, Amsterdam, Netherlands

Vanishing white matter (VWM) is a leukodystrophy with predominantly early-childhood onset. Affected children display various neurological signs, including ataxia and spasticity, and die early. VWM patients have bi-allelic mutations in any of the five genes encoding the subunits of the eukaryotic translation factor 2B (eIF2B). eIF2B regulates protein synthesis rates under basal and cellular stress conditions. The underlying molecular mechanism of how mutations in eIF2B result in VWM is unknown.

Previous studies suggest that brain white matter astrocytes are primarily affected in VWM. We hypothesized that the translation rate of certain astrocytic mRNAs is affected by the mutations, resulting in astrocytic dysfunction. Here we subjected primary astrocyte cultures of wild type (wt) and VWM (2b5ho) mice to pulsed labeling proteomics based on stable isotope labeling with amino acids in cell culture (SILAC) with an L-azidohomoalanine (AHA) pulse to select newly synthesized proteins. AHA was incorporated into newly synthesized proteins in wt and 2b5ho astrocytes with similar efficiency, without affecting cell viability. We quantified proteins synthesized in astrocytes of wt and 2b5ho mice. This proteomic profiling identified a total of 80 proteins that were regulated by the eIF2B mutation. We confirmed increased expression of PROS1 in 2b5ho astrocytes and brain. A DAVID enrichment analysis showed that approximately 50% of the eIF2B-regulated proteins used the secretory pathway. A small-scale metabolic screen further highlighted a significant change in the metabolite 6-phospho-gluconate, indicative of an altered flux through the pentose phosphate pathway (PPP). Some of the proteins migrating through the secretory pathway undergo oxidative folding reactions in the endoplasmic reticulum (ER), which produces reactive oxygen species (ROS). The PPP produces NADPH to remove ROS. The proteomic and metabolomics data together suggest a deregulation of ER function in 2b5ho mouse astrocytes.

Keywords: vanishing white matter, eIF2B, AHA, SILAC, astrocytes, metabolomics, endoplasmic reticulum
INTRODUCTION

Vanishing white matter (VWM) is one of the more prevalent leukodystrophies (van der Knaap et al., 1999). Patients with VWM display chronic neurological deterioration and additionally episodes of stress-provoked rapid and severe deterioration. Neuropathology of post-mortem brain shows cystic degeneration of the cerebral white matter with lack of appropriate astrogliotic scar formation, profound lack of myelin, increased numbers of oligodendrocyte precursor cells and immature astrocytes. The morphology of especially astrocytes in cerebral white matter is abnormal. They look coarse and have fewer, thicker processes than normal (van der Knaap et al., 2006; Bugiani et al., 2011). A small proportion of the oligodendrocytes look foamy (Wong et al., 2000).

Recessive mutations in the eukaryotic translation factor 2B (eIF2B) cause VWM (Leegwater et al., 2001; van der Knaap et al., 2002). eIF2B is an enzyme composed of five different subunits (eIF2Bα, β, γ, δ, ε), encoded by five genes (EIF2B1-5). Mutations reduce the activity of eIF2B (van Kollenburg et al., 2006; Horzinski et al., 2009; Liu et al., 2011).

The eIF2B complex functions as a guanine nucleotide exchange factor (GEF), mediating the exchange of GDP for GTP on eIF2. eIF2-GTP binds to methionine-charged initiator tRNA (Met-tRNAMet), thereby forming the ternary complex eIF2-GTP.Met-tRNAMet. The ternary complex together with the small ribosomal subunit binds the 5′-end of the mRNA and scans the 5′-untranslated region (5′-UTR) until it encounters a start codon in a suitable context, whereupon translation of the open reading frame (ORF) starts. Simultaneously, GTP on eIF2 is hydrolyzed to GDP which makes the complex inactive (Konieczny and Safer, 1983; Kleijn et al., 1998; Proud, 2001). GEF activity is needed to recharge eIF2 with GTP.

Translation initiation is a complex process, involving multiple eukaryotic initiation factors (eIFs) (Voorma et al., 1994; Sonenberg and Hinnebusch, 2009). Translation initiation efficiency is profoundly influenced by the nucleotides flanking the start codon (usually AUG), the Kozak consensus sequence (Kozak, 1987). Purines at the −3 and +4 position relative to the AUG start codon are most important in determining translation initiation efficiency (Kozak, 1987; Nederer et al., 2014). The architecture of the 5′-UTR with regard to various sequences, structural motifs and length also determines the translation efficiency of an mRNA (Hinnebusch et al., 2016). eIF2B mutations are expected to reduce ternary complex levels and thus overall mRNA translation. However, this reduced activity can actually enhance translation of certain mRNAs with 5′-upstream open reading frames (uORFs) in their 5′-UTR. uORFs can inhibit translation of the main ORF (mORF) and translation of these mORFs depends on translation reinitiation, which is regulated by ternary complex levels (Meijer and Thomas, 2002; Hinnebusch et al., 2016).

A VWM mouse model, homozygous for the Arg191His mutation in the eIF2Be subunit has been developed, representative of the human disease (2b5ho mice) (Dooves et al., 2016). The Arg191His mutation corresponds to the Arg195His mutation in patients. This mutation reduces eIF2B activity in vitro and gives a severe VWM phenotype (Fogli et al., 2004; Li et al., 2004). Astrocytes in brains of 2b5ho mice are positive for the immaturity marker nestin. In vitro experiments show that 2b5ho astrocytes inhibit maturation of wild type (wt) oligodendrocytes (Dooves et al., 2016). The mechanism by which mutations in eIF2B lead to astrocytic dysfunction and disease remains unclear. Here we aim to improve understanding of the molecular mechanism underlying VWM. We expected that mutant eIF2B would not have a general effect on mRNA translation but rather affects translation of a small number of specific mRNAs, leading to disruption of cellular balances and dysfunction of astrocytes in particular. To identify these translational differences, we subjected adult mouse astrocytes to high-resolution quantitative proteomics after a pulse labeling with AHA (L-azidohomoalanine) combined with SILAC (stable isotope-labeling by amino acids in cell culture) (Dieterich et al., 2006; Eichelbaum et al., 2012; Kenney et al., 2016).

MATERIALS AND METHODS

Mice

All experiments were carried out under the Dutch and European law with approval of the local Institutional Animal Care and Use Committee (IACUC) of the VU University (Amsterdam, Netherlands). Wt and 2b5ho animals were used. All animals were weaned at P21 and had ad libitum access to food and water. The mice were housed with a 12 h light and dark cycle.

