GD1a Overcomes Inhibition of Myelination by Fibronectin via Activation of Protein Kinase A: Implications for Multiple Sclerosis

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Remyelination failure by oligodendrocytes contributes to the functional impairment that characterizes the demyelinating disease multiple sclerosis (MS). Since incomplete remyelination will irreversibly damage axonal connections, treatments effectively promoting remyelination are pivotal in halting disease progression. Our previous findings suggest that fibronectin aggregates, as an environmental factor, contribute to remyelination failure by perturbing oligodendrocyte progenitor cell (OPC) maturation. Here, we aim at elucidating whether exogenously added gangliosides (i.e., cell surface lipids with a potential to modulate signaling pathways) could counteract fibronectin-mediated inhibition of OPC maturation. Exclusive exposure of rat oligodendrocytes to GD1a, but not other gangliosides, overcomes aggregated fibronectin-induced inhibition of myelin membrane formation, in vitro, and OPC differentiation in fibronectin aggregate containing cuprizone-induced demyelinated lesions in male mice. GD1a exerts its effect on OPCs by inducing their proliferation and, at a late stage, by modulating OPC maturation. Kinase activity profiling revealed that GD1a activated a protein kinase A (PKA)-dependent signaling pathway and increased phosphorylation of the transcription factor cAMP response element-binding protein. Consistently, the effect of GD1a in restoring myelin membrane formation in the presence of fibronectin aggregates was abolished by the PKA inhibitor H89, whereas the effect of GD1a was mimicked by the PKA activator dibutyryl-cAMP. Together, GD1a overcomes the inhibiting effect of aggregated fibronectin on OPC maturation by activating a PKA-dependent signaling pathway. Given the persistent presence of fibronectin aggregates in MS lesions, ganglioside GD1a might act as a potential novel therapeutic tool to selectively modulate the detrimental signaling environment that precludes remyelination.

Key words: fibronectin; ganglioside; multiple sclerosis; oligodendrocyte; remyelination

Significance Statement

As an environmental factor, aggregates of the extracellular matrix protein fibronectin perturb the maturation of oligodendrocyte progenitor cells (OPCs), thereby impeding remyelination, in the demyelinating disease multiple sclerosis (MS). Here we demonstrate that exogenous addition of ganglioside GD1a overcomes the inhibiting effect of aggregated fibronectin on OPC maturation, both in vitro and in vivo, by activating a PKA-dependent signaling pathway. We propose that targeted delivery of GD1a to MS lesions may act as a potential novel molecular tool to boost maturation of resident OPCs to overcome remyelination failure and halt disease progression.

Introduction

Multiple sclerosis (MS) is characterized by inflammation, multiple demyelinated lesions, axonal degeneration, and reactive glial scar formation (Compston and Coles, 2008). Remyelination, involving the regeneration of myelin sheaths and the restoration of functional deficits, is a common feature at early stages of MS, but...
generally fails at later stages, resulting in secondary axonal degeneration and disease progression (Franklin and ffrench-Constant, 2008; Goldschmidt et al., 2009). Current therapies rely primarily on immunomodulating strategies, thereby delaying the time and severity of new lesion formation (Compston and Coles, 2008; Franklin and ffrench-Constant, 2008; Irvine and Blakemore, 2008; Winkelmann et al., 2016). However, these therapies often fail in patients with progressive MS, indicating that once the cascade of events leading to axonal degeneration has been initiated, the suppression of inflammation does not disrupt and/or interfere with clinical disease progression. Hence, the development of treatments that preserve axons by means of remyelination is an essential therapeutic goal.

