Regular Article

Voltage-gated Ca$$^{++}$$ entry promotes oligodendrocyte progenitor cell maturation and myelination in vitro

V.T. Cheli, D.A. Santiago González, V. Spreuer, P.M. Paez *

Hunter James Kelly Research Institute, Department of Pharmacology and Toxicology, School of Medicine and Biomedical Sciences, SUNY, University at Buffalo, NYS Center of Excellence, 701 Ellicott St., Buffalo, New York 14203, USA

A B S T R A C T

We have previously shown that the expression of voltage-operated Ca$$^{++}$$ channels (VOCCs) is highly regulated in the oligodendroglial lineage and is essential for proper oligodendrocyte progenitor cell (OPC) migration. Here we assessed the role of VOCCs, in particular the L-type, in oligodendrocyte maturation. We used pharmacological treatments to activate or block voltage-gated Ca$$^{++}$$ uptake and siRNAs to specifically knock down the L-type VOCC in primary cultures of mouse OPCs. Activation of VOCCs by plasma membrane depolarization increased OPC morphological differentiation as well as the expression of mature oligodendrocyte markers. On the contrary, inhibition of L-type Ca$$^{++}$$ channels significantly delayed OPC development. OPCs transfected with siRNAs for the Cav1.2 subunit that conducts L-type Ca$$^{++}$$ currents showed reduced Ca$$^{++}$$ influx by ~75% after plasma membrane depolarization, indicating that Cav1.2 is heavily involved in mediating voltage-operated Ca$$^{++}$$ entry in OPCs. Cav1.2 knockdown induced a decrease in the proportion of oligodendrocytes that expressed myelin proteins, and an increase in cells that retained immature oligodendrocyte markers. Moreover, OPC proliferation, but not cell viability, was negatively affected after L-type Ca$$^{++}$$ channel knockdown. Additionally, we have tested the ability of L-type VOCCs to facilitate axon–glial interaction during the first steps of myelin formation using an in vitro co-culture system of OPCs with cortical neurons. Unlike control OPCs, Cav1.2 deficient oligodendrocytes displayed a simple morphology, low levels of myelin proteins expression and appeared to be less capable of establishing contacts with neurites and axons. Together, this set of in vitro experiments characterizes the involvement of L-type VOCCs on OPC maturation as well as the role played by these Ca$$^{++}$$ channels during the early phases of myelination.

© 2014 Elsevier Inc. All rights reserved.

Introduction

Several studies have addressed the importance of Ca$$^{++}$$ signaling in oligodendrocyte progenitor cell (OPC) differentiation and myelination (Soliven, 2001), as well as in processes extension and OPC migration (Simpson and Armstrong, 1999; Yoo et al., 1999), and in retraction of oligodendrocyte plasma membrane can occur through a number of routes: (1) directly through a variety of ligand-operated channels, such as the $$\alpha$$-adrenergic, P2Y, P2X and glutamate receptors (Kastritsis and McCarthy, 1993; Kirchhoff and Kettenmann, 1992; Patneau et al., 1994); (2) through voltage-operated Ca$$^{++}$$ channels (VOCCs) activated in response to cell membrane depolarization, e.g. increased extracellular K$$^{-}$$; and (3) through other routes such as the opening of store-operated Ca$$^{++}$$ channels in the membrane by the depletion of Ca$$^{++}$$ stores in the endoplasmic reticulum (Alberdi et al., 2005; Belachew et al., 2000; Deitmer et al., 1998; Simpson et al., 1997). This work will focus on Ca$$^{++}$$ influx mediated by VOCCs and the role that these Ca$$^{++}$$ channels play during OPC development and the initial stages of myelination.

VOCCs are a vehicle for impulse generation and propagation in neurons and muscle cells, and thus, their expression in non-excitable cells was surprising. Six types of VOCCs (P/Q, N, L, R and T) have been classified on the basis of electrophysiological and pharmacological properties. Chen et al. (2000) found strong, transient expression of VOCCs in CNS white matter. The immunoreactivity appeared in glial cells along specific pathways of the brainstem, cerebellum and telencephalon. Ultrastructural analysis confirmed that VOCC immunoreactivity was located in oligodendroglial somata, projections, paranodal wraps and loose myelin sheaths (Chen et al., 2000). Electrophysiological recordings performed in our lab have identified low-voltage and high-voltage activated currents in corpus callosum OPCs. The low-voltage and high-voltage activated currents were found to possess the pharmacological and voltage-dependent properties of T-type and L-type VOCCs respectively (Fulton et al., 2010). These

Grant sponsors: NIH/NINDS grant 1R01NS078041-01A1 and National Multiple Sclerosis Society Grant RG4554-A-2.

* Corresponding author at: Hunter James Kelly Research Institute, Department of Pharmacology and Toxicology, School of Medicine and Biomedical Sciences, SUNY at Buffalo, NYS Center of Excellence, 701 Ellicott St., Buffalo, New York 14203, USA.

Fax: +1 716 849 6651.

E-mail address: ppaez@buffalo.edu (P.M. Paez).

http://dx.doi.org/10.1016/j.expneur.2014.12.012
0014-4886/© 2014 Elsevier Inc. All rights reserved.
We have shown that VOCCs affect many Ca<sup>2+</sup> dependent functions in OPCs as a consequence of their ability to modulate intracellular Ca<sup>2+</sup> concentrations. We have found that L-type VOCCs regulate extension/retraction of OPC processes (Paez et al., 2007). Additionally, we have provided direct evidence that L-type VOCC activation increases the amplitude of spontaneous Ca<sup>2+</sup> oscillations in the soma and in the leading process of migrating OPCs leading to an accelerated cell migration by promoting Ca<sup>2+</sup> dependent soma translocation and leading processes formation (Paez et al., 2009a). This mechanism illustrates a key role for VOCCs in the regulation of the rate of OPC migration through spontaneous Ca<sup>2+</sup> oscillations.

The overall goal of this work was to test the hypothesis that voltage-gated Ca<sup>2+</sup> entry promotes OPC maturation and the preliminary steps of myelination in vitro. We used different pharmacological treatments to activate or block voltage-gated Ca<sup>2+</sup> uptake and siRNAs to specifically knock down the L-type VOCC in OPCs. We have found that VOCCs are important regulators of the initial stages of myelination, from process extension to the initial contact with axons. This study will lead to new insights into the factors that govern OPC behavior and could lead to novel approaches to intervene in neurodegenerative diseases in which myelin is lost or damaged.

Materials and methods

Animal experimentation

All animals used in the present study were housed in the UB Division of Laboratory Animal Medicine vivarium, and procedures were approved by UB's Animal Care and Use Committee, and conducted in accordance with the guidelines in the "Guide for the Care and Use of Laboratory Animals" from the National Institutes of Health.