Astrocyte Culture

Four-month-old mice were sacrificed by cervical dislocation. Brains were taken out and the olfactory bulbs, cerebella and forebrains were removed. Astrocytes from the remaining structures (gray and white matter structures, including striatum, hippocampi and basal nuclei) were isolated. Brain tissue was minced with a scalpel in Hank’s balanced salt solution (HBSS) without magnesium and calcium (Gibco) at 4°C. The tissue was dissociated with a papain solution containing 20 mM PIPES (pH 7.4), 120 mM NaCl, 5 mM KCl, 1.1 mM EDTA, 5.5 mM L-cysteine-HCl, 40 U/ml DNase and 20 U/ml papain for 30 min at 37°C. Cells were plated in poly-L-ornithine (PLO)-coated flasks and cultured in DMEM/F12 with 15% fetal bovine serum (FBS) (Hyclone), 1% sodium pyruvate (Gibco), 100U penicillin, 100 µg/ml streptomycin (Gibco) and 10 µg/ml gentamicin (Gibco). The cells were passaged twice before they were used for experiments. All chemicals were purchased from Sigma–Aldrich unless otherwise stated. Every experiment was replicated in independent cultures derived from different mice (number of experiments is indicated in figure legends as, e.g., n = 3).

Rate of Protein Synthesis Assay

Astrocytes were plated in 6 cm dishes (~250,000 cells/dish) and cultured until 80–90% confluent. Cells were starved for methionine for 15 min in AHA medium [DMEM/F12 without methionine, lysine, arginine and phenol red (Thermo Fisher Scientific)] supplemented with 91.3 mg/L L-lysine, 147.5 mg/L...
The amount of AHA-staining was corrected for subjection to Western blot analysis. Samples were loaded on 12% lysate were coupled to biotin. Proteins were precipitated by a final concentration of 2 mM. Cells were harvested in lysis buffer (Bachem) was added to AHA medium for the indicated time in a culture medium. The plate was shaken for 2 min and left standing for 10 min before measuring luminescence with a Victor2 plate reader (PerkinElmer Life Sciences).

Cell Viability Assay
Astrocytes were plated in 1/2 area 96 well plates (~5000 cells/well) and cultured for 2 days in AHA-SILAC medium (Thermo Fisher Scientific, custom-made DMEM/F12 without methionine, arginine, l-lysine and phenol red), with addition of 15% dialyzed serum, 0.005% phenol red, SILAC amino acids (94.2 mg/L [13C6]-l-lysine 152 mg/L [13C6]-l-arginine or 96 mg/L [13C6],[15N2]-l-lysine and 154.3 mg/L [13C6],[15N4]-l-arginine) and 17.2 mg/L L-methionine. The cells were starved for methionine from 15 min and cultured with 2 mM AHA (Bachem) or methionine overnight in half well area plates. Cell viability (ATP levels, CellTiter-Glo) was measured according to manufacturer’s instructions (Promega). In short, cells were kept at room temperature for 30 min. CellTiter-Glo was added in the same volume as the culture medium. The plate was shaken for 2 min and left standing for 10 min before measuring luminescence with a Victor2 plate reader (PerkinElmer Life Sciences).

AHA Enrichment and On-Bead Digestion
Astrocytes were plated in 10 cm dishes (~750,000 cells/dish) and cultured until 80% confluent. The culture medium was replaced with AHA-SILAC medium. Cells were grown for 4 days. SILAC labels were reversed in biological duplicates. Cells were subsequently starved for methionine for 15 min and AHA was added as described in the previous section. Astrocytes were further cultured for 2 h or otherwise indicated. The cells were subsequently washed with cold PBS and lysed in urea lysis buffer (supplied with the Click-IT Protein enrichment Kit, Invitrogen).

For the protein analysis of secreted proteins conditioned medium samples from wt and 2b5h0 astrocytes (2 h) were concentrated using Amicon Ultra-15 centrifuge filter tubes (3 kDa, Merck). Samples were diluted with urea lysis buffer to a volume of 400 µl per sample.

The AHA-labeled proteins from cell lysates as well as from the conditioned medium were enriched using the Click-IT® Protein Enrichment Kit according to the manufacturer’s protocol (Invitrogen) with some minor modifications. In short, the AHA-labeled proteins were bound to the resin (16 h), following an iodoacetamide treatment and several washing steps. The AHA-labeled proteins bound to the resin were dissolved in 50 mM ammonium bicarbonate with 3 M urea. Digestion was performed at 37°C by adding 0.1 µg Lys-C for 4 h followed by addition of 1 µg trypsin overnight. The peptides were separated from the resin by briefly centrifuging them through a 0.8 ml spin columns (Thermo Fisher Scientific). The flow through contains the peptides. Peptide samples were stored at −80°C until further use. See also Figure 1 for an overview of the enrichment procedure.

LC-MS/MS Analysis
Peptides were analyzed using a Q ExactiveTM Hybrid Quadrupole-OrbitrapTM Mass Spectrometer (Thermo Fisher Scientific), which was connected to a 1290 Infinity II LC System (Agilent). The trap column was made of C18 (Dr. Maisch ReproSil) material and the analytical column was a 50 cm, 50 µm inner diameter Poroshell C18 (Agilent) column. Both the trap and analytical columns were packed in-house. Solvent A consisted of 0.1% formic acid (Merck) in deionized water (Merck) and Solvent B of 0.1% formic acid in 80% acetonitrile (Biosolve). Peptides were first trapped at 50 µl/min with solvent A and then eluted with solvent B in a 120 min gradient at 100 nl/min: 0–10 min, 100% solvent A; 10.1–10 min, 13–40% solvent B; 105–108 min, 40–100% solvent B; 108–109 min, 100% solvent B; 109–110 min, 0–100% solvent A; 110–120 min, 100% solvent A. The Orbitrap was operated in a data-dependent manner, with the following settings: ESI voltage, 1700 V; inlet capillary temperature 320°C; full-scan automatic gain control (AGC) target, 3 × 106 ions at 35000 resolution; scan range, 350–1500 m/z; Orbitrap full-scan maximum injection time, 250 ms; MS2 scan AGC target, 5 × 104 ions at 17500 resolution; maximum injection, 120 ms; normalized collision energy, 25; dynamic exclusion time, 30; isolation window 1.5 m/z; 10 MS2 scans per full scan.