Remyelination in MS is often insufficient, despite the presence of oligodendrocyte progenitor cells (OPCs) in 70% of the lesions (Lucchinetti et al., 1999; Chang et al., 2002; Franklin and ffrench-Constant, 2008; Münzel et al., 2013). The failure of OPC maturation can be attributed to numerous factors and may include axonal damage and/or dysregulation of the cellular and extracellular signaling environment within the lesions (Franklin and ffrench-Constant, 2008). Recent progress has highlighted the importance of the aberrant expression of distinct extracellular matrix (ECM) components in the injured environment that contributes to remyelination failure in MS lesions (Back et al., 2005; Lau et al., 2012; Stoffels et al., 2013). Our previous findings revealed that the persistent presence of fibronectin (Fn) aggregates in MS lesions constitutes a most perturbing environmental factor as it impairs OPC maturation and therefore remyelination. Thus, while transiently expressed (dimeric) Fn is a physiologically relevant factor, associated with the regulation of OPC proliferation following CNS demyelination (Stoffels et al., 2015), in MS, Fn aggregation occurs, causing its persistent presence in the lesions, and failure of remyelination is observed (Stoffels et al., 2013). Together, these findings thus prompt for the development of therapeutic approaches, relying on the application of Fn (aggregate) antagonists, capable of overcoming the devastating aggregate-induced signaling defect that causes remyelination failure.

It has been well documented that gangliosides, (largely) cell surface localized glycosphingolipids, act as (signaling) modulators of cell functions, including cell growth, apoptosis, and differentiation (Lopez and Schnaar, 2009; Sonnino and Prinetti, 2010). In addition, gangliosides are known to interfere with the interaction of Fn and integrins, their cell surface receptors (Merzak et al., 1995; Wang et al., 2001). Hence, we hypothesized that gangliosides may counteract the perturbing effect of Fn on (re)myelination. Here, we demonstrate that exogenous addition of ganglioside GD1a, but not other gangliosides, eliminates the inhibition of (re)myelination by Fn aggregates via a protein kinase A (PKA)-dependent signaling pathway that promotes OPC maturation. Accordingly, these findings therefore highlight the exciting possibility of considering the application of GD1a as a novel tool to promote remyelination in MS lesions.

Materials and Methods

Cell cultures

Primary oligodendrocyte cultures. Primary rat OPCs were isolated and cultured as described previously (Bibisi et al., 2012). OPCs were plated on the indicated substrates [5 μg/ml poly-1-lysine (PLL), 10 μg/ml plasma Fn (Sigma-Aldrich), or 10 μg/ml aggregated Fn (see below)] and cultured for 2 d in defined Sato medium (Maier et al., 2005) containing 10 ng/ml PDGF-AA (Peprotech) and FGF-2 (Peprotech). OPC differentiation was induced by switching to Sato medium containing 0.5% FCS (Bodico). Cells at the indicated developmental stage were treated with vehicle (ethanol), gangliosides (10 μM, Dr. S. Sonnino, University of Milan, Milan, Italy), dibutyryl-cAMP (dbcAMP; 1 mM, Sigma-Aldrich), and/or H89 (20 μM; Merck Millipore).

Primary dorsal root ganglion neuron–oligodendrocyte cocultures. Dorsal root ganglion neurons (DRGNe) were dissected from 15- to 17-d-old Wistar rat embryos and cultured as described previously (Stancic et al., 2012). Fn aggregates (10 μg/ml) were added 1 h before OPCs were seeded on 16 d in vitro DRGNe, and gangliosides (2 μM) and vehicle (ethanol) were added 1 h after seeding and at every medium change. OPCs were cocultured with the DRGNe for 14 d with medium changes every second day.

