Vitamin D receptor–retinoid X receptor heterodimer signaling regulates oligodendrocyte progenitor cell differentiation

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The mechanisms regulating differentiation of oligodendrocyte (OLG) progenitor cells (OPCs) into mature OLGs are key to understanding myelination and remyelination. Signaling via the retinoid X receptor γ (RXR-γ) has been shown to be a positive regulator of OPC differentiation. However, the nuclear receptor (NR) binding partner of RXR-γ has not been established. In this study we show that RXR-γ binds to several NRs in OPCs and OLGs, one of which is vitamin D receptor (VDR). Using pharmacological and knockdown approaches we show that RXR–VDR signaling induces OPC differentiation and that VDR agonist vitamin D enhances OPC differentiation. We also show expression of VDR in OLG lineage cells in multiple sclerosis. Our data reveal a role for vitamin D in the regenerative component of demyelinating disease and identify a new target for remyelination medicines.

Introduction

Remyelination involves the generation of new myelin sheath–forming oligodendrocytes (OLGs) after primary demyelination in the central nervous system (Franklin and ffrench-Constant, 2008). OLGs are generated from multipotent progenitors referred to as OLG progenitor cells (OPCs), which, in response to demyelination, divide, migrate, and differentiate into mature OLGs (Levine and Reynolds, 1999; Zawadzka et al., 2010; Moyon et al., 2015). In common with other regenerative processes, remyelination efficiency declines with aging, with the result that in chronic demyelinating diseases such as multiple sclerosis (MS), remyelination becomes ineffective (Shields et al., 2000; Goldschmidt et al., 2009). Several lines of experimental evidence indicate impairment of OPC differentiation as the key determinant of the aging effects in remyelination (Wolswijk, 1998; Sim et al., 2002; Woodruff et al., 2004; Kuhlmann et al., 2008; Shen et al., 2008). Thus, the mechanisms regulating OPC differentiation are key to identifying targets for remyelination-enhancing therapy (Kotter et al., 2011).

We previously identified the nuclear receptor (NR) retinoid X receptor (RXR) γ as a positive regulator of OPC differentiation (Huang et al., 2011). The acceleration of remyelination in aged rodents after systemic delivery of an RXR-γ agonist identified RXR-γ as an attractive target for therapeutic remyelination, especially given the drug development activity around this NR in the context of other diseases (de Lera et al., 2007; Pérez et al., 2012). RXR-γ functions when bound to another NR as a heterodimer. Therefore, understanding which RXR-γ heterodimers are involved in the control of OPC differentiation is critically important for fully exploiting the therapeutic potential of RXR signaling.

Results and discussion

RXR-γ-Vitamin D receptor (VDR) complex in OLG lineage cells

RXR-γ binding partners were identified using whole cell lysates obtained from cultures of OPCs or OLGs. OPCs isolated from a mixed glial culture (MGC) were maintained in the presence of PDGF-AA and basic FGF (bFGF). OLGs were generated as described at http://creativecommons.org/licenses/by-nc-sa/3.0/.
by growing OPCs in medium without these two growth factors for 5 d in vitro (DIV). RXRs α and γ, retinoid acid receptors (RARs) α and β, VDR, thyroid hormone receptor β, and peroxisome proliferator activated receptor (PPAR) γ were detected in OPCs and OLGs (Fig. 1 A).

To identify the NRs bound to RXR-γ within OLG lineage cells, we performed coimmunoprecipitation (CoIP) assays. VDR, together with RAR-β and PPAR-γ (unpublished data), were pulled down with RXR-γ in OPC and OLG lysates (Fig. 1, B and D). Because of the association of hypovitaminosis D and MS, we focused on VDR (Ascherio et al., 2012; Burton and Costello, 2015). CoIP assays revealed no differences in RXR-γ–VDR binding in nuclear and cytoplasmic fractions from either OPCs or OLGs (Fig. 1 C). Using a proximity ligand assay (PLA; Duolink), which enables protein binding to be visualized in cells, we detected binding of VDR to RXR-γ in NG2 + OPCs (Fig. 1 E) and O4 + OLG lineage cells (Fig. 1 F).