Primary cultures of cortical OPCs

Primary cultures of cortical OPCs were prepared as described by Amur-Umarjee et al. (1993). First, cerebral hemispheres from 1 day old mice were mechanically dissociated and were plated on poly-D-lysine-coated flasks in Dulbecco's modified Eagle's medium and Ham's F12 (1:1 v/v) (Invitrogen), containing 100 μg/ml gentamicin and supplemented with 4 mg/ml dextrose anhydrous, 3.75 mg/ml HEPES buffer, 2.4 mg/ml sodium bicarbonate and 10% fetal bovine serum (FBS) (Omega Scientific). After 24 h the medium was changed and the cells were grown in DMEM/F12 supplemented with insulin (5 μg/ml), human transferrin (50 μg/ml), sodium selenite (30 nM), β-Biotin (10 mM), 0.1% BSA (Sigma), 1% FBS (Omega Scientific) and 1% horse serum (Omega Scientific). After 9 days, OPCs were purified from the mixed glial culture by the differential shaking and adhesion procedure of Suzumura et al. (1984) and allowed to grow on poly-o-lysine coated coverslips in defined culture media (Agresti et al., 1996) plus PDGF (10 ng/ml) and bFGF (10 ng/ml) (Peprotech). OPCs were kept in mitogens (PDGF and bFGF) for 2 days and then induced to exit from the cell cycle and differentiate by switching the cells to a mitogen-free medium (mN2) (Oh et al., 2003) (Fig. 1A). mN2: DMEM/F12 supplemented with n-glucose (4.5g/l), insulin (5 μg/ml), human transferrin (50 μg/ml), sodium selenite (30 nM), T3 (15 nM), β-Biotin (10 mM), hydrocortisone (10 nM), 0.1% BSA, 1% horse serum and 1% FBS.

siRNA knockdown of Cav1.2 and Cav1.3

Twenty-four hours after plating, the cells were transiently transfected with a combination of three different Stealth RNAi™ siRNA duplexes (Invitrogen) specific for Cav1.2 and Cav1.3 (see Table 1 for siRNA duplex sequences). Briefly, 6 pmol of each siRNA duplex was mixed with Lipofectamine™ RNAiMAX (Invitrogen) and the mixture was added to 35 mm Petri dishes containing 80% confluent OPCs. After transfection OPCs were further cultured for 24 h in defined culture media (Agresti et al., 1996) plus PDGF (10 ng/ml) and bFGF (10 ng/ml) and then the cells were switched to a mitogen-free medium (mN2) to induce differentiation (Oh et al., 2003) (Fig. 2A). Control cells were transfected with siRNA designed to minimize sequence homology to any known vertebrate transcript and with a similar GC content as our siRNAs. Additionally, following the transfection protocol described above, OPCs were treated with fluorescein-labeled dsRNA oligomers (BLOCK-iTTM Fluorescent Oligos, Invitrogen) to determine siRNA transfection efficiency.

Immunocytochemistry

Cells were stained with antibodies against several oligodendrocyte markers and examined by confocal microscopy. Briefly, the cells were rinsed in PBS and fixed in 4% buffered paraformaldehyde for 30 min at room temperature. After rinsing in PBS, the cells were permeabilized with 0.1% Triton X-100 in PBS for 10 min at room temperature and then processed for immunocytochemistry following the protocol as outlined by Reyes and Campagnoni (2002). Essentially, fixed cells were incubated in a blocking solution (5% goat serum in PBS) followed by an overnight incubation at 4 °C with the primary antibody. Cells were then incubated with the appropriate secondary antibodies (1:200; Jackson), nuclei were stained with the fluorescent dye DAPI (4',6-diamidino-2-phenylindole; Invitrogen), mounted onto slides with Aquamount (Lerner Laboratories), and fluorescent images were obtained using an Olympus spinning disc confocal microscope (Olympus, IX83-DSU). Quantitative analysis of the results was done counting the antigen-positive and DAPI-positive cells (total number of cells) in 20 randomly selected fields, which resulted in counts of >2000 cells for each experimental condition. The primary antibodies used for immunocytochemistry were against: CC1 (1:300; Calbiochem), MBP (1:1000; Covance), MOG (1:200; Millipore), NG2 (1:400; Millipore), neurofilament M (1:250; Millipore), Olig1 (1:500; Millipore), Olig2 (1:500; Millipore), PDGF (1:200; Millipore), phospho-histone H3 (1:500; Millipore) and PLP (1:200; Thermo Scientific).

Incorporation of bromo-deoxyuridine

Twenty-four hour pulses of 10 μM bromo-deoxyuridine (BrdU) (BD Pharmingen) were applied at 24, 48 and 72 h post siRNA transfection (Fig. 5A). After each BrdU pulse, cells were fixed and immunostained in order to determine the number of positive cells. Cells were fixed in 4% paraformaldehyde in PBS. After treatment with 6 N HCl and 1% Triton X-100 to denature nuclear DNA, the cells were incubated in 0.1 M sodium borate (in PBS and 1% Triton X-100) for 10 min. Immunocytochemistry was done using anti-BrdU antibody (1:1000; BD Pharmingen) and anti-NG2 (1:400; Millipore) with the corresponding fluorescent secondary antibodies. The percentage of BrdU positive cells was estimated on the basis of the total number of NG2 positive cells.

Cell cycle time analysis

To examine cell cycle time, primary cultures of GFP-labeled OPCs were used (Malon et al., 2002). These cultures were incubated in a stage top chamber with 5% CO2 at 37 °C, which was placed on the stage of a spinning disc confocal inverted microscope (Olympus, IX83-DSU) equipped with a motorized stage and an atmosphere regulator. Fluorescent images were obtained at 6 min intervals. Individual clones of mouse OPCs were followed for a period of 30 h. In these time-lapse experiments, ~20 clones were analyzed per experimental condition.
MetaMorph (Molecular Devices) was used in the analysis of video microscopic image sequences. The MetaMorph software allows an investigator to cycle back-and-forth through the movie files frame-by-frame (minute-by-minute), facilitating the accurate determination of event time. To quantitatively analyze the dynamics of cell division, 120 cytokinetic events were randomly selected from movies at different time points. Cell proliferation was assessed by calculating the average cell cycle time (time between birth cytokinesis and division cytokinesis) in different OPC clones. Tracking of cells was performed by visual observation of image sequences as described above. Tracking was performed forwards and backwards in time from each identified cytokinesis event to maximize the number of lineally related cytokinetic events identified. Furthermore, the percentage of cycling OPCs during the first 12 h of the time-lapse experiment was performed by visual observation.