Organotypic cerebellar slice cultures. Rat organotypic cerebellar slice cultures were generated from neonatal cerebellum as described previously (Miron et al., 2010). Briefly, cerebella were dissected from 1- to 2-d-old Wistar rats and cut into 300 μm sagittal sections using a tissue chopper. Slices were plated on Millipore-Millicel-CM culture inserts (Merck Millipore) with four slices per insert. Culture media were composed of 50% MEM (Invitrogen), 25% heat-inactivated horse serum (Invitrogen), 25% Earle’s balanced salt solution (Invitrogen), and 6.5 mg/ml glucose (Sigma-Aldrich). Inserts were transferred into fresh media every 2–3 d. Slices were left untreated to allow the clearance of debris and remyelination to occur. To induce demyelination, 21-d-old slice cultures were treated with lyssolecithin (0.5 mg/ml; Sigma-Aldrich) for 17 h and allowed to recover for 2 d. To induce endogenous aggregation of Fn, demyelinated cultures were treated with the Toll-like receptor 3 (TLR3) agonist polyinosine-polycytidylic acid (poly(I:C); 50 μg/ml; GE Healthcare) for 2 d. The next day, slice cultures were treated with gangliosides (10 μM) or vehicle (ethanol) and at every medium change (every 2–3 d) for 21 d.

Cuprizone-induced demyelination

Animal protocols were approved by the Institutional Animal Care and Use Committee of the University of Groningen (Groningen, The Netherlands). Nine-week-old male C57BL/6 mice (Harlan Laboratories) received a stereotactically placed intracranial cannula (brainlink) located on top of the corpus callosum (coordinates relative to bregma: antero-posterior ~0.94 mm; lateromedial ~0.80 mm; dorsoventral ~1.55 mm) under isoflurane anesthesia. The mice received perioperative and postoperative analgesia for 2 d with xylazine and carprofen, respectively. Animals were individually housed and subjected to a 0.2% (w/w) cuprizone (Sigma-Aldrich) diet inducing robust and reproducible demyelination of the corpus callosum (Gudi et al., 2014). Animals received a fresh supply of cuprizone-containing food ad libitum on a daily basis. After 5 weeks, the animals received an injection of aggregated Fn (1.5 μg/3 μl) through the cannula and subsequently returned to standard chow. Two and three days after injection of aggregated Fn, vehicle or gangliosides (10 μM) were injected through the same cannula (3 μl). Animals were killed 5 d after Fn aggregate injection by means of transcardial perfusion under isoflurane anesthesia with buffered paraformaldehyde solution (4%). All buffers were pretreated with diethylpyrocarbonate to prevent RNA degradation. The brains were cryosectioned in the coronal plane at 12 μm thickness. Demyelination of the corpus callosum was confirmed by Sudan Black (0.1% in 70% ethanol) for 5 min.

Generation of fibronectin aggregates

Fn aggregates were generated from primary rat astrocytes as described previously (Stoffels et al., 2013). Briefly, astrocytes were treated with 50 μg/ml poly(I:C) for 48 h and lysed in water for 3 h at 37°C, and the remaining deposits were scraped in DOC buffer [2% deoxycholate, 2 mM EDTA, and 5 mM Tris-HCl pH 7.4, containing a protease inhibitor cocktail (Complete Mini, Roche)]. Fn aggregates, verified by Western blot, were obtained by centrifugation at 16,300 × g for 30 min and dissolved in PBS by sonication.

Adhesion assay

Adhesion assays were performed as described previously (Baron et al., 2014). Non-tissue culture 96-well plates were coated with Fn or PLL. For “masking” experiments, the coating was followed by 1 h with coating with gangliosides (10 μM) or vehicle (ethanol). For “blinding” experiments, OPCs were preincubated with the indicated gangliosides (10 μM) or vehicle (ethanol) for 30 min at 37°C. OPCs were seeded at a density of
10,000 cells/well and left to adhere for 1 h at 37°C. Adhesion is determined as a percentage of the 100% adhesion value.