To determine VDR expression in OLG lineage cells during remyelination, we induced focal demyelination by injecting ethidium bromide into the caudal cerebellar peduncle (CCP) of 2-mo-old rats (Fig. 1 G; Woodruff and Franklin, 1999). We used antibodies to adenomatous polyposis coli (APC; a marker of mature OLGs), Olig2 (an OLG lineage marker), and VDR to analyze VDR cellular localization in normal-appearing white matter (NAWM) and at 5 d post lesion (dpl), when OPC recruitment is maximal, at 14 dpl, when recruited OPCs are undergoing differentiation and new myelin sheaths appear, and at...
21 dpl, when remyelination is complete. APC+ and Olig2+ cells both expressed nuclear VDR (Fig. 1, H and I). The proportion of Olig2+ cells expressing nuclear VDR was decreased at 5 dpl compared with NAWM, but at 14 and 21 dpl, there were no differences with NAWM (Fig. 1 J). Olig2+ cells that did not express APC were taken to be OPCs and immature OLGs. At 5 dpl, there was a significant increase in the density of Olig2+/APC− cells with nuclear VDR expression, indicating a high level of VDR expression in activated recruited OPCs. As the number of OPCs declined at 14 and 21 dpl, there was a corresponding increase in the number of APC+ cells expressing VDR (Fig. 1 M). The majority of APC+ cells (~90%) expressed nuclear VDR expression at all the time points and in NAWM (Fig. 1 L).

### Vitamin D increases OPC differentiation through VDR signaling

We next tested activation of VDR with vitamin D. We treated OPCs for 24 h with vitamin D (1,25-dihydroxyvitamin D3 or calcitriol) and examined proliferation and differentiation using antibodies to Ki67, MBP, and Olig2. Compared with untreated cultures, there was no difference in OPC survival, proliferation, and differentiation or in Olig2 expression (Fig. S2, A–F). Treatment with vitamin D and 9cRA had no additional effect compared with 9cRA treatment alone (Fig. S2, G and H).

We hypothesized that the lack of effect of vitamin D was caused by the presence of saturating concentrations of vitamin D in the culture media resulting from the exposure of OPCs to FBS (which contains traces of steroidal molecules including vitamin D) before isolation from MGCs. To test this, FBS was treated with activated charcoal to remove steroids by adsorption before its addition to the MGC media. We then exposed OPCs grown in charcoal-stripped serum to vitamin D for 24 h. Treatment with 0.1-µM vitamin D increased the number of Olig2+ cells expressing MBP by 80.1% (Fig. 3, A and B) and decreased the number of Olig2+ cells expressing Ki67 by 47.0% (Fig. 3, E and F) without affecting Olig2 expression (Fig. 3 H). MBP protein levels increased by 58.8% after vitamin D treatment (Fig. 3, C and D), and there was no difference in cell survival (Fig. 3 G).

To determine whether the vitamin D effects on OPCs were mediated via VDR, we treated OPCs with vitamin D and the VDR antagonist ZK159222 simultaneously. Vitamin D addition canceled the effect of ZK159222 on proliferation in OPCs derived from MGCs cultured with conventional serum (Fig. 3 I), whereas ZK159222 canceled the effect of vitamin D on differentiation in OPCs derived from MGCs cultured with charcoal-stripped serum (Fig. 3 J). These data indicate that vitamin D enhances OPC differentiation at the expense of OPC proliferation through VDR activation. 9cRA alone did not increase OPC differentiation in charcoal-stripped serum, supporting a key role for VDR in the prodifferentiating effect of 9cRA (Fig. S1 B).