Data Processing
The LC-MS/MS data were processed with MaxQuant (v1.5.2.8) and MS2 spectra were searched with the Andromeda search engine against the mouse proteome in UniProt (17155 entries, downloaded on 2015-10-27). Enzyme specificity was set to Trypsin/P and two missed cleavages were allowed. SILAC labels (K6/8, R6/10), methionine for AHA substitution and methionine oxidation were set as variable modifications and cysteine carbamidomethylation was set as fixed modification. Minimum peptide length was 7 amino acids. Mass tolerance was set to 20 ppm for peptide masses and 0.6 Da for fragmentation masses. The false discovery rate (FDR) threshold was set to 1% for identifications. Minimal ratio count was set to 2 for protein quantification and the functions “match between runs” and “requantify” were enabled. Data were further analyzed in Perseus (v1.5.0.0). Protein groups were kept for further analysis if they were detected in at least three of the biological replicates.
A t-test was performed and protein groups were considered significant if the p-value was <0.05.

**Signal Peptide Analysis**

To assess the presence of an N-terminal signal peptide in the proteins, we subjected the proteins from the proteomic screen to the SignalP 4.1 server (Bendtsen et al., 2004; Petersen et al., 2011). For all UniProt IDs the FASTA files were downloaded from the UniProt website and all Fasta files are automatically subjected to the SignalP 4.1 server. The presence or absence of a signal peptide for each UniProt ID was predicted with the SignalP 4.1 algorithms and the results were saved to Excel. The UniProt IDs linked back to the list of all identified protein names. Biomart was used to identify proteins containing a transmembrane domain. The number of proteins with a signal peptide was calculated for the proteins, the synthesis of which was affected by eIF2Be\textsubscript{Arg191His} as well as for the other proteins that are not regulated by the same mutation. Next the number of regulated proteins with a signal peptide were subdivided in ‘upregulated’ and ‘downregulated’.

**In Silico Analysis of mRNA Features of eIF2Be\textsubscript{Arg191His}-Regulated Proteins**

The sequences of the mRNA variants encoding the proteins identified in the AHA-SILAC proteomes were downloaded from the NCBI database including the start and stop codon location. The 5′-UTR features “uORFs, %GC and thermodynamic stability (ΔG in kcal/mol)” were determined for all mRNAs (Babendure et al., 2006). We compared each mRNA characteristic for all proteins found in the AHA-SILAC proteomic screen and compared the characteristics for the proteins regulated by eIF2Be\textsubscript{Arg191His} and non-regulated proteins. The 5′-UTR length,
Chilled microfuge tubes. Protein concentrations were determined through a 23G needle. The samples were centrifuged (10 min, pH 7.4, 100 mM KAc, 3 mM MgAc$_2$, 4°C) and the supernatant was used for Western blot analyses.

**DAVID Analysis**

Overrepresentation of specific pathways was analyzed with DAVID [based on gene ontology (GO)-terms] between the proteins that are affected by the eIF2Be$^{Arg191His}$ mutation and all the proteins found in the AHA-SILAC proteome (Huang et al., 2009a,b).

**Cell Lysates for Western Blot Experiments**

Astrocytes were plated in 10 cm dishes (~750,000 cells/dish) and cultured until 80% confluent. Cells were washed with cold PBS, collected by scraping in PBS and pelleted by centrifugation (5 min, 1000 × g, 4°C). The method is based on the protocol described. Cells were lysed in harvesting buffer containing 10 mM HEPES pH 7.9, 50 mM NaCl, 0.5 M sucrose, 0.1 mM EDTA, 0.5% Triton, 1 mM DTT, phosphatase inhibitors (0.5 mM activated NaVO$_3$), 25 mM β-glycerophosphate, 50 mM NaF) and protease inhibitors (Roche). Lysates were centrifuged (10 min 500 × g, 4°C) and the supernatant was used for Western blot analyses.

**Whole Brain Lysates for qPCR and Western Blot Experiments**

Mice were sacrificed by cervical dislocation at 4 months of age. Brains were removed, snap-frozen in liquid nitrogen and stored at −80°C until further use. Lysates were prepared by grinding the brain samples with a pestle and mortar under liquid nitrogen. The powder was lysed in cytoplasmic lysis buffer (20 mM Tris pH 7.4, 100 mM KAc, 3 mM MgAc$_2$, 2 mM DTT, 15% IGEPAL, 1.5% sodium deoxycholate (SODC), 1x HALT (protease and phosphatase inhibitory cocktail, Thermo Fisher Scientific). The samples were homogenized with a pestle followed by trituration through a 23G needle. The samples were centrifuged (10 min, 10,000 × g, 4°C) and the supernatant was aliquoted into pre-chilled microfuge tubes. Protein concentrations were determined using a Quick Start™ Bradford Protein Assay (Bio-Rad). For RNA isolation, TRIzol™ Reagent (Invitrogen) was added to 50 µl of the supernatant and total RNA was isolated as described under RNA isolation and cDNA synthesis.

**Western Blot**

For cell lysates approximately 10 µg and for brain lysates 50–60 µg of protein was loaded on a 12% SDS-polyacrylamide gel with 2,2,2-trichloroethanol (TCE) which allows detection of total protein load (Ladner et al., 2004). Proteins were transferred onto a PVDF membrane (Bio-Rad). Membranes were blocked in 5% (w/v) milk powder and stained with the antibody against PROS1 (16910-1-AP, Proteintech) or SLC3A2 (LS-C334231, LSBio) overnight (16 h) at 4°C. Membranes were washed with TBS-Tween20 (0.1%) and incubated with an HRP-conjugated secondary goat anti-rabbit IgG antibody (Dako, P0448) for 2 h at room temperature. The membranes were washed three times with TBS-Tween (0.1%) and once with TBS, incubated with SuperSignal™ West Femto (Thermo Fisher Scientific) and imaged (Odyssey® Fc, LI-COR). Protein expression was corrected for total amount of protein determined by Gel Doc™ EZ System (Bio-Rad).

**RNA Isolation, cDNA Synthesis and qPCR**

Astrocytes were washed twice with PBS and collected in TRIzol™ Reagent (Invitrogen). RNA was isolated according to the manufacturer’s protocol. In short, 1/5 volume chloroform was added to the TRIzol™ Reagent. The samples were centrifuged for 20 min at 4°C and 12000 × g. The pellet was washed twice with 70% ethanol and resuspended in non-DEPC treated water (Ambion). RNA was precipitated with 175 mM sodium acetate (pH 5.2) and 70% ethanol and incubated for 30 min at −20°C. The samples were centrifuged (30 min, 4°C, 12000 × g) and the pellets were washed twice with 70% ethanol and resuspended in non-DEPC-treated water.