**MTT and LDH assay**

Mitochondrial dehydrogenase activity (MTT assay) was measured as previously described (Bisbi et al., 2012). Briefly, OPCs were seeded into plasma Fn-coated 24-well plates in Sato medium at 50,000 cells/well. At the immature stage (3 d after initiating differentiation), the cells were treated with the indicated gangliosides (10 µM) or vehicle (ethanol, controls) (1). After 4 d, the cells were incubated for 3–4 h with MTT (0.5 mg/ml; Sigma-Aldrich) in Sato medium. The medium was removed, and the MTT-formazan crystals were collected in DMSO (Sigma-Aldrich), and the absorption at 520 nm was determined. In parallel, a commercially available lactate dehydrogenase (LDH) assay kit (Roche) was used to quantify LDH release into the medium from the same cells as used for the MTT assay. To this end, the medium was centrifuged for 7 min at 7000 rpm, and LDH release was assayed according to the manufacturer instructions.

**Proliferation assay**

Cell proliferation was measured by determining the incorporation of the thymidine analog bromodeoxyuridine (BrdU), as described previously, with minor modifications (Stancic et al., 2012). Cells were plated in PLL- or Fn-coated eight-well chamber slides at a density of 25,000 cells/well. After 1 h, cells were exposed to the indicated gangliosides (10 µM) or vehicle [ethanol, control (ctrl)] in the presence of growth factors PDGFAA (10 ng/ml) and FGF-2 (10 ng/ml) and 10 µM BrdU. After 16 h, BrdU incorporation was detected using a BrdU proliferation detection kit (Roche) according to the manufacturer instructions. Nuclei were stained with DAPI (1 µg/ml, Sigma-Aldrich). Proliferation is expressed as the percentage of BrdU-positive cells of total (DAPI-stained) cells, with at least 500 cells counted per condition.

**Migration assay**

Migration assays were performed in 24-well Falcon transwell cell culture inserts containing membranes with a pore size of 8 µm, as described previously (Stoffels et al., 2015). Both sides of the transwell membrane were precoated with PLL or Fn. OPCs were plated at 100,000 cells/well and allowed to migrate for 24 h. The upper chamber contained gangliosides (10 µM) or vehicle (ethanol, controls). Nonmigrated cells were removed with a cotton swab from the top compartment of the chamber. Remaining cells on the bottom of the membrane were fixed with 4% paraformaldehyde in PBS for 15 min at room temperature. DAPI-stained migrated cells were visualized with a TissueGnostics system, and the number of migrated cells per square millimeter were analyzed using TissueQuest Cell Analysis Software (TissueGnostics).

**[^3]H-GD1a**

Tritium-labeled GD1a ([^3]H-GD1a; 1.28 Ci/mmol) was added to cells for 24 or 96 h before collection. Gangliosides were isolated from the cell pellet as described previously (Senn et al., 1992), spotted onto high-performance thin layer chromatography plates, and separated in CHCl3/CH2OH/0.2% (w/v) CaCl2 (11:9:2, v/v/v). Gangliosides were visualized by exposure to iodine vapor, scraped from the plate, dispersed in an UltimateGold (PerkinElmer) scintillation cocktail, and counted in a liquid scintillation counter (LS 6500, Beckman Coulter). Data are shown as the percentage of total activity.

**PamChip peptide array analysis**

Serine/threonine kinase (STK) activity profiles were generated with the STK PamChip peptide microarray system (PamGene International; Eriksson et al., 2014; Shi et al., 2016). Cells were lysed in M-PER Mammalian Extraction Buffer containing phosphatase and protease inhibitors (Termo Fisher Scientific), incubated on ice for 15 min, and centrifuged at 10,000 × g for 15 min at 4°C. The PamChip assay was performed on the PamStation12 according to manufacturer instructions. In short, the PamChip arrays were blocked with 2% bovine serum albumin (BSA). The reaction mixture contained 0.4 mM ATP and 2 µg of protein lysate per array supplemented with protein kinase buffer, BSA, and STK antibody mix (FITC-labeled). To account for differences in protein input, signal intensities were normalized to the array mean. After a visual check of correct peptide spot identification, data analysis and quantification were performed using Bionavigator software (PamGene International) and the BRB-array tools version 4.5.0. The Phospho-ELM BLAST tool version 9.0 was applied to determine potential upstream kinases.