**Table 1**
Sequences of stealth RNA™ siRNA duplexes used for Cav1.2 and Cav1.3 knockdown.

<table>
<thead>
<tr>
<th>siRNA duplex oligoribonucleotide sequence</th>
<th>Cav1.2 siRNA1</th>
<th>Cav1.2 siRNA2</th>
<th>Cav1.2 siRNA3</th>
<th>Cav1.3 siRNA1</th>
<th>Cav1.3 siRNA2</th>
<th>Cav1.3 siRNA3</th>
</tr>
</thead>
<tbody>
<tr>
<td>GCCAUGCAAUUAUCUCAAGUCGUUU</td>
<td>UGACAGUCUGACAGCACGACGUUU</td>
<td>UUAGACAGUGAGUGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>UCAUGCAAUUAUCUCAAGUCGUUU</td>
<td>AACACAGUCUGAGUAUAUCUCAAGGC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AAUAGCAUAUUAUCUCAAGUCGUUU</td>
<td>AAACGACGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>UGACAGUCUGACAGCACGACGUUU</td>
<td>UUAGACAGUGAGUGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AAUAGCAUAUUAUCUCAAGUCGUUU</td>
<td>AAACGACGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AAACGACGACAGCACGACGUUU</td>
<td>UUAGACAGUGAGUGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AAACGACGACAGCACGACGUUU</td>
<td>AAACGACGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AAACGACGACAGCACGACGUUU</td>
<td>AAACGACGACAGCACGACGUUU</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Caspase-3 assay**

NucView 488 caspase-3 substrate, a cell membrane-permeable fluorogenic caspase substrate designed for detecting caspase-3 activity within live cells in real time, was used in accordance with the manufacturer’s recommendations (Biotium Inc). Briefly, OPC primary cultures were incubated in medium containing NucView 488 caspase-3 substrate (final concentration 5 μM) in a stage top chamber with 5% CO2 at 37 °C, which was placed on the stage of a spinning disc confocal inverted microscope (Olympus, IX83-DSU). Fluorescent field images were obtained with a specific GFP filter at 6 min intervals for a total of 24 h. MetaMorph software was used to assess apoptotic cell death by calculating the percentage of caspase-3 positive cells in a total of five experiments on four random fields.

**MIT assay**

The MITT survival assay was performed as described by Mosmann (1983). The sterile solution of MITT (Molecular Probes) was added to all wells and the micro plate was incubated at 37 °C for 45 min. The reaction was stopped by addition of SDS and the product was quantified by spectrophotometry at 570 nm.

**Co-culture of cortical neurons and OPCs**

Cortical neurons were prepared from the brains of 1- to 2-day-old mouse pups using established procedures (Echeverria et al., 2005) with minor modifications. Briefly, brains were removed aseptically...
and placed into DMEM/F12 (Invitrogen). After the brains were dissected, the blood vessels and meninges were carefully removed under a dissecting microscope. Brain cortices were isolated and dissociated by digestion with a solution of 0.05% trypsin (Sigma) containing DNase I (0.06%) in Neurobasal medium for 10 min at 37 °C. The digestion reaction was stopped with Neurobasal medium containing 10% fetal bovine serum (Omega Scientific) and triturated by repeated passages (20 times) through a 10 ml pipette. The cell suspension was filtered through a sterile cell strainer (70 μm; BD Biosciences) into a 50 ml centrifuge tube. The cells were pelleted by centrifugation at 200 g for 5 min, and resuspended in Neurobasal medium plus 2% (v/v) B27 supplemented with 0.25 mM GlutaMax I (Invitrogen), 0.25 mM glutamine (Invitrogen), and 100 μg/ml gentamicin (Omega Scientific). High-density cultures (5 × 10^5 cells, ~2500 cells/mm^2) were plated onto a sterile cell strainer (70 μm; BD Biosciences). The cells were pelleted by centrifugation at 200 g for 5 min, and resuspended in Neurobasal medium plus 2% (v/v) B27 supplemented with 0.25 mM GlutaMax I (Invitrogen), 0.25 mM glutamine (Invitrogen), and 100 μg/ml gentamicin (Omega Scientific). High-density cultures (5 × 10^5 cells, ~2500 cells/mm^2) were plated onto 20 mm^2 tissue culture wells coated with poly-D-lysine (Sigma). The neurons were kept at 37 °C in 95% air 5% CO2 for 7 days in vitro and used for co-cultures. After 7 days in vitro, cortical neuron cultures consisted of neurons essentially free from non-neural cells. Co-cultures were prepared by the addition of OPCs to the cultures of cortical neurons at a density of 3 × 10^5 cells/ml. These cultures were maintained in DMEM/F12, 1% FBS for 7 and 14 days.

Cell morphology assessment

Oligodendroglial cell morphology was evaluated using the procedure described by Sperber and McMorris (2001) with slight modifications. Individual MBP-positive cells were scored according to their morphological complexity in four categories, based on the length and number of primary processes, whether the processes were radially distributed, the relative development of secondary and tertiary processes, and the overall size of the cell, including the process arbor.

RT-PCR

Total RNA was isolated using TRIzol reagent (Invitrogen) according to the instructions of the manufacturer. RNA content was estimated by measuring the absorbance at 260 nm and the purity was assessed by measuring the ratio of absorbance: 260/280 nm. PCR primers for Cav1.2 (F-CAGCTCATGCCAACATGAAT, R-TGCTTCTTGGGTTTCCCATA) and for Cav1.3 (F-AATGGCACGGAATGTAGGAG, R-GACGAAAAATGAGCAGGA) were designed based on published sequences by Rosati et al. (2011) and Xu et al. (2007) respectively. First-strand cDNA was prepared from 1 μg of total RNA using SuperScript™ III RNase H-reverse transcriptase (Invitrogen) and 1 μg of oligo(dT). The mRNA samples were denatured at 65 °C for 5 min. Reverse transcription was performed at 50 °C for 55 min and was stopped by heating the samples at 85 °C for 5 min. The cDNA was amplified by PCR using the Cav1.2 and Cav1.3 isoform-specific primers listed above and PCR Platinum Supersmix reagent (Invitrogen). PCR conditions were as follows: 94 °C for 2 min, 40 cycles of 94 °C for 30 s, 58 °C for 30 s followed by 68 °C for 2 min. After completion of the 40 cycles, samples were incubated at 72 °C for 10 min. A β-actin positive control was run alongside the experimental samples, as well as a negative control with no reverse transcriptase. The PCR products were visualized on an ethidium bromide-stained agarose gel and the bands digitized using a Gel Doc™ EZ System (BioRad).
Quantitative real-time PCR

Total RNA was extracted from primary cultures of pure oligoden-droglial cells and co-cultures of OPCs with cortical neurons using TRIZOL Reagent (Invitrogen), following the manufacturer's protocol. Samples were further purified with treatment by TURBO DNA-free™ (Ambion), followed by a second extraction with phenol-chloroform. RNA was reverse transcribed using iScript cDNA Synthesis Kit (BioRad) then analyzed for various transcript expressions. qPCR was set up using iQ SYBR Green Supermix (BioRad) and performed in triplicate as previously described (Ghiani et al., 2006) on an iCycler MyiQ Real-Time PCR machine (BioRad). Negative controls (samples in which reverse transcriptase was omitted) were amplified individually using the same primer sets to ensure the absence of genomic DNA contamination. PCR amplification resulted in the generation of single bands. Amplification specificity was assessed by melting curves and standard curves made from serial dilutions of control RNA and were used for quantification. Data were normalized to the internal control glyceraldehyde-3-phosphate dehydrogenase (GAPDH). Primers used for real-time PCR were designed based on published sequences by Mattan et al. (2010), Li et al. (2008) and Conrad et al. (2009) (see Table 2 for primer sequences).