### Blocking VDR signaling impairs OPC differentiation

To test the role of VDR signaling in the OLG lineage, we suppressed VDR activity in OPCs using ZK159222, a synthetic VDR antagonist. ZK159222 binds to VDR with less affinity than vitamin D and prevents binding of coactivators such as SRC1, preventing transcription of target genes (Herdick et al., 2000; Schmitz et al., 2015). The cells were then characterized using antibodies to Olig2 and Ki67 to study the effect of ZK159222 on OPC cell cycle (Fig. 2 A). After 24 h of exposure to ZK159222, there was a twofold increase in the proportion of Olig2+ cells expressing Ki67 (Fig. 2 B). We next asked whether the increase in proliferation was associated with impaired OPC differentiation because there is a mutual exclusivity between OPC proliferation and differentiation (Casaccia-Bonnefil and Liu, 2003). To address this, we cultured OPCs in medium without PDGF-AA and bFGF and exposed them to ZK159222. We then stained for Olig2 and myelin basic protein (MBP), a protein expressed by mature OLGs (Fig. 2 C). After a 24-h treatment with 0.2-µM ZK159222, we observed a 47.6% decrease in the proportion of MBP+ cells within the Olig2+ population compared with controls (Fig. 2 D), whereas 0.5-µM ZK159222 decreased levels by 31.0% (Fig. 2, E and F). Cell death was assessed by propidium iodide staining and showed no differences between control and ZK159222 treatment (Fig. 2 G).

To validate these data, we transfected OPCs with siRNA against VDR and a control nontargeting siRNA. Western blotting showed that transfection with VDR siRNA caused a significant decrease in VDR protein levels compared with the control (Fig. 2 J). After transfection with VDR siRNA, the proportion of Olig2+ cells expressing MBP decreased to 63.4% of that in cells transfected with control siRNA, whereas the proportion of cells expressing Ki67 increased by 80.4% (Fig. 2, H, I, and K). Transfection did not affect cell death (Fig. 2 L). Thus, blocking VDR impaired OPC differentiation while increasing proliferation.

### RXR agonists promote OPC differentiation via VDR heterodimers

Because 9-cis-retinoic acid (9cRA), an RXR agonist, enhances OPC differentiation (Huang et al., 2011), we next asked which component of the RXR−γ−VDR heterodimer was dominant. When OPCs in differentiation medium were treated with the VDR antagonist ZK159222 together with 9cRA, the proportion of MBP+ cells decreased by 51.60% when compared with 9cRA treatment alone (Fig. S1 A). Thus, blocking signaling via VDR affects RXR signaling, and VDR needs be activated for 9cRA to exert its prodifferentiating effect.

### Blocking VDR signaling impairs myelination and remyelination

We next tested the effects of the VDR antagonist in an ex vivo cerebellar slice model of toxin-induced demyelination and remyelination. Cerebellar slices, cultured for 7 d before the addition of the VDR antagonist, were stained for MBP and neurofilament H (NFH), an axonal marker (Fig. 4 A), to assess myelination. Exposure to 0.2-µM ZK159222 decreased myelination by 37%, whereas 2-µM ZK159222 decreased myelination by 52% when compared with controls (Fig. 4 B). We then tested the effects of the VDR antagonist on remyelination. After 7 d in culture, slices were demyelinated with lysolecithin (Birgbauer et al., 2004). The lysolecithin-containing medium was then replaced with normal medium with 1% DMSO as the vehicle control, or with 0.2-µM or 2-µM ZK159222 (Fig. 4 C). We quantified remyelination as the area of MBP and NFH colocalization normalized to the total area of NFH staining. At 6 dpl, only 2-µM ZK159222 caused a decrease in remyelination compared with controls, whereas at 8 dpl, 0.2-µM ZK159222 and 2-µM ZK159222 had 42% and 48% less remyelination, respectively (Fig. 4 D). Thus, VDR activation plays a role in both myelination and remyelination.
VDR is highly expressed in active MS plaques