RNA quality and quantity were determined by measuring the A$_{260}$ and A$_{280}$ (NanoDrop 2000, Thermo Fisher Scientific). cDNA was synthesized in a 20 µl reverse transcription reaction: 1x first strand buffer (Invitrogen), random hexamers (0.02 µg/µl; Qiagen), oligoDT (0.02 µg/µl; Qiagen), dNTPs (1 mM each; Roche), DTT (1 mM; Invitrogen), RNaseOUT (0.25 U/µl; Invitrogen), Superscript III (5 U/µl; Invitrogen) and 1.5 µg total RNA were incubated for 2 h at 50°C. RNaseH (62.5 U/µl; Invitrogen) was added and incubated for 30 min at 37°C followed by 15 min at 70°C.

mRNA levels were determined with qPCR using a LightCycler® 480 II Instrument (Roche). For each 10 µl sample a mixture of LightCycler® 480 SYBR Green I Master (Roche), primers (1 pmol/µl) and cDNA (0.1 µl) was used. Used primers are listed in Table 1. Gapdh mRNA was used as reference.

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1. http://unafold.rna.albany.edu/?q=DINAMelt/Quickfold
2. http://ron.cimr.cam.ac.uk/protocols/NucCyto.html
Constructs

Construct to Determine Secretion

The pNL1.3 plasmid (Promega) expresses the nanoluciferase protein with an N-terminal signal peptide from interleukin-6 (IL-6; Secluc). The signal peptide promotes secretion of the nanoluciferase into the culture medium. The pNL1.3-6 (IL-6; Secluc). The signal peptide promotes secretion of the nanoluciferase into the culture medium. The pNL1.3 Gapdh construct contains the promoter including 5'-UTR of Gapdh, which was inserted in the pNL1.3 vector using an infusion reaction, according to the manufacturer’s protocol (In-Fusion HD Cloning Kit, Clontech). The PCR-amplified sequence of the constructs was confirmed by sequence analysis.

Constructs to Determine Translation

The pNL1.1 plasmids express the nanoluciferase protein (Nluc). Promoter and 5'-UTR sequences of candidate genes were taken from NCBI and Ensembl databases (Table 2). Infusion primers were designed to amplify promoter (approximately 2000 bps upstream of the transcription start site) and 5'-UTR-encoding sequences using the primer design tool on the Clontech website. Internal primers were designed to merge the Gapdh promoter region and the 5'-UTR of the candidate. The promoter and 5'-UTR sequences were amplified using the infusion primers and the overlapping internal primers on gDNA from mouse liver or cDNA from cultured mouse astrocytes (Table 2) according to the manufacturer’s protocol (ClonAmp, In-Fusion HD, Clontech). For the chimeric constructs comprising the murine Gapdh core promoter and 5'-UTR of the candidate mRNA a triple infusion reaction was performed with the pNL1.1 vector, the amplified promoter product and the amplified 5'-UTR product (Table 2). All promoter and 5'-UTR sequences were inserted into the pNL1.1 vector (Promega) using an infusion reaction according to the manufacturer’s protocol (In-Fusion HD Cloning Kit, Clontech). The resulting plasmids are listed in Table 2. PCR-amplified sequence of all constructs was confirmed by sequence analysis.

Construct Used As Internal Control

The pGL3 plasmid expresses the firefly luciferase protein (Fluc). The murine Gapdh promoter and 5'-UTR were digested from the pNL1.1 vector with KpnI and Ncol, purified from agarose gel (High Pure PCR Cleanup, Roche) and inserted into an empty pGL3 vector digested with the same restriction enzyme using a T4 DNA ligase (Promega). The resulting plasmid pGL3-Gapdh was used as internal standard in transfection studies. PCR-amplified sequence of the constructs was confirmed by sequence analysis.

Measurement of Secretion and Secretory Pathway Flux

Astrocytes were plated in half-area 96 well plates (~3000 cells/dish, CELLSTAR) to 80% confluency. pNL1.3-Gapdh (80 ng) was transfected into the cells using 0.24 µl FuGENE 6 according to the manufacturer’s protocol (Promega). Luciferase activity in cells and culture medium was measured using the protocol of Promega and a Wallac 1420 Victor2 Microplate Reader (Perkin Elmer). The ratio of extracellular/intracellular luciferase activity determines the relative secretion.

Transfections

Astrocytes were plated in half-area 96 well plates (~3000 cells/dish, CELLSTAR). Each pNL1.1 promoter/5'-UTR construct (40 ng) was co-transfected with internal control pGL3-Gapdh (20 ng) using FuGENE 6 Transfection Reagent (0.24 µl) according to the manufacturer’s protocol. Luciferase activity was measured using the protocol of Promega and a GloMax® Discover System (Promega). The ratio Nluc:Fluc determines the expression of Nluc corrected for well-by-well differences.

Metabolomics

Astrocytes were cultured in 6 cm dishes (~250,000 cells/dish) until 70% confluent. Medium was replaced 72 and 24 h before harvesting. At the time of harvesting culture medium was removed from the cells and stored at −80°C until analysis. Cells were washed with ice-cold PBS and metabolites were extracted from cells in 0.5 ml lysis buffer containing methanol/acetonitrile/H2O (2:2:1). Samples were spun at 16,000 × g for 15 min at 4°C. Supernatants were collected for LC-MS analysis. 10 µl of conditioned medium was added to 1 mL of lysis buffer containing methanol/acetonitrile/H2O (2:2:1) and prepared as above.

LC-MS analysis was performed on an Exacta mass spectrometer (Thermo Fisher Scientific) coupled to a Dionex Ultimate 3000 autosampler and pump (Thermo Fisher Scientific). The MS operated in polarity-switching mode with spray voltages of 4.5 and −3.5 kV. Metabolites were separated using a SeQuant® ZIC®-pHILIC HPLC Columns (2.1 mm × 150 mm, 5 µm, guard column 2.1 mm × 20 mm, 5 µm; Merck) using a linear gradient of acetonitrile and eluent A [20 mM (NH4)2CO3, 0.1% NH4OH in ULC/MS grade water (Biosolve)]. Flow rate was set at 150 µl/min. Metabolites were identified and quantified using LCQUAN™ Quantitative Software (Thermo Fisher Scientific) on the basis of exact mass within 5 ppm and further validated by concordance with retention times of standards. Metabolites were quantified.