**Western blot analysis**

Total protein was collected from primary cultures of pure oligoden-droglial cells and co-cultures of OPCs with cortical neurons. The final protein pellet was homogenized in lysis buffer containing 50 mM Tris–HCl, 0.25% (w/v) sodium deoxycholate, 150 mM NaCl, 1 mM EDTA, 1% (w/v) Triton X-100, 0.1% (w/v) SDS, 1 mM sodium vanadate, 1 mM PMSF, 10 μg/ml aprotinin, 10 μg/ml leupeptin, 10 μg/ml pepstain, and 4 μM sodium fluoride. Western blots were performed as previously described (Ghiani et al., 2010). Twenty-five to thirty-five micrograms of total protein was loaded onto a 4–20% Tris-glycine gel (Invitrogen). Protein bands were detected by chemiluminescence using the Amersham ECL kit (GE Healthcare) with horseradish peroxidase-conjugated secondary antibodies (Cell Signaling). Relative intensities of the protein bands were quantified by scanning densitometry using the NIH Image Software Image J. Equal protein loading was proven to be an effective way to fully activate OPC VOCCs by plasma membrane depolarization (Paez et al., 2010), we used small interfering RNAs (siRNAs) for the Cav1 family of α1 subunits that conducts L-type Ca2+ currents. The α1 subunit of the L-type Ca2+ channel is encoded by Cav1–genes consisting of four different subtypes referred to as Cav1.1 (α1D), Cav1.2 (α1C), Cav1.3 (α1D), and Cav1.4 (α1F); only Cav1.2 and Cav1.3 are expressed in the brain (Hell et al., 1993), and we have found by RT-PCR and Western blot that both Cav1.2 and Cav1.3 are expressed in the brain.

**Calcium imaging**

Methods were similar to those described previously (Paez et al., 2007). Briefly, primary cultures of OPCs were washed in serum and phe-nol red-free DMEM containing a final concentration of 4 μM Fura-2 (AM) (TefLabs) plus 0.08% Pluronic F127 (Molecular Probes) to load dye into the cells, incubated for 45 min at 37 °C, 5% CO2, then washed four times in DMEM and stored in DMEM for 10 min before being imaged. Resting Ca2+ levels were measured in serum-free HBSS containing 2 mM Ca2+ but no Mg2+. Other measurements were made in HBSS. Calcium influx and resting Ca2+ levels were measured on individual cells, and the results were pooled from five separate experiments for each condition. The fluorescence of Fura-2 was excited alternatively at wavelengths of 340 and 380 nm by means of a high-speed wavelength-switching device (Lambda DG4; Sutter Instruments). A spinning disc confocal inverted microscope (Olympus, IX83-DSU) equipped with a CCD camera (Hamamatsu ORCA-R2) measured the fluorescence. Image analysis software (MetaFluor, Molecular Devices) allowed for the selection of several “regions of interest” within the field from which measurements are taken. To minimize bleaching, the intensity of excitation light and sampling frequency was kept as low as possible. In these experiments, measurements were made once every 2 s.

**Statistical analysis**

Normal distributions were tested in each data set using Kolmogorov–Smirnov tests. For data with normal distributions, single between-group comparisons were made by the Student paired t-test, and multiple comparisons were investigated by one-way ANOVA followed by Bonferroni’s multiple comparison tests to detect pair-wise between-group differences. All statistical tests were performed in Graphpad Prism (Graphpad Software). A fixed value of p < 0.05 for one tailed tests was the criterion for reliable differences between groups. Data are presented as mean ± SEM unless otherwise noted.

**Results**

The role of VOCCs on OPC differentiation

The role of VOCCs on OPC differentiation was analyzed in primary cultures of mouse OPCs subjected to different pharmacological treatments in order to activate or block voltage-gated Ca2+ uptake. Calcium influx mediated by VOCCs was induced by depolarization via elevated external potassium (K+). Increasing external K+ to 20 mM has been proven to be an effective way to fully activate OPC VOCCs by plasma membrane depolarization (Paez et al., 2007, 2010). The effects of blocking VOCC dependent Ca2+ influx was studied with the specific L-type channel inhibitor verapamil. Using this in vitro system we followed the morphological differentiation of OPCs (processes formation and branching) and the expression of oligodendrocyte stage specific markers. Our data show that the morphological differentiation as well as the expression of mature oligodendrocyte markers was significantly inhibited in cells treated with verapamil (Figs. 1B, C and D). In contrast, OPCs depolarized with high K+ displayed a more complex morphology and showed a significant increase in the expression of mature markers such as MBP, MOG and CC1 (Figs. 1B, C and D). Interestingly, the maturation of OPCs treated with high K+ in the presence of verapamil was similar to that observed in verapamil treated cultures (Figs. 1B, C and D), suggesting that plasma membrane depolarization have no effect on OPC development if L-type VOCCs are block by specific inhibitors. In order to obtain a direct correlation between these channels and OPC differentiation we knocked down the expression of VOCCs in primary cultures of mouse OPCs. Since more than 90% of the Ca2+ influx after plasma membrane depolarization in OPCs is mediated by L-type Ca2+ channels (Paez et al., 2010), we used small interfering RNAs (siRNAs) for the Cav1 family of α1 subunits that conducts L-type Ca2+ currents. The α1 subunit of the L-type Ca2+ channel is encoded by Cav1–genes consisting of four different subtypes referred to as Cav1.1 (α1S), Cav1.2 (α1C), Cav1.3 (α1D), and Cav1.4 (α1F); only Cav1.2 and Cav1.3 are expressed in the brain (Hell et al., 1993), and we have found by RT-PCR and Western blot that both Cav1.2 and