To assess the relevance of these findings to human disease, we examined the expression of VDR in lesions from individuals with MS. Lesions were classified as periplaque white matter (PPWM), active lesions with remyelination capacity, and chronic lesions with either impaired or complete remyelination (Lucchinetti et al., 2000). Immunostaining was performed on snap-frozen postmortem brain sections obtained from randomly chosen individuals with MS and equivalent controls using an antibody against VDR together with antibodies that stain for the major central nervous system cell types. VDR was expressed in every cell type examined, including Sox10+ and Olig1+ OLG lineage cells (Fig. 5 A and B), major histocompatibility complex class II (MHC-II)–expressing microglia/macrophages (Fig. 5 C), myelin OLG glycoprotein (MOG)–expressing OLGs (Fig. 5 D), NeuN+ neurons (Fig. 5 E), and glial fibrillary acidic protein (GFAP)–expressing astrocytes (Fig. 5 F). The density of VDR-expressing cells in lesion tissue was compared with that of comparable NAWM regions in control subjects without MS. Active lesions contained a 3.6-fold increase in VDR+ cell density when compared with NAWM controls and a 1.3-fold increase when compared with PPWM. Chronic lesions had 71.1% fewer VDR-expressing cells than active lesions. The only remyelinated lesion identified was full of VDR-expressing cells (Fig. 5 J). The density of cells with nuclear VDR localization was 74.7% higher in active lesions than in chronic lesions, 60.2% higher than in PPWM, and 81.5%...
higher than in NAWM controls, whereas no differences in cytoplasmic VDR were detected (Fig. 5 J). Within active lesions, 30.2% of VDR+ cells were also Sox10+ OLG lineage cells, whereas the remaining cells were mainly GFAP+ astrocytes or MHC-II+ monocytes/macrophages. This expression pattern, resembling that of RXR-γ (Huang et al., 2011), suggests a role for VDR in regeneration of MS lesions.

Here, we identify vitamin D and its cognate NR VDR as a positive regulator of OPC differentiation. Although VDR expression has previously been reported in cultured OPCs and in human MS tissue (Baas et al., 2000; Eyles et al., 2005; Smolders et al., 2013), this study reveals that in OLG lineage cells VDR heterodimerizes with RXR-γ, a determinant of OPC differentiation (Huang et al., 2011), and is expressed in OPCs during remyelination (Fig. 1). Using pharmacological and knockdown approaches, we show that blocking VDR impairs OPC differentiation in vitro and during myelination and remyelination (Figs. 2 and 4; Birgbauer et al., 2004; Zhang et al., 2011; Jarjour et al., 2012). Moreover, activating VDR via vitamin D increases OPC differentiation (Fig. 3) in line
with a recent study demonstrating enhanced differentiation of neural stem cells toward the OLG lineage by vitamin D (Shirazi et al., 2015).

Epidemiological data have established an indirect link between the risk of developing MS and vitamin D deficiency (McLeod and Cooke, 1989; Munger et al., 2004, 2006; Soilu-Hänninen et al., 2005, 2008; Ascherio, 2013; Mokry et al., 2015). Thus far, the focus has been on the protective and immunomodulatory role of vitamin D in experimental autoimmune encephalomyelitis, an animal model for MS (Lemire and Archer, 1991; Cantorna et al., 1996; Nataf et al., 1996; Garcion et al., 2003). Our data suggest that hypovitaminosis D in MS patients may be a contributor to remyelination failure.

Previous studies have indicated a role for vitamin D in myelination and remyelination (Goudarzvand et al., 2010; Wergeland et al., 2011; Chabas et al., 2013; Nystad et al., 2014; Montava et al., 2015). These data, together with ours, highlight its potential for remyelination therapies. VDR is also expressed in microglia/macrophages (Fig. 5; Zhang et al., 2014). Because vitamin D decreases inducible nitric oxide synthase expression in microglia (Garcion et al., 1998), it may be one of the mechanisms that shifts the balance between pro- and antiinflammatory states known to be important for controlling remyelination (Miron et al., 2013). Vitamin D also increases microglial activation (Nystad et al., 2014) and amyloid-β phagocytosis (Masoumi et al., 2009), suggesting that it may also facilitate myelin debris clearance and thus remyelination (Kotter et al., 2006). Further investigation into the molecular mechanisms of VDR in remyelination will open up new opportunities for the development of regenerative medicines for demyelinating diseases.