### Table 1 | Primers used for qPCR.

| Gene Name | Forward (5'-3') | Reverse (5'-3') |
|-----------|----------------|----------------|
| Gapdh     | GTGCTGAGTATGTCGTGGAAG | TCGTGCTCTACCCCATCAAC |
| Akt       | AAAGGACGCGGATCGTCGCGG | GGTGCGTGGGCTGAATGGAAG |
| Gas6      | CTAAAACGTCCCAAGACAT | GTGACAGGCTCTACCGCTT |
| Nrxn1     | CCCCACAAGAAGAAGCCCATCA | GTTGACCTAACCCACGGACG |
| Sarcs1    | GCGACATCGCCGAGCTCATC | AACCCGCAAGCAAGATG |
| Fndc3c1   | AAGGCGAGCTTTTGGAGAAGA | TGCCACTGTTGAGGATTTCA |
| Pros1     | TTCGTCAGGCTACTCACCTTGT | AGGCTCAAGGAAGGCTCAC |
| Scrn1     | GTCTGCTGCTGCTCAGAAAC | TTTGCGTGGCTCCTGCTCA |
| Fh1       | GGGAGAACAACTTCTCGATGT | TCGGATAGGTTACCGCCA |
| Tnixp     | GTCTCAAGCGAGCTCAGAAACGAC | CTTGCCAGGATAGCTACGG |
| Daf1      | CTGCTGCTCTCCTCATCATC | GTGACGCTCTCCCTCCAAAC |
| Slc3a2    | GAAAGCGTCATGACGGACCC | CAATTTGCGAGGTCGAGG |
| Sec31A    | GAGAGCTCGTCGTCGTGGAAG | AGAACCTCAATACCCCATC |
using LCQUAN\textsuperscript{TM} Quantitative Software (Thermo Fisher Scientific). Peak intensities were normalized based on median peak intensity.

**Statistical Analyses**

The program Factor was used to correct for differences between experiments (qPCR, Western blot, cell viability) but not between other conditions (genotype, treatments) (Ruijter et al., 2006). Statistical analysis of qPCR, Western blot and cell viability experiments were performed with a two-way ANOVA followed by a Sidak’s multiple comparison test, using GraphPad Prism software. Statistical analysis of the transfection data was performed using a T-test per construct.

A Chi-square analysis was used to measure significant differences in the presence of signal peptides, transmembrane regions and SP-targeting proteins as well as for the analysis on the 5’-UTR features using GraphPad Prism software.

The secretion assay was statistically tested with a multilevel analysis in SPSS. Differences were significant when \( p < 0.05 \).

| Gene number | Construct name | Primer Sequence |
|-------------|----------------|-----------------|
| NM_008084   | pNL1.3-Gapdh   | Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_008084   | pNL1.3-Gapdh   | Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_011173   | pNL1.1-Pros1    | Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_015921   | pNL1.1-Gas6     | Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_027268   | pNL1.1-Scrn1    | Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_010206.3 | pNL1.1-Fgfr1    | Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_011173   | pNL1.1-Pros1    | 5’UTR Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_015921   | pNL1.1-Gas6     | 5’UTR Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_010206.3 | pNL1.1-Fgfr1    | 5’UTR Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_020252.3 | pNL1.1-Nrxn1    | 5’UTR Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_021377   | pNL1.1-Sorcs1   | 5’UTR Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_001198833| pNL1.1-Ddr1     | 5’UTR Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |
| NM_011173   | pNL1.1-Pros1    | promoter Infusion FW primer TGGCCTAAGTCGGCGGTAACCTGTGTGTACTACCGAAGAAACAAGAGGAGAGGAGAT | Infusion RV primer GGCTAGCGAGCTCACTCGTCTGTTGCTGAGGCTGGACGCCACAGAAAG |

TABLE 2 | Primer sequences used for cloning.
RESULTS

AHA-SILAC Incorporation Is Not Affected by Mutations in eIF2B and Does Not Affect Astrocyte Viability

L-azidohomoalanine incorporation was measured in wt and 2b5ho astrocytes to investigate if the general protein synthesis rate is affected by the homozygous Arg191His mutation in eIF2Bε (Figure 1). Label incorporation increased linearly for a period of 4 h (Figure 1B). The occurrence occurred with similar kinetics in 2b5ho and wt astrocytes (Figure 1B), indicating that the eIF2Bε Arg191His mutation did not significantly affect protein synthesis rate. The viability of both wt and 2b5ho astrocytes was not influenced by AHA or SILAC treatment (Figure 1C). Expression of stress-related mRNAs was unaffected by the AHA-labeling protocol (Supplementary Figure 1). These results allow further proteomic analyses with an AHA incorporation pulse of 2 h.

Pulsed Labeling Proteomics of Astrocytes Reveals 80 Proteins Regulated by the eIF2Bε Arg191His Mutation

The AHA proteomic labeling approach was performed to identify and quantify proteins that are differentially translated in 2b5ho astrocytes. SILAC-labeled wt and 2b5ho astrocytes were subjected to AHA labeling for 2 h. After AHA labeling cells were harvested and subjected to bead-based enrichment using a Click-iT chemistry approach. Bound proteins were digested and resulting peptides were analyzed by LC-MS/MS. In four biological replicates, we identified a total of 2888 proteins across both wt and 2b5ho astrocytes. 1240 proteins were detected in all proteins, proteins measured in wt and 2b5ho astrocytes; regulated, proteins that differ significantly in intracellular accumulation between wt and 2b5ho astrocytes (divided in upregulated and downregulated).