**Table 2**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward</th>
<th>Reverse</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDCGF</td>
<td>5′-TCTTACGTGAGTCCACGCCCA</td>
<td>5′-CGGTGCTCTTCTCTACCC</td>
</tr>
<tr>
<td>MAG</td>
<td>5′-GGTGCTGACGGACGAGTTG</td>
<td>5′-GGTITGTCGCTGACGCAGCA</td>
</tr>
<tr>
<td>CGT</td>
<td>5′-TCTCGAAGCCTTCCCTTCTCC</td>
<td>5′-CTATTGTCCTTACGCAACAC</td>
</tr>
<tr>
<td>CNPase</td>
<td>5′-AACTTTGACGGCCTCAGTGC</td>
<td>5′-TCAGGACCCAGCAGAAGTAAA</td>
</tr>
<tr>
<td>PFLP</td>
<td>5′-CCACCCCTCTCCGCTTCTT</td>
<td>5′-CAGGGAAAAAGACGACCATGTC</td>
</tr>
<tr>
<td>MBP</td>
<td>5′-ATTATCTCTTGCCCGATAG</td>
<td>5′-CTGGTCTGCTGACGCAGCAACAC</td>
</tr>
<tr>
<td>GAPDH</td>
<td>5′-CTTCATGCCTGCTGCTGCTGCTGCTGC</td>
<td>5′-CTTCATGCTGCTGCTGCTGC</td>
</tr>
</tbody>
</table>
Cav1.3 are expressed in cultured OPCs (Fig. 2C). Twenty-four hours after plating, OPCs were transiently transfected with a combination of three different validated siRNA duplexes specific for Cav1.2 and Cav1.3, the sole pore-forming subunits of the voltage-dependent L-type Ca\(^{2+}\) channel in OPCs (Zhang et al., 2014). siRNAs were selected to target three distinct Cav1.2 and Cav1.3 mRNA regions to enhance silencing (Table 1). A negative siRNA control was designed to minimize sequence homology to any known vertebrate transcript and with a similar GC content as our siRNAs and used for sequence independent effects in all the siRNA experiments performed. To determine the percentage of siRNA-transfected cells, OPCs were treated with fluorescein-labeled dsRNA oligomers. Twenty-four hours after transfection, we found that 100% of the NG2 and Olig2 positive cells displayed some degree of siRNA fluorescent signal (Fig. 2B). The function of Cav1.2/1.3 siRNAs was validated using semi-quantitative RT-PCR and western blotting. Fig. 2C shows a representative RT-PCR and western blot demonstrating that Cav1.2/1.3 siRNA decreased Cav1.2/1.3 mRNAs and proteins in OPCs 72 h after transfection. Additionally, Fig. 2D shows the effect of Cav1.2/1.3 siRNA on L-type Ca\(^{2+}\) uptake in OPCs. Consistent with the RT-PCR and western blot data, Cav1.2 siRNA reduced L-type Ca\(^{2+}\) influx by approximately 75% after plasma membrane depolarization, whereas Cav1.3 siRNA reduced Ca\(^{2+}\) influx by ~20% (Fig. 2D). These data suggest that Cav1.2 is the principal pore-forming subunit of the voltage-dependent L-type Ca\(^{2+}\) channel in OPCs.

Previous reports have shown that gap junctions can mediate the spread of Ca\(^{2+}\) waves between oligodendrocytes and that primary oligodendrocyte cultures, in the absence of astrocytes, display gap junction-mediated calcium signaling (Takeda et al., 1995). To test whether gap junctions play a role in OPC calcium response after plasma membrane depolarization, we performed calcium imaging experiments in the presence of a broad-spectrum gap junction blocker, carbenoxolone (Davidson and Baumgarten, 1988). Under these conditions, the amplitude of the Ca\(^{2+}\) response as well as the percentage of responding cells were similar to those found in control conditions (Fig. 2E). Thus, these data suggest that gap junctions between oligodendrocytes are not involved in the propagation of Ca\(^{2+}\) transients after high K\(^{+}\) stimulation in our experimental culture system.

Using the in vitro system described in Fig. 3A we have monitored the morphological differentiation of OPCs and the expression of oligodendrocyte stage specific markers after mitogen removal. Our data show that morphological differentiation as well as expression of mature oligodendrocyte markers was significantly inhibited in Cav1.2 deficient cells (Figs. 3B-F). Under conditions where OPCs were allowed to differentiate for 48 h, Cav1.2 siRNAs induced a decrease in the proportion of cells that expressed the mature markers MBP and CC1, and an increase in cells that retained immature oligodendrocyte markers such as PDGFr, Olig1 and NG2 (Figs. 3B and C). In addition, real-time PCR experiments performed with total mRNA extracted from Cav1.2 transfected OPCs showed a significant reduction in mRNA levels of several myelin proteins including MAG, PLP, MBP, and ceramide-galactosyl transferase (CGT), a key enzyme involved in the generation and maintenance of the myelin sheath (Fig. 3D). Similar results were obtained in western blot experiments (Fig. 3F). In contrast, the maturation of Cav1.3 knockdown OPCs as well as their expression level of myelin proteins was similar to that observed in control cells (Figs. 3B-F). OPC morphology was evaluated using the procedure described by Sperber and McMorrison (2001). Seventy-two hours after siRNA transfection, individual MBP-positive OPCs were scored according to their morphological complexity in four different categories. In control cultures, the percentage of cells of low and medium complexity was 54% and 36% respectively, while the percentage of cells with medium-high complexity was 12% (Fig. 3E). After 72 h of Cav1.2 siRNA transfection, the percentage of cells with low complexity increased markedly (68%), while there was a significant decrease in the percentage of cells with medium (25%) and medium-high morphology (5%) (Fig. 3E). In agreement with the immunocytochemistry results, the morphological differentiation of Cav1.3 knockdown OPCs was similar to that observed in control cells (Fig. 3E). Furthermore, 24 h after siRNA transfection, OPCs were treated with high K\(^{+}\) in order to activate voltage-gated Ca\(^{2+}\) uptake in Cav1.2 knockdown cells. Two days after high K\(^{+}\) treatment we found no changes in the maturation of Cav1.2 knockdown OPCs compared to non-stimulated transfected cells (Fig. 3C), suggesting that Ca\(^{2+}\) influx mediated by Cav1.2 channels is essential for the effect of plasma membrane depolarization on OPC development.

In order to explore the role of VOCCs in the development of premyelinating oligodendrocytes, cells were transfected with Cav1.2/1.3 siRNAs after two days of mitogen deprivation (Fig. 4A). We have found that after shifting to the differentiation medium (mN2) there is a sharp increase in mature (e.g. CC1, MBP) markers consistent with differentiation of the OPCs (Paez et al., 2009b). Suggesting a role for VOCCs during the first steps of OPC development, our data show that the morphological differentiation and the expression of mature oligodendrocyte markers were not affected in newly generated oligodendrocytes after 72 h of being treated with siRNAs for L-type VOCCs (Figs. 4B-D).