Materials and methods

OPC culture
OPC cultures were prepared from neonatal (postnatal day [P] 0–2) Sprague-Dawley rats as described previously (McCarthy and de Vellis, 1980). MGCs were kept for 11 DIV in DMEM (Gibco), 10% FBS (Biosera), and Mycozap PR-Plus (Lonza) before OPC selection. OPCs were isolated by shaking off nonadherent or loosely adherent cells (rich in OPCs) from the underlying adherent cell monolayer. Upon isolation, a relatively pure population of OPCs (71% Olig2+ [OLG lineage marker], 16% cell death [generally OPCs], 7% GFAP+ [astrocytes], and 5% CD11b+ cells [microglia]) were seeded onto poly-d-lysine–coated (Sigma-Aldrich) 13-mm coverslips for immunostaining, or T25 cell culture flasks for immunoprecipitation and Western blotting. Isolated OPCs were cultured at 37°C in 5% CO₂ in SATO medium (DMEM [Gibco], 1% penicillin/streptomycin, 10 µg/ml BSA, 0.06 µg/ml progesterone, 16.10 µg/ml putrescine, 0.005 µg/ml sodium selenite, 5 µg/ml insulin, and 50 µg/ml holotransferrin [Sigma-Aldrich]). A high proportion of OPCs within the cell culture was maintained by daily addition of 10 ng/ml PDGF-AA and 10 ng/ml bFGF (PeproTech). OLG-rich cultures were obtained by culturing OPCs in SATO media without growth factors. For both OPC and OLG cultures, thyroxine and triiodothyronine were absent from the culture media.

Serum charcoal stripping
For charcoal treatment, FBS was incubated with 2.5% (wt/vol) activated charcoal (Sigma-Aldrich) for 18 h at 4°C and filtered before...
adding it to the MGC medium. Depletion of vitamin D was verified by mass spectrometry using previously described methods (Table S2; Bruce et al., 2013; van den Ouweland et al., 2014). In brief, a standard calibration curve was generated by spiking vitamin D into horse serum previously determined to give no signal. 150 µl of 0.2-M ZnSO₄ was added to each sample followed by vortexing for 1 min. 600 µl methanol was then added with vortexing for a further 5 min. The precipitate was sedimented by centrifugation. A solid phase extraction plate (Oasis HLB µElution; Waters) was conditioned with 200 µl methanol followed by 60% (vol/vol) methanol. 600 µl of each supernatant from sample pretreatment was loaded followed by two wash steps: 200 µl of 5% (vol/vol) methanol and then 200 µl of 60% (vol/vol) methanol. Retained analytes were eluted and transferred to amber glass vials before liquid chromatography (LC)–mass spectrometry analysis. The LC–mass spectrometry/mass spectrometry system was comprised of an ultra-performance LC system (Acquity I-Class; Waters) coupled to a triple quadrupole mass spectrometer (Xevo TQ-S; Waters) operating in positive electrospray ionization mode and using MassLynx software version 4.1 (Waters). 20 µl of each sample was injected into a 2.1 × 50-mm ultra-performance LC ethylene-bridged hybrid phenyl column (Acquity; Waters) with 1.7 µl maintained at 35°C. The data were independently verified by SAS Bone Metabolic Laboratories.