TABLE 3 | Analysis of the proteins that are synthesized and migrate in the secretory pathway (SP).

| AHA-SILAC data set* | % of proteins with a signal peptide (absolute #) | % of proteins with a transmembrane domain (absolute #) | % of SP-targeting proteins (absolute #) |
|---------------------|-----------------------------------------------|---------------------------------------------------|---------------------------------------|
| All proteins (1240) | 19.1% (237)                                  | 14.3% (177)                                      | 25.6% (317)                           |
| Regulated (80)      | 33.8% (27)                                   | 30.0% (24)                                      | 47.5% (38)                           |
| Upregulated (72)    | 37.5% (27)                                   | 33.3% (24)                                      | 52.7% (38)                           |
| Downregulated (8)   | 0% (0)                                       | 0% (0)                                           | 0% (0)                               |

*All proteins, proteins measured in wt and 2b5ho astrocytes; regulated, proteins that differ significantly in intracellular accumulation between wt and 2b5ho astrocytes (divided in upregulated and downregulated).
proteins (Figure 2B). This outcome suggests that many regulated proteins follow the secretory pathway. Since signal peptides and transmembrane domains are associated with targeting to the secretory pathway (von Heijne, 1990; Petersen et al., 2011; Borgese, 2016), we tested all 1240 identified proteins for signal peptides or transmembrane domains. Signal peptides were found in 237 proteins and transmembrane domains in 177. The number of proteins predicted to pass through the secretory pathway was significantly enriched in the 2b5ho proteome (Table 3). 27 out of the 80 regulated proteins harbor a signal peptide and 24 have at least one transmembrane domain; 13 of the proteins harbor both a signal peptide and a transmembrane domain (in total 38 proteins out of 80). Intriguingly, the proteins annotated as migrating through the secretory pathway were all upregulated in 2b5ho astrocytes. It is possible that these proteins accumulate more in 2b5ho astrocytes during the 2-h labeling pulse due to a secretory pathway flux difference and not to an actual increase in translation rate. To investigate this, we analyzed the AHA-enriched secretome from wt and 2b5ho cultures after a 2-h AHA labeling. We detected 22 labeled proteins in the secretome, of which three (APOE, CST3 and POSTN) were significantly reduced in the 2b5ho astrocyte-conditioned medium. Intracellularly, APOE and POSTN were not changed by eIF2BεArg191His. CST3 was significantly increased intracellularly in 2b5ho astrocytes. No other protein from the original 27 eIF2BεArg191His-regulated proteins with a signal peptide was detected in the secretomes of wt and 2b5ho astrocytes. To further address the hypothesis that 2b5ho astrocytes have an altered secretory pathway flux, we evaluated the flux of proteins migrating through the secretory pathway with a reporter assay. The assay showed robust secretion of the Secluc reporter (Supplementary Figure 2) and secretion differences between wt and 2b5ho astrocytes were not detected (Figure 3).

Validation of eIF2BεArg191His-Regulated Proteins

To investigate if the proteins that were found in the screen are indeed differentially expressed at the translational level, we selected eleven candidates that are either increased or decreased in wt vs. 2b5ho cultures. We quantified their mRNA levels to discriminate whether differences in protein amounts are regulated at the transcriptional or the translational level. Only one of the selected candidates differed in mRNA expression between wt and 2b5ho astrocytes (Figure 4A), suggesting that the increased accumulation of the other 10 proteins was not due to increased transcription or mRNA stability. Of the 11 candidates two were further investigated at the protein level by Western blot. We found that PROS1 and SLC3A2 were indeed significantly increased at the total protein level (Figure 4B). This result confirms the data from the proteomic screen and also demonstrates increased accumulation of these candidate proteins.

We next investigated several targets in mouse brain. At the mRNA level seven out of nine targets were similar between wt and 2b5ho brains (Supplementary Figure 3A). Two proteins were further assessed using Western blot (Supplementary Figure 3B). SLC3A2 protein abundance was also increased in brain lysates; however, this correlated with increased Slc3a2 mRNA levels, suggesting transcriptional regulation in brain. PROS1 protein abundance was also increased in brain lysates, but Pros1 mRNA abundance was not. This finding suggests translational upregulation of PROS1 both in cell culture and brain.

In Silico Analysis of mRNAs
eIF2B plays an essential role in the regulation of protein synthesis. For this reason, we investigated if the mRNAs of the eIF2BεArg191His-regulated proteins share specific features that could explain the translational regulation by eIF2B. We investigated the 5’-UTRs for length, structural stability (%GC and ΔG) and number of uORFs with a Kozak sequence. The 5’-UTR length was significantly increased for mRNAs encoding eIF2BεArg191His-regulated proteins (Figure 5). Of note, the mRNAs for the eight proteins decreased in the eIF2BεArg191His-regulated proteome have relatively short 5’-UTRs (29-343 bases, median 133). This analysis showed that the eIF2BεArg191His mutation differentially influences expression of proteins translated from mRNAs with short 5’-UTRs (<150 bases), which were underrepresented, compared to those with lengthy 5’-UTRs (>550 nucleotides), which were overrepresented; those translated from mRNAs with short 5’-UTRs, if regulated, tend to go down in expression in the mutant cells. The %GC in the 5’-UTRs did not significantly differ between eIF2BεArg191His-regulated proteins and other proteins (Figure 5). The thermodynamic stability (ΔG in kcal/mole) of the 5’-UTR seemed to be higher for mRNAs encoding the eIF2BεArg191His-regulated proteins, although not significantly (Figure 5). The
overall number of uORFs did not significantly differ for mRNA encoding the eIF2BεArg191His-regulated proteins compared to the non-regulated proteins (Figure 5). However, mRNAs with high numbers of uORFs (8 or more) were clearly overrepresented in the group of significantly altered proteins.

**Translation Efficiency Was Assessed in Reporter Assays**

We investigated the potential translational regulation of some eIF2BεArg191His-regulated proteins using reporter constructs encoding nanoluciferase driven by promoter sequences (including the 5′-UTR) of the candidates GAS6, PROS1, SCRN1 and FGFR1. A Gapdh promoter construct was included as non-regulated control. Transfection of each construct yielded similar luciferase expression in wt and 2b5ho cultures (Figure 6). Transfection of the pNL1.1-Fgfr1 promoter construct did not yield a reliable level of nanoluciferase (approximately two–threefold over the background signal in non-transfected cells) and was therefore omitted from analysis. The levels of nanoluciferase activity expressed with the other constructs were still low in comparison to the Gapdh-predictor-driven expression. Because expression could be increased by replacing the candidate promoter with the Gapdh promoter, we therefore constructed Gapdh-candidate chimeras. With these chimeras, we tested the 5′-UTR efficiency of candidates GAS6, SORCS1, PROS1, FGFR1, NRXN1, and DDR1 in wt and 2b5ho cultures.

Neither the pNL1.1-Fgfr1-5′-UTR nor the pNL1.1-Nrxn1-5′-UTR constructs yielded quantifiable nanoluciferase expression. These constructs were omitted from further analyses. The expression from the chimeric promoter-5′-UTR constructs of GAS6 and Pros1 was increased by the Gapdh core promoter in both wt and 2b5ho cells. Still, none of these constructs yielded an increased nanoluciferase expression in 2b5ho compared to wt astrocytes (Figure 6).