**PKA activity assay**

OPCs were plated at a density of 350,000 cells/35 mm Fn-coated dish and allowed to differentiate for 6 d. Cells were exposed to 10 µM GD1a and lysed at 4°C in MOPS-[α-(n-morpholino)propanesulfonic acid]-lysine buffer (20 mM MOPS, 5 mM EGTA, 2 mM EDTA, 1% NP-40, 1 mM DTT, and MS-SAFE protease and phosphatase inhibitor (Sigma-Aldrich)). The cytosolic fraction was obtained upon centrifugation at 13,000 rpm for 15 min at 4°C. The cytosolic fraction (20 µg) was subjected to a nonradioactive PKA activity assay (Enzo Life Sciences) according to manufacturer instructions. Protein determination was performed by a DC Protein Assay (BIO-RAD), using BSA as a standard.

**Immunochemistry**

**Monocultures.** Cells were cultured in Fn-coated eight-well chamber slides (Nalge Nunc) at a density of 25,000 cells/well, fixed 7 d after initiating differentiation, and stained for MBP (1:250; catalog #MCA4095, BIO-RAD/A bd Serotec; RRID: AB_325004) as described previously (Baron et al., 2014). The cells were analyzed with a conventional fluorescence microscope (ProVis AX70, Olympus). In each experiment, at least 500 cells were scored as MBP-positive or MBP-negative (“differentiation”). In addition, cells bearing MBP-positive membranous structures spread between the cellular processes were identified as myelin membrane forming (“myelination”).

**Cocultures.** Cocultures were fixed and costained for MBP (myelin) and NF-H (axon; 1:5000; Cat# CPCA-NF-H RRID: AB_2149761, EnCor Bio- technology), as described previously (Stancic et al., 2012), and analyzed by confocal laser scan microscopy (TCS SP2 or SP8 AOBs Microscope, Leica Microsystems) using Leica Confocal Software. The percentage of myelinated axons was calculated using MATLAB software programmed to recognize only linear structures [myelin and axons; i.e., excluding oligodendrocyte (OLG) cell bodies]. In each experiment, three images at 20× magnifications per coverslip and two to three coverslips per condition were analyzed.

**Immunohistochemistry**

**Cerebellar slice cultures.** Cerebellar slices were fixe d for 15 min in 4% paraformaldehyde and blocked for 3 h in block solution (1 mmol/L HEPES, 2% heat-inactivated horse serum, 10% heat-inactivated normal goat serum, 1% BSA, and 0.25% Triton X-100 in PBS). Primary antibodies (anti-MBP and anti-NF-H) were applied for 4 h at 4°C. Slices were subsequently incubated with appropriate Alexa Fluor-conjugated secondary antibodies (1:500) overnight at 4°C. Nuclei were stained with DAPI. Analyses were performed as described for cocultures (see Immunocytochemistry).

**Brain tissue.** Sections were blocked and permeabilized with PBS containing 5% normal donkey serum (Olig2) or 5% horse serum (Ki67, PDGF receptor (PDGFRα)) and 0.3% Triton X-100. Sections were incubated with anti-Olig2 (1:1000; catalog #AB9610, Millipore; RRID: AB_10140147) or a mixture of anti-Ki67 (1:800; catalog #PA5-19462, Thermo Fisher Scientific; RRID: AB_10981523) and anti-PDGFRα (1:2000; catalog #AF1062, R&D Systems; RRID: AB_2238987) diluted in PBS containing 3% serum overnight at 4°C, followed by a 2 h incubation with appropriate Alexa Fluor-conjugated secondary antibodies. Nuclei were visualized by DAPI staining. Myelin was stained with 0.1% (in 70% ethanol) Sudan Black solution, which did not interfere with immunofluorescence. Sections were analyzed with a confocal laser scan microscope (SP8 AOBs, Leica). The numbers of Olig2-positive cells of DAPI-stained cells, and the numbers of Ki67-positive/PDGFRα-negative, Ki67-positive/PDGFRα-positive, and Ki67-negative/PDGFRα-positive cells at the base of the injection site were manually counted.