CoIP

CoIP was performed according to the instructions on the cross-linking CoIP kit (Thermo Fisher Scientific). In brief, OPCs and OLGs were grown for 5 DIV and lysed with immunoprecipitation lysis wash buffer (0.025-M Tris, 0.15-M NaCl, 0.001-M EDTA, 1% NP-40, and 5% glycerol; Thermo Fisher Scientific) with Halt protease and phosphatase inhibitor cocktail (Thermo Fisher Scientific). Protein concentration was measured by bicinchoninic acid assay (Thermo Fisher Scientific), and immunoprecipitation was performed following the Pierce cross-link immunoprecipitation kit protocol (Thermo Fisher Scientific). In brief, 10 µg RXR–γ antibodies (ab15518; Abcam) were bound to protein A–coated agarose beads. Between 200 and 600 µg of protein were incubated overnight at 4°C with the antibody. Elution was analyzed by Western blotting.
Western blotting
Protein lysate was mixed with NuPAGE loading buffer and NuPAGE reducing agent (Invitrogen) and boiled for 10 min at 95°C. 5–15 µg of protein (lysed as described in the CoIP section) or total CoIP elution was loaded in 4–12% Bis-Tris gels (NuPAGE Novex; Invitrogen). After, electrophoresis gels were transferred to nitrocellulose membranes (GE Healthcare) and blocked with 5% milk in 0.1% PBS with Tween 20 (PBST). Membranes were incubated overnight at 4°C with the previously mentioned antibodies diluted in 5% nonfat dry milk in PBST. Upon three PBST washes, membranes were incubated with peroxidasel conjugated secondary antibodies (Dako) for 2 h at 20–26°C and developed with ECL (GE Healthcare) or Supersignal (Thermo Fisher Scientific) depending on the antibody. Western blot analysis was performed measuring the integrated density of the corresponding bands and normalizing them to β-actin (Sigma-Aldrich) control or CoIP control antibody, which is an antibody against the protein for which the pull-down has been done (RXR-γ in most cases).

Duolink
After fixation in 4% PFA, cells were stained using the Duolink in situ CoIP protocol (Olink; Sigma-Aldrich). Cells were blocked with Olink blocking solution for 1 h at 20–26°C. Primary antibodies against NRs were diluted in antibody diluent solution and incubated for 1 h at 20–26°C. PLA probe minus and PLA probe plus conjugated secondary antibodies were added and incubated for 1 h at 37°C. Next, PLA probes for the secondary antibodies were ligated by a 30-min incubation at 37°C with ligation solution, and the DNA signal was amplified by incubating the cells for 100 min at 37°C with DNA polymerase and fluorescent deoxynucleotide solution. OLG lineage marker antibodies were then added following the usual immunocytochemistry protocol.

siRNA transfection
After shake off, OPCs were kept for 2 DIV with growth factors, and then the medium was changed to SATO medium without penicillin/streptomycin overnight. Cells were transfected with 50-nM siRNA for VDR or for nontargeting control (GE Healthcare) using 1% Lipofectamine siRNAMAX (Invitrogen) diluted in Opti-MEM (Gibco) according to the manufacturer’s protocol. After 6 h of transfection, the medium was replaced by normal SATO medium, with or without growth factors and without thyroxine or triiodothyronine. After 48 h of transfection, cells on coverslips were fixed with 4% PFA or lyzed with previously mentioned immunoprecipitation lysis/wash buffer and subjected to Western blot analysis.

Cell death assessment
Cells were incubated for 20 min at 37°C with 10 µg/ml propidium iodide (Invitrogen) after the corresponding treatment. Cells were then washed with PBS and fixed for 10 min with 4% PFA as previously described (Leeceur, 2002).

Immunocytochemistry
Cells were fixed for 10 min with 4% PFA and blocked with 5% normal goat serum (NGS; Sigma-Aldrich) with 0.1% Triton X-100 in PBS for 1 h at 20–26°C. Then, cells were incubated for 1 h at 20–26°C with antibodies for Olig2 (1:500; EMD Millipore), NG2 (1:200; EMD Millipore), MBP (1:500; Serotec), Klf6 (1:200; Vector Laboratories), or O4 (1:1,000; R&D Systems) in blocking solution. Upon primary antibody washes, fluorescently labeled secondary antibodies (1:500; Alexa Fluor (AF) 488 and AF568; Invitrogen) diluted at 1:500 in 5% NGS with 0.1% Triton X-100 were incubated for 1 h at 20–26°C. Nuclei were stained with 2 µg/ml Hoechst (Sigma-Aldrich) for 5 min, and two washes with PBS were performed before mounting the coverslips with Fluoromount G (SouthernBiotec).