**Metabolic Screen of Astrocytes**

**Conditioned Medium and Lysates**

Because the proteomic data only gave small differences, the VWM phenotype remained difficult to explain. Tiny differences in protein levels and post-translational modifications occasionally lead to metabolic shifts. Thus we checked whether they affected cellular metabolism, using a mass spectrometry-based metabolic screen. We measured intra- and extracellular metabolites 24 and 72 h after replacing the culture medium. We looked at general energy consumption by measuring the uptake and secretion of metabolites from the culture medium (Figure 7A). As expected, both cultures had taken up glutamate (Martin-Jimenez et al., 2017) and aspartate (which use the same transporters) (Bender et al., 1997), reflecting specific astrocyte function. The uptake of glutamate and aspartate did not influence their intracellular levels (Supplementary File 2). Also, both wt and 2b5ho cultures showed uptake of pyruvate. Astrocytes convert...
FIGURE 5 | In silico 5′UTR analysis of the eIF2β<sup>Arg191His</sup>-regulated proteins. The 5′UTRs of the mRNAs encoding the proteins regulated by eIF2β<sup>Arg191His</sup> mRNAs are significantly longer than of those encoding non-regulated proteins. They have an overall similar percentage GC content to the non regulated proteins. However, they seem to tend to higher ΔG's. These findings combined are indicative of relatively structured 5′-UTRs in the mRNAs of eIF2β<sup>Arg191His</sup>-regulated proteins. The number of uORFs is similar between the significant and not significant proteins, though mRNAs with very high numbers (8 or more) of predicted uORFs are overrepresented in the group of significantly altered proteins.

dysregulation in VWM astrocytes.

pyruvate into lactate and glutamate/aspartate into glutamine, cis-aconitate and α-ketoglutarate (Martin-Jimenez et al., 2017). Indeed, the wt as well as the 2b<sup>ho</sup> cultures secreted lactate, glutamine, cis-aconitate and α-ketoglutarate (Figure 7A). In addition, we investigated intracellular metabolites (Figure 7B). 6P-glucuronate, an intermediate of the pentose phosphate pathway (PPP), was detected in 2b<sup>ho</sup> but not in wt cells at both time points (Figure 7B). A fluctuation over time in NADPH levels was observed. The remaining metabolites tested did not show consistent differences over time between wt and 2b<sup>ho</sup> astrocytes (Figure 7B and Supplementary File 2).

**DISCUSSION**

Here we aimed to identify whether specific proteins are translationally regulated by eIF2B<sub>ε</sub><sup>Arg191His</sup> in astrocytes. We used primary astrocyte cultures of the 2b<sup>ho</sup> mouse model. Both wt and 2b<sup>ho</sup> cells display an astrocyte morphology, synthesize CD44, nestin and vimentin (Supplementary Figure 5 and File 1), supporting astrocyte identity. We first established that cultures of primary astrocytes are suitable for sensitive labeling of newly synthesized proteins. Consistent with previous studies (Howden et al., 2013; Genheden et al., 2015), we found that AHA-labeling did not affect murine astrocyte viability for at least 16 h. No differences in AHA-labeling efficiency were observed between wt and 2b<sup>ho</sup> astrocytes. These results confirm that the pulsed AHA-SILAC labeling protocol is suitable to study differences in proteins synthesized in wt and 2b<sup>ho</sup> astrocytes, enabling us to investigate whether the Arg191His mutation in eIF2B<sub>ε</sub> affects translation of specific mRNAs.

We identified 80 proteins that accumulate differentially in wt vs. 2b<sup>ho</sup> astrocyte cultures upon a 2 h AHA pulse. The majority (72 out of 80) is upregulated in 2b<sup>ho</sup> astrocytes. We investigated the proteomic results by qPCR (11 candidates) and Western blot (4 candidates) (Figure 4). The tested antibodies detecting PROS1 and SLC3A2 showed consistent differences between wt and 2b<sup>ho</sup> while TXNIP gave variable results and SCRN1 was not detected. Changes observed at the proteomic level were not due to altered mRNA levels, indicating that differences arose at the translational level. PROS1 and SLC3A2 protein levels were also increased in 2b<sup>ho</sup> mouse brain lysates, supporting the findings in astrocyte cultures. Of these, Pros1 was not increased at the mRNA level.
Our analyses indicate significant enrichment for proteins containing a signal peptide and motifs for N-glycosylation and disulfide bond formation in 2b5ho astrocytes (Figure 2B). These motifs highlight involvement of the endoplasmic reticulum (ER), the first organelle of the secretory pathway (Farhan and Thomas, 2002). Our analyses highlighted some ER proteins (Sen et al., 2016). To further characterize the 5'-UTR efficiency in wt and 2b5ho astrocytes, we performed transient transfection assays for several candidates, some of which were longer than 700 nucleotides (Fgfr1 and Nrxn1). Unfortunately, transfection of the Fgfr1 and Nrxn1 constructs did not yield reliable expression, which precluded testing whether they are translated more efficiently in 2b5ho than in wt astrocytes. The transfection assay is not robust enough to detect differences that were picked up with the pulsed AHA-SILAC proteomics and in silico 5'-UTR analyses.

PROS1 was shown to be upregulated in both 2b5ho astrocytes and brain without increased mRNA level. In brain, PROS1 is expressed by astrocytes and microglia (Zhang et al., 2014). Interestingly, PROS1 and GAS6 (which was also found to be upregulated) bind the same class of receptors (Hafizi and Dahlback, 2006). PROS1 and GAS6 enhance myelin formation and support oligodendrocyte survival in mice in vitro and in vivo (Tsiperson et al., 2010; Gruber et al., 2014; Goudarzi et al., 2016; Akkermann et al., 2017). Recently, PROS1 was described to function in neural stem cells (NSCs) as a regulator for NSC quiescence, proliferation and NSC development into neurons or astrocytes (Zelentsova et al., 2016).