**In situ hybridization**

The generation and the detection of the DIG-labeled proteolipid protein (PLP) probe are described (Chari et al., 2006; Stoffels et al., 2013). PLP
mRNA reactivity was assessed in the corpus callosum directly at the base of the injection site. Scores were related to PLP positivity in the corpus callosum well outside the lesion area. At minimum, 100 cells were scored in each assessed area.

Western blot analysis
Equal amounts of protein were subjected to Western blot analysis as described previously (Bislisbi et al., 2012). The antibodies used were anti-CREB-S133 (1:1000; catalog #9198 also #9198S and #9198L, Cell Signaling Technology; RRID: AB_2561044), anti-Fn (1:500; catalog #AB2033, Millipore; RRID: AB_2105702), anti-MBP (1:100; catalog #MCA4095, BIO-RAD/AbD Serotec; RRID: AB_325004), anti-actin (1:500; catalog #A-5441, Sigma-Aldrich; RRID: AB_476744), and appropriate IRdye-conjugated secondary antibodies (1:5000; LI-COR Biosciences). The signals were detected using the Odyssey Infrared Imaging System (LI-COR Biosciences) and analyzed using Scion image software.

Experimental design and statistical analysis
Details of the experimental design are described in the Materials and Methods section, the Results section, and the figure legends along with the individual statistical analyses and the number of independent experiments or animals (n). For cell and slice culture experiments, independent experiments were performed with newborn rats of different litters. Data are presented as the mean ± SD. Statistical analysis was performed using a one-sample t test when compared relative to vehicle-treated or control cells or slices (set to 100% in each independent experiment). When relative values of more than two means were compared, statistical significance was calculated by a one-way ANOVA followed by a Dunnett’s post-test. When absolute values of more than two means were compared, statistical significance was calculated by a one-way ANOVA followed by a Newman–Keuls post-test. A p value of <0.05 was considered significant. Statistical analysis was performed using GraphPad Prism software.

Results
Ganglioside GD1a overcomes inhibition of myelin membrane formation by fibronectin
To assess whether gangliosides may act as Fn antagonists by precluding their inhibitory effect on myelination, we first examined the effect of various gangliosides on OPC maturation in primary OLG cultures, which were grown on Fn or inert PLL. To this end, gangliosides were added to immature OLGs (Fig. 1A) and the percentages of MBP-positive cells and those that elaborated myelin membranes, reflecting OLG differentiation and myelin membrane formation respectively, were determined. Consistent with previous work by us and others (Buttery and Silence-Constant, 1999; Maier et al., 2005; Sisková et al., 2006, 2009; Lafrenaye and Fuss, 2010; Stoffels et al., 2013), OLGs plated on Fn are morphologically immature when compared with OLGs plated on PLL (Fig. 1B). Thus, an ~60% reduction was observed in the number of MBP-positive OLGs that form myelin membranes on Fn compared with PLL (38.3 ± 5.3% on PLL vs 15.1 ± 2.3% on Fn), while the number of MBP-positive cells was similar on either substrate (40.7 ± 5.7% on PLL vs 38.4 ± 5.6% on Fn). Intriguingly, while the extent of differentiation of OPCs remained similar (Fig. 1C), treatment of immature OLGs with 10 μM GD1a essentially reversed the Fn-induced inhibitory effect on myelin formation (Fig. 1D, black bars; p = 0.049, n = 3). In fact, upon GD1a treatment, the number of OLGs forming MBP-positive myelin membranes was very similar to that obtained on PLL (38.3 ± 5.3% on PLL vs 36.2 ± 6.8% on Fn plus GD1a). Although a nonsignificant enhancement of myelin membrane formation on Fn was also seen in the presence of GD3, other investigated gangliosides were without any effect, emphasizing the selectivity of the stimulatory effect of GD1a. Moreover, the effect of GD1a appears to be specific in conjunction with the presence of Fn, since no apparent effect on myelin formation was seen in cells grown on PLL (Fig. 1C, white bars) or another ECM compound, laminin-2 (data not shown). Notably, the exposure of immature OLGs to GD1b and GT1b slightly reduced the number of myelin membranes on PLL (Fig. 1D; GD1b, p = 0.049; GT1b, p = 0.013; n = 3). Importantly, GD1a, but not its precursor GM1, similarly enhanced myelin membrane formation when the cells were plated on aggregated Fn (Fig. 1E; nearly significant; p = 0.068; n = 4). The treatment of immature OLGs with 10 μM gangliosides was not toxic, as no significant release of cytosolic LDH was observed (Fig. 1F, white bars). Consistently, cell viability, as monitored with an MTT assay, was not affected either (Fig. 1F, black bars).