Focal demyelination
Bilateral focal demyelination lesions were induced in the CCP of 2–3-mo-old female Sprague-Dawley rats in compliance with United Kingdom Home Office regulations (project license 70/7715). This involved stereotactic injections of 4 µl of 0.01% ethidium bromide (vol/vol) into the CCP. Animals were then perfused with 4% PFA at 5, 14, and 21 dpl. Once perfused, the removed brains were kept in 20% sucrose and embedded in optimal cutting temperature compound (Taab). Sections were cut to 12 µm and frozen at −80°C until further analysis.

Immunohistochemistry
12-µm sections were dried for 1 h at 20–26°C and washed with PBS before antigen retrieval. Antigen retrieval was performed by incubating the sections for 10 min at 75°C with preboiled 1x citrate buffer, pH 6.0, and antigen retriever (Dako). Then, slides were incubated for 6 h at 20–26°C with 5% normal donkey serum with 0.1% Triton X-100 and anti-Olig2 antibodies (1:200; R&D Systems). Slides were then incubated overnight at 4°C with antibodies against APC (1:200; EMD Millipore) and VDR (1:100; Santa Cruz Biotechnology, Inc.) in blocking solution. The remaining primary antibodies were washed away, and sections were incubated for 2 h at 20–26°C with 1:500 AF-conjugated secondary antibodies (1:500 donkey AF488 anti-goat, 1:500 donkey AF568 anti-rabbit, and 1:500 donkey AF647 anti-mouse [Invitrogen]) in 5% normal donkey serum with 0.1% Triton X-100. Nuclei were stained with 1 µg/ml Hoechst G for 5 min, and slides were mounted after washes with Fluoromount G.

Organotypic cerebellar slices
Cerebellar slice cultures were prepared as previously described (Birg-bauer et al., 2004). In brief, P9 rat cerebellum was isolated, and 300-µm sections were obtained using a tissue chopper (Mcllwain). Slices were then incubated at 37°C with 5% CO2 in organotypic slice medium composed of 50% HBSS (Gibco), 25% heat-inactivated horse serum (Gibco), 25% basal Eagle medium (Gibco), 5 mg/ml glucose, 1x glutamax (Gibco), and 1x Mycozap Plus-PR (Lonza). After 1 wk in culture, slices were demyelinated with 0.5 mg/ml lyssolecithin (Sigma-Aldrich) for 16 h, and then medium was replaced with medium with the corresponding treatment. Treatment was replaced every other day, and slices were collected at 6 and 8 dpl and fixed with 4% PFA for 1 h. Then, slices were stored in 1x PBS at –20°C for immunostaining. In the case of myelination experiments, medium was replaced at 8 DIV with medium with the corresponding treatments, and no lyssolecithin was added. For immunostaining, slices were blocked for 1 h with 10% NGS and 0.5% Triton X-100 in agitation and incubated overnight with antibodies against MBP (1:500; Serotec) and NFH (1:500; Abcam) in blocking solution. Slices were washed three times with 0.1% PBS and Triton X-100, and then incubated with secondary antibodies (1:300; AF488 and AF568; Invitrogen) for 2 h at 20–26°C. The secondary antibodies were removed by washing with 0.1% PBS and Triton X-100 and then Hoescht (1 µg/ml final concentration) was used for nuclei staining. Slices were mounted with Fluoromount G onto poly-n-lysine slides and covered with a 20-mm coverslip (VWR).