Other proteins enriched in the eIF2Be<Arg191His>-regulated proteome are implicated in astrocyte development. For example Nrxn1 (Zeng et al., 2013) and Sorcs1, that transports Nrxn1, can induce differentiation toward astrocytes (Savas et al., 2015). Moreover TGFB2 (Park et al., 2012) and its binding partners, ERBIN and latent TGFB-binding protein 3 (LTBP3) are increased during astrocyte differentiation (Robertson et al., 2015). All these results suggest that 2b5ho astrocytes have a slightly altered differentiation state as shown before in 2b5ho mice and VWM patients (Bugiani et al., 2011; Dooves et al., 2016).
FIGURE 7 | Intracellular and extracellular metabolic analysis after 24 and 72 h of culture showed increased levels of intracellular levels of a pentose phosphate pathway intermediate (6P-gluconate). The astrocytes were cultured until 70% confluent. Media of the cultures were replaced with fresh medium. The cells as well as the media were collected after 24 and 72 h. The graphs show the mean ± SD. (A) The metabolite concentrations in the culture media of wt and 2b5ho astrocytes were compared to unconditioned media (n = 2). Glucose levels and pyruvate levels in conditioned medium of wt and 2b5ho astrocytes were decreased as well as glutamate and aspartate levels. Lactate, glutamine, cis-aconitate and α-ketoglutarate were increased in the conditioned medium of both cultures. (B) 6P-gluconate levels were consistently detected in 2b5ho cultures but not in wt cultures (24 and 72 h after replacing the culture medium). Four other metabolites of the pentose phosphate pathway were detected in the metabolic screen (erythrose-4P and sedoheptulose-7P, NADPH and NADP^+), but these were not consistently altered between wt and 2b5ho cultures.

To investigate other functional changes, we compared the metabolomes of wt and 2b5ho cells. We detected glutamate uptake in both cultures, which is a typical function of astrocytes, (Martin-Jimenez et al., 2017). This observation further confirms the astrocytic identity of the cultured cells. The most consistent difference observed was an increase in 6P-gluconate, an intermediate of the PPP. The PPP shunt is an alternative route for the metabolism of glucose which can supply ribose for nucleotide production in the non-oxidative part and NADPH in the oxidative part of the pathway. The reducing equivalents provided by NADPH are used both for biosynthesis and repair of oxidative damage (Stincone et al., 2015). The oxidative intermediate 6P-gluconate was increased in 2b5ho astrocytes while the non-oxidative intermediates sedoheptulose 7-phosphate and erythrose 4-phosphate were not consistent over time. These results suggest an increased activity of the PPP, especially at earlier time points. Interestingly, the observed PPP signature has been found in lung cancer cells in response to...
FIGURE 8 | Overview of differentially regulated functions in 2b5ho astrocytes. Proteins with a signal peptide or transmembrane domain enter the ER to be folded. Proteins going through the secretory pathway that are subjected to N-glycosylation and disulfide bond formation are enriched in the set of eIF2β Arg191His-regulated proteins. Disulfide bridge formation through oxidation of these proteins can induce H₂O₂. 6-Phospho-gluconate is involved in the production of NADPH which can be used to remove H₂O₂. PDI, protein disulfide isomerase; GSH, reduced glutathione; GSSG, oxidized glutathione; Ero1-ox, oxidized Ero1; Ero1-red, reduced Ero1; PPP, pentose phosphate pathway.

reduced 6P-gluconate dehydrogenase (6PGDH) activity leading to reactive oxygen species (ROS) production (Patra and Hay, 2014).

Could there be a link between the two main observations, i.e., the deregulated transport of some proteins in the ER and the increased 6P-gluconate concentrations? We think this is likely; although PPP is cytosolic, there are links to the ER. Part of the pathway appears to be associated with the ER (Stockwell et al., 2012) and NADPH produced by the PPP might need to combat ROS formation, which can be produced in the ER by processes related to oxidative protein folding. On the basis of the combined proteomic and metabolomic findings, we hypothesize that 2b5ho astrocytes may accumulate ROS as a result of an increased flux of proteins (e.g., requiring correct disulfide bond formation) through the secretory pathway (Figures 2, 8). Correct disulfide bond formation is dependent on protein disulfide isomerases (PDIs) and Ero1 in the ER (Bechtel and Weerapana, 2017). This process can result in H₂O₂ as by-product and increased PPP flux might compensate by producing NADPH (Espinosa-Diez et al., 2015). 6P-gluconate is the only metabolite in the metabolic screen that was changed in 2b5ho astrocytes and it is not completely clear whether this reflects increased production or decreased breakdown or a combination of both. There are indications that G6PDH is the only NADPH-producing enzyme activated upon oxidative stress and therefore it has been called “guardian of the cell redox potential” (Filosa et al., 2003). Lack of consistent changes over time in NADPH in 2b5ho astrocytes suggest that the balance between H₂O₂ and NADPH in steady state is “successfully” compensated (Figure 8). On the basis of our results we cannot discriminate between a compensated system or a deregulated system, the latter leading to pathology (Bechtel and Weerapana, 2017).

CONCLUSION

The 80 proteins differentially detected in the AHA-SILAC proteome are most likely translationally regulated as their mRNA levels are similar. Regulation thus must occur via increased synthesis or accumulation in 2b5ho astrocytes. We observed that the function of some of these differentially affected proteins is linked to astrocyte differentiation, which is known to be disturbed in VWM. As we would expect from deregulated protein synthesis in 2b5ho cells, the regulated proteins differ from their non-regulated counterparts in aspects of their 5’-UTRs. Altered synthesis may lead to an increase of proteins in the ER and secretory pathway without an overt effect on overall secretion. Possibly, increased demands of protein folding in the ER may result in generation of H₂O₂ or other ROS by-products (Espinosa-Diez et al., 2015). Thus, the increased synthesis of proteins that undergo disulfide formation and elevated 6P-gluconate might point to increased ROS in astrocytes, which may affect their function (Figure 8).
AUTHOR CONTRIBUTIONS

Substantial contributions to the conception or design of the work: LW, RP, CP, MvdK, and TA. The acquisition, analysis: LW, RP, EZ, Cyb, TtB, and EP. Interpretation of data for the work: LW, RP, EZ, JK, CP, CB, MA, DS, and TA. Drafting the work or revising it critically for important intellectual content: LW, RP, EZ, Cyb, TtB, EP, JK, CP, CB, MA, DS, MvdK, and TA. Final approval of the version to be published: LW, RP, EZ, Cyb, TtB, EP, JK, CP, CB, MA, DS, MvdK, and TA. Agreement to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved: LW, RP, EZ, Cyb, TtB, EP, JK, CP, CB, MA, DS, MvdK, and TA.

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SUPPLEMENTARY MATERIAL

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