The function of L-type VOCCs on the proliferation and survival of OPCs

We have performed experiments to assess the effect of knocking down the expression of L-type VOCCs on OPC proliferation. Since actively proliferating cells duplicate their DNA content, we labeled proliferating OPCs from control and siRNA-transfected cultures with the thymidine analogue bromo-deoxyuridine (BrdU) over a period of three days. Twenty-four hours pulses of 10 \(\mu\)M BrdU were given at 24 h, 48 h and 72 h after transfection (Fig. 3E). After each BrdU pulse, proliferating progenitors were identified by double immunofluorescence for BrdU and NG2 and the relative number of BrdU\(^{+}\)/NG2\(^{+}\) cells was quantified in each cell population (Figs. 5B and D). We found that OPC proliferation was lower in the Cav1.2 transfected populations than control cultures (Figs. 5B and D). For example, after 48 h the average number of proliferating cells in the Cav1.2 transfected cultures (12%) was significantly lower than that of the control group (23%, \(P < 0.01\)) (Fig. 5D). Similar results were found at 24 and 72 h after transfection (Fig. 5D). Since growth factors played an essential role in OPCs division, parallel experiments were done to assess the proliferation of siRNA-transfected OPCs in the presence of PDGF. OPCs were transfected with siRNA duplexes specific for Cav1.2/1.3 and cultured during three days in defined culture media (Agresti et al., 1996) containing PDGF (10 \(\mu\)g/ml). Under these experimental conditions and in agreement with our previous results after growth factors withdrawal, OPCs lacking Cav1.2 showed decreased levels of proliferation compare to control cells (Fig. 5D). Confirmation of the BrdU cell proliferation results was performed using an anti Phospho-Histone-H3 antibody as a marker of mitotic cells (Wei et al., 1999). In agreement with our previous findings, Cav1.2 knockdown was able to significantly decrease the total number of Phospho-Histone H3\(^{+}\) OPCs at all the time points evaluated (Figs. 5C and E). Moreover, no significant differences were found in the proliferation rate of Cav1.3 transfected OPCs versus control cells (Figs. 5D and E).

Next, we measured the cell cycle times of Cav1.2/1.3 transfected OPCs by performing time-lapse imaging in primary cultures. The OPCs used in these experiments were isolated from transgenic mice expressing GFP under control of the PLP promoter (Mallon et al., 2002). In these mice, GFP expression provides a convenient marker for OPCs in time-lapse experiments. We imaged individual clones of GFP-labeled OPCs for a period of 30 h. In these time-lapse experiments, cell proliferation was assessed by calculating the average cell cycle time in different OPC clones. Cell cycle time was defined as the period between when a cell was first generated by cytokinesis and when that cell subsequently divided, giving two daughter cells (Paez et al., 2009b). Examples of cytokinetic events in GFP-labeled control OPCs tracked for 24 h are shown in Fig. 6A. One hundred twenty cytokinetic events were
randomly selected from movies at different time points. Tracking of cells was performed by visual observation of image sequences as described previously (Paez et al., 2009b). Control OPCs showed a cell cycle time of ~23 h, whereas the cell cycle time of OPCs transfected with Cav1.2 siRNAs was increased to ~28 h and ~27 h after 48 h and 72 h of siRNA transfection respectively (Fig. 6B). Similar experiments were performed in the presence of PDGF. Although there was an increase in the cell replication speed of control OPCs in the presence of PDGF, the cell cycle time of Cav1.2 deficient cells was similar to what we found after growth factors withdrawal (Fig. 6B). Furthermore, as shown in Fig. 6C, the percentage of cycling OPCs during the first 12 h of the time-lapse experiment was lower in the Cav1.2 transfected cultures suggesting that the fraction of proliferating OPCs is decreased after L-type VOCC knockdown. These results indicate by several independent measures that Cav1.2 knockdown diminished OPC proliferation, i.e. decreased the number of mitotic cells and increased the duration of the cell cycle. Additionally, Cav1.3 transfected OPCs showed no significant changes in cell division relative to control (Figs. 6B and C), suggesting that this particular L-type \( \alpha_1 \) subunit is not involved in modulating OPC proliferation.

Fig. 3. Cav1.2 knockdown prevents OPC maturation. (A) OPCs were transfected with siRNA duplexes specific for Cav1.2 and Cav1.3 (siCav1.2/1.3) and grown as described in Fig. 2. 24 h after siRNA transfection, a group Cav1.2 deficient OPCs were treated with high K\(^+\). High K\(^+\) (20 mM) was applied in five consecutive pulses (5 min/each) distributed during the first 5 h of the third day in vitro. (B–C) 72 h after siRNA transfection, OPCs were stained with antibodies against PDGF, Olig1, NG2, MBP and CC1 and the percentage of positive cells in each experimental condition was examined by confocal microscopy. Scale bar = 60 \( \mu \)m (Olig1, CC1); 80 \( \mu \)m (PDGF, NG2, MBP). (D) Real-time PCR experiments were performed using total mRNA extracted from primary cultures of OPCs and specific primers for PDGF, CGT, MAG, PLP and MBP (Table 2). (E) Morphological complexity of MBP-positive cells was scored in four categories. (F) Western blot analysis of myelin proteins expression in OPC primary cultures was performed 72 h after siRNA transfection using P84 as internal standard. Data from three independent experiments are summarized based on the relative spot intensities and plotted as percent of controls. Values are expressed as mean \( \pm \) SEM of four independent experiments. *\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.001 \) vs. respective controls. ICC: immunocytochemistry, qPCR: quantitative real-time PCR, WB: Western blot.
The consequence of L-type VOCC knockdown on OPC survival was examined using the MTT quantitative colorimetric method for cell viability. The results revealed no significant differences between groups 48 h after siRNA transfection (Fig. 7C). Cav1.2 siRNA-transfected OPCs showed no changes in cell viability relative to control (Fig. 7C), suggesting that the decrease in cell maturation and proliferation found in these cultures are not a consequence of cell death. We also examined the role of L-type VOCC on OPC apoptotic cell death in Cav1.2/1.3 deficient OPCs. We used a real-time caspase-3 assay, which detects caspase-3 activation within individual living cells. This assay is

![Fig. 4. Cav1.2 knockdown has no effect on the development of more mature premyelinating oligodendrocytes.](image)

(A) OPCs were transfected with Cav1.2/1.3 siRNAs after being induced to differentiate by mitogen deprivation. (B–C) Premyelinating oligodendrocytes were stained with antibodies against PDGFr, NG2, MBP and CC1 and the percentage of positive cells in each experimental condition was examined by confocal microscopy. Scale bar = 80 μm. (D) Morphological complexity of MBP-positive cells was scored in four categories. Values are expressed as mean ± SEM of four independent experiments. ICC: immunocytochemistry.