MS tissue samples and immunohistochemistry
MS patient brain samples were obtained from the UK MS Tissue Bank (Table S1). Control brain samples from individuals without evident neurological disease were also obtained from the same source. Tissues were collected with the donor’s fully informed consent after ethical approval. Fixed frozen MS sections were hydrated in PBS and microwaved in antigen retrieval solution (Vector Laboratories) according to the manufacturer’s protocol. Sections were preincubated in blocking buffer containing 10% NGS and 0.1% Triton X-100 in PBS for 1 h. Primary antibodies were incubated overnight at 4°C: GFAP (1:100; EMD
Millipore), Olig1 (1:50; R&D Systems), MOG (1:100; a gift from C. Linnington, University of Glasgow, Scotland, UK), NeuN (1:50; EMD Millipore), MHC-II (1:50; Dako), Sox10 (1:50; R&D Systems), and VDR (1:100; Santa Cruz Biotechnology, Inc.). Primary antibodies were extensively washed with 0.1% PBS and Triton X-100 and incubated with appropriated secondary antibodies. Quantification of VDR+ cells was performed on high magnification fluorescent images of studied regions of interest using ImageJ version 1.44 software (National Institutes of Health) in at least three serial MS sections 100 μm apart from five MS and five control cases (Fig. 3 A). 6–10 pictures were taken for each MS region of interest, including active chronic lesions, PPWM, as well as NAWM from nonneurological controls. The number of immunopositive cells per square millimeter was calculated.

Quantification

For in vitro data obtained with OPC cultures, five randomly chosen areas of the coverslip were imaged per condition with the 20x objective (NA 0.30; Ph 1), or 40x (NA 0.50; Ph 1) for Duolink, using a fluorescence microscope (AxioVision Observer A1; Carl Zeiss) (Axio-Cam HRC; Carl Zeiss), and counting was done manually using the cell counter plugin in ImageJ. Organotypic cerebellar slices and in vivo CCP demyelination lesions were imaged using a confocal (SP5; Leica) and Advanced Fluorescence software, version 2.7.3.9723 (Leica). In the case of cerebellar slices, four independent slices were analyzed per condition and per experiment. Five images of randomly chosen areas of each slice were imaged using the 40x objective (NA 1.25) with 1.5x digital zoom, 1,024 x 1,024 resolution, and 1.75–2-μm thickness stacks. For analysis, a macro was created in ImageJ that measured the MBP–NFH colocalization area and the total NFH area. For CCP lesion images, the whole lesion of three sections was imaged per animal with the 20x objective (NA 0.70), 1,024 x 1,024 resolution, and 1.5–2-μm stacks. Using ImageJ software, the lesion area was delineated and measured, and the number of different cell types within the lesion was counted manually with the cell counter plugin. All the images were acquired at room temperature.

Statistics

Statistical analysis was performed using Prism 5.0 (GraphPad Software) and SPSS Statistics 20.0 (IBM). Mean ± SEM are shown in all the graphs. Agonist and antagonist treatments were analyzed by a one-way analysis of variance (ANOVA) followed by a Dunnett’s post-hoc test, and VDR characterization in CCP lesions was analyzed by an one-way ANOVA followed by a Bonferroni post-hoc test or a Kruskal–Wallis test in cases of lack of normality (assessed using a Shapiro–Wilk test). siRNA experiments and charcoal treatment effect in MGCs and OPCs were analyzed using a bilateral unpaired t test or a Kruskal–Wallis test in cases of lack of normality (assessed using a Shapiro–Wilk test). All the images were acquired at room temperature.

Online supplemental material

Fig. S1 shows the effect of blocking VDR with the antagonist ZK159222 in the presence or absence of 9cRA, an RXR agonist. Fig. S2 shows the absence of effect of vitamin D in OPCs derived from MGCs exposed to normal serum in proliferation and differentiation. Table S1 provides the clinical data of the MS patients and matching controls used for this study. Table S2 shows the two independent measurements of vitamin D performed in the serum before and after charcoal treatment. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201505119/DC1. Additional data are available in the JCB DataViewer at http://dx.doi.org/10.1083/jcb.201505119.dv.

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