![Fig. 5. Decreased cell division in OPCs lacking Cav1.2.](image)

(A) OPCs were transfected with siRNA duplexes specific for Cav1.2 and Cav1.3 (siCav1.2/1.3) and grown either in the presence or absence of PDGF. 24 h pulses of BrdU were begun at 24 h, 48 h and 72 h after siRNA transfection. After each BrdU pulse, cells were fixed and immunostained with anti-BrdU and anti-NG2 antibodies. (B–C) Microphotographs showing NG2+/BrdU+ and P-Histone H3+ cells grown in mN2 at different time points after siRNA transfection. Arrows indicate P-Histone H3+ cells. Scale bar = 60 μm. (D–E) The percentage of NG2+/BrdU+ and P-Histone H3+ cells in each experimental condition was compared with respective controls. Values are expressed as mean ± SEM of four independent experiments. *p < 0.05, **p < 0.01, ***p < 0.001 vs. respective controls.
bi-functional in that it is able to detect both intracellular caspase-3 activity and also stain the cell nucleus, which undergoes morphological changes during the apoptotic process (Paez et al., 2009b). Examples of such measurements in Cav1.2 transfected OPCs are shown in Fig. 7A. Consistent with the MTT results, the combined use of real-time confocal microscopy and the caspase-3 indicator revealed that OPCs transfected with siRNAs for Cav1.2/1.3 showed no increase in the percentage of apoptotic cells relative to controls (Fig. 7B).

The role of VOCCs on oligodendrocyte–neuron interaction

In this group of experiments we used a co-culture system of OPCs with cortical neurons to analyze the role of L-type VOCCs on the early phases of myelination. Twenty-four hours after siRNA transfection, OPCs were added to cortical neuron cultures prepared from the brains of mouse pups (Echeverria et al., 2005). Using this in vitro model of myelination, we analyzed the interaction of OPCs with neurites and the elaboration of myelin membrane structures by these cells as described previously (Paez et al., 2005). Initially, GFP-labeled OPCs (Mallon et al., 2002) were co-cultured with cortical neurons for 7 days and the number of contacts with neurites, including the elaboration of myelin membrane structures, was analyzed by confocal microscopy. As shown in Fig. 8A there was a noticeable change in the morphology of control OPCs, from rounded cells with few processes to a highly branching, reticulated structures. Many cell extensions seemed to be in close contact with neurites (Fig. 8A). Although in Cav1.2 transfected cells, there was a change in cell morphology when co-cultured with cortical neurons, processes formation and elaboration of membranous sheets were much less extensive than in the control cells (Fig. 8B).

Analysis of the morphological complexity according to Sperber and McMorris (2001) showed that knocking down the expression of Cav1.2 markedly decreased the percentage of cells with medium, medium-high and high complexity, and increased the percentage of cells with low complexity (Fig. 8C). Furthermore, the average number of contact points with neurites was significantly reduced when the
co-culture included Cav1.2 transfected OPCs (Fig. 8D). To establish whether siRNA treated OPCs were able to myelinate axons, we evaluated the expression of oligodendrocyte differentiation markers and myelin related proteins by immunocytochemistry, real-time PCR and western blot. Fig. 9 shows examples of NG2 and PLP positive cells after 7 days of co-culture. In the control group nearly all the NG2 expressing cells as well as the PLP positive oligodendrocytes were interacting with neurons (Figs. 9A and C). Although we found a small proportion of NG2 transfected cells interacting with neurons, none of the PLP positive Cav1.2 transfected cells established contact with cortical neurons (Figs. 9B and D). Similarly, control cells expressing MBP interact extensively with cortical neurons and show complex morphology, whereas Cav1.2 transfected cells display a much simpler morphology and nearly no contact points with neurons (Figs. 10A and B). Real-time PCR experiments performed with total mRNA extracted from co-cultures of Cav1.2 deficient cells showed a significant reduction in mRNA levels of myelin proteins (Fig. 10C). In the same line, we found a significant decline in the expression of MBP, MOG, PLP and CNPase and a substantial rise in NG2 expression by western blot in Cav1.2 deficient co-cultures (Fig. 10E). Furthermore, OPC maturation was analyzed in co-cultures at 14 days in vitro. Figs. 11A and B show examples of NG2 and MBP-positive cells grew for 14 days in the presence of neurons. Compared with 7 days old co-cultures, control cells displayed a more complex morphology, higher levels of myelin proteins expression and more contact points with neurons (Figs. 11A–E). On the contrary, two weeks of co-culture did not significantly improve the maturation of OPCs lacking L-type VOCCs, which showed low percentage of cells expressing MBP and PLP and few contact points with neurons (Figs. 11A–E). In agreement with our previous experiments in primary cultures of OPCs, high K+ treatment increases the number and the morphological maturation...
of control MBP-positive cells but do not promote the maturation of Cav1.2 knockdown OPCs in co-culture (Figs. 11F–H). These data suggest that depolarization of neurons and OPCs in co-culture stimulate the maturation of only L-type VOCC expressing OPCs. In summary, this set of in vitro experiments established a key role for the L-type VOCC subunit Cav1.2 in the development and differentiation of OPCs and in the early phases of the oligodendrocyte-neuron interaction.

Discussion

L-type VOCCs are essential for OPC maturation

L-type calcium channels are essential for the normal development of neuronal progenitors playing established roles in regulating gene expression, cell survival and synaptic plasticity (Gallin and Greenberg, 1995; Mao et al., 1999; Marshall et al., 2003; Murphy et al., 1991; Norris et al., 1998; Weisskopf et al., 1999; Yagami et al., 2012). We hypothesized that L-type VOCCs are also important for OPC development, probably by the activation of similar Ca++-dependent intracellular pathways. Clearly indicating that VOCCs are essential for OPC maturation, the morphological differentiation as well as the expression of mature oligodendrocyte markers was significantly repressed in cells treated with specific VOCC inhibitors. In contrast, OPCs treated with high K+ displayed a more complex morphology and showed a significant increase in the expression of mature markers. In the same line, blocking the expression of the α1.2 subunit that conducts L-type Ca++ currents significantly prevents OPC processes extension, morphological maturation and as a consequence, myelin proteins expression. The effect of knocking down the expression of Cav1.2 on OPC differentiation could be interpreted either as inhibition or delay of the normal maturation process. However, since the effect of knocking down Cav1.2 on OPC differentiation persisted three days after growth factor withdrawal in a pro-differentiation culture media, we believe that the absence of L-type VOCCs is inhibiting OPC maturation.

We have found that the two main α1 subunits of the L-type Ca++ channel present in the brain are expressed in OPCs. Using calcium imaging analysis and specific siRNAs we have shown that over 75% of the Ca++ uptake after plasma membrane depolarization is mediated by Cav1.2 containing channels and less than 20% is associated with Cav1.3 expressing VOCCs. Suggesting that Cav1.3 is not involved in modulating oligodendrocyte development, Cav1.3 deficient OPCs show normal expression of myelin proteins, regular morphological maturation and no significant differences in the proliferation rate versus control cells. These data suggest that Cav1.2 is the principal pore-forming subunit of the voltage-dependent L-type Ca++ channel in OPCs and unlike Cav1.3, is essential for proper OPC maturation.

OPCs from the mouse cortex (Verkhratsky et al., 1990; von Blankenfeld et al., 1992) exhibit both low-voltage-activated (LVA) and high-voltage activated (HVA) Ca++ currents (Williamson et al., 1997). Recordings from early postnatal slices revealed the presence of both HVA and LVA Ca++ currents in OPCs located in white matter regions (Berger et al., 1992; Fulton et al., 2010). Several studies have reported that Ca++ responses through VOCCs appear to diminish with maturation of oligodendrocytes from progenitors to mature cells in culture (Berger et al., 1992; Takeda et al., 1995). We have previously shown that OPCs responded to depolarization with large increases in intracellular Ca++ whereas more mature oligodendrocytes responded with a

![Fig. 9. VOCC deficient oligodendrocytes fail to interact with neurons at different developmental stages. OPCs were transfected with siRNA duplexes specific for Cav1.2 (siCav1.2). 24 h after siRNA transfection, control and transfected OPCs were co-cultured with cortical neurons for 7 days. (A–B) Examples of NG2 immunostaining in 7 days old co-cultures. Scale bar = (A) 40 μm; (B) 60 μm. (C–D) Examples of PLP immunostaining in 7 days old co-cultures. Scale bar = 40 μm. Yellow arrows indicate oligodendrocyte-neurite contact points.](image-url)
significantly smaller increase in the Fura-2 signal under the same high K⁺ treatment (Paez et al., 2010). Additionally, voltage-operated Ca²⁺ influx was significantly greater in OPCs from the sub-ventricular zone compared to more mature oligodendrocytes found in the corpus callosum (Paez et al., 2010). A recently published RNA-sequencing transcriptome database of glial, neurons, and vascular cells of the cerebral cortex, reveal high expression of L-type Ca²⁺ channel subunits in OPCs and downregulation of L-type VOCCs in newly formed and myelinating oligodendrocytes (Zhang et al., 2014). The in vitro and in situ data reveal that voltage-operated Ca²⁺ influx present in OPCs and immature oligodendrocytes disappear as the cells matured. It is likely that voltage-operated Ca²⁺ influx plays a role during the first steps of OPC maturation because expression of VOCCs decreases during development. In this work we have established that the development of newly generated oligodendrocytes, unlike OPCs was not affected by the L-type Ca²⁺ channel knockdown indicating that there is a precise and narrow time window in which VOCCs affect OPC maturation and myelination.

Fig. 10. L-type VOCCs are necessary for neuron-glia interaction and myelination in vitro. Co-cultures were prepared as described in Fig. 9. (A–B) Examples of MBP immunostaining in 7 days old co-cultures. Scale bar = 40 μm. Yellow arrows indicate oligodendrocyte-neurite contact points. (C) Real-time PCR experiments were performed using total mRNA extracted from 7 days old co-cultures and specific primers for CNPase, CGT, MAG, PLP and MBP (Table 2). (D) Immunocytochemical analysis of oligodendrocyte differentiation markers and myelin protein expression in 7 days old co-cultures. OPCs were stained with antibodies against PDGFr, NG2, CC1, PLP and MBP and the percentage of positive cells in each experimental condition was examined by confocal microscopy. (E) Western blot analysis of myelin proteins expression in 7 days old co-cultures was performed using P84, GAPDH and β-actin as internal standards. Data from three independent experiments are summarized based on the relative spot intensities and plotted as percent of controls. Values are expressed as mean ± SEM of four independent experiments. *p < 0.01, **p < 0.001 vs. respective controls. ICC: immunocytochemistry, qPCR: quantitative real-time PCR, WB: Western blot.
L-type Ca++ channels modulate OPC proliferation

L-type calcium channels play a key role in the proliferation and differentiation of non-excitable cells such as bone marrow mesenchymal stem cells and T-lymphocytes (Kotturi et al., 2003; Wen et al., 2012). Blocking L-type VOCC significantly inhibited bone marrow mesenchymal stem cell proliferation and the osteogenic differentiation of this cells in vitro (Wen et al., 2012). Furthermore, L-type Ca++ uptake is required for T-lymphocyte activation and proliferation during an immune response (Kotturi et al., 2003). Our data reveal that Cav1.2 knockdown decreased OPC proliferation, as the proportion of NG2 positive OPCs that incorporated BrdU significantly decreased compared with control cells. Cav1.2 deficient OPCs proliferated less robustly and showed a significant increase in the length of the cell cycle after growth factor withdrawal. The differences we noted in cell proliferation are likely explained by changes in the cell cycle duration, since the length of the cell cycle time could progressively increase the probability that an OPC stops dividing (Gao et al., 1998; Tang et al., 2000). Interestingly, the mitotic activity of OPCs lacking Cav1.2 was also diminished in the presence of PDGF. PDGF is the best characterized and the most active proliferative
factor for OPCs. It exerts its effect through interaction with receptors bearing intrinsic tyrosine kinase activity (Taniuchi, 1995). Ca\(^{++}\) signals activated after membrane receptor recruitment is one of the most conserved responses triggered in the target cells in which PDGF exert a mitogenic action (Fatatis and Miller, 1997). In a previous work we have shown that store-operated Ca\(^{++}\) entry as well as Ca\(^{++}\) release from intracellular stores are essential components of the PDGF mitotic mechanism in OPCs (Paez et al., 2009b). Our present data suggest that the absence of Cav1.2 disrupts OPC response to PDGF and more importantly that Ca\(^{++}\) influx mediated by L-type VOCCs is critical for the effect of this growth factor on the cell cycle. Our current hypothesis is that Ca\(^{++}\) uptake mediated by L-type channels modulates OPC division and maturation through independent intracellular pathways. VOCCs seem to be essential for the normal progression of the cell cycle in mitotic OPCs, but after cell cycle exit, the same channels are playing an important role in the maturation of oligodendrocytes.

**L-type Ca\(^{++}\) channels seem to be required in the communication between oligodendrocytes and neurons**


References