To reveal whether GD1a affects other cell behavioral aspects relevant for OPC recruitment to the demyelinated areas, OPCs were subjected to a BrdU incorporation (“proliferation”) and transwell migration assay. As shown in Figure 2, GD1a exposure enhanced OPC proliferation on Fn (p = 0.033, n = 3) whereas GM1 decreased it (p = 0.008, n = 3), while both gangliosides decreased OPC migration (GM1, p = 0.033; GD1a, p = 0.009; n = 3). In contrast, in the case of cells plated on PLL, neither GM1 nor GD1a hardly affected OPC proliferation compared with the control, while GD1a, but not GM1, slightly enhanced rather than decreased OPC migration. Hence, exogenous addition of ganglioside GD1a to immature OLGs overcomes the inhibition of myelin membrane formation by Fn in OLG cultures, while at the same conditions GD1a treatment of OPCs increases proliferation and decreases migration. This effect appears to be dependent on the presence of Fn, because on PLL GD1a hardly affected either myelin membrane formation or proliferation, while OPC migration appeared to be slightly increased rather than decreased.

Ganglioside GD1a overcomes inhibition of (re)myelination by aggregated fibronectin
To mimic more closely the myelinating conditions, we next examined whether GD1a also overcomes the inhibition of myelination by Fn aggregates in DRG–OLG cocultures. The addition of aggregated Fn inhibited myelination of the DRGNs (Fig. 3A,B; p = 0.0001, n = 3), whereas the percentage of myelinated axons increased upon exogenous addition of GD1a, but not GM1. In the absence of aggregated Fn, exposure to either ganglioside was without any effect. To further explore the effect of GD1a in a more in vivo-like environment, we next examined the potential effect of GD1a on Fn-inhibited remyelination in demyelinated organotypic cerebellar slice cultures. As shown in Figure 3C and by others (Miron et al., 2010), at defined conditions lyssolecithin-induced demyelination can be triggered in these cultures (p = 0.036, n = 3), which is eventually followed by remyelination. Exposure to GD1a upon lyssolecithin-induced demyelination, slightly enhanced remyelination both at 14 and 21 d after lyssolecithin treatment (Fig. 3D). This is likely due to the fact that Fn expression upon lyssolecithin-induced demyelination is only transient (Stoffels et al., 2013), and thus remyelination is also robust in untreated cultures (Miron et al., 2010). Therefore, to expose the cells to the more persistent remyelination-inhibiting Fn aggregates, the demyelinated cerebellar slices were treated with the TLR3 agonist poly(I:C). Transient exposure to poly(I:C) induces astrocyte-mediates aggregation of Fn after lyssolecithin-induced demyelination and, as the Fn aggregates persist in this case (Fig. 3E), remyelination is blocked (Sikkema AH, unpublished observation; Fig. 3E–H; **p = 0.0002; n = 3). Of note, when continuously and subcutaneously applied, poly(I:C) stimulates myelin gene expression and OPC recruitment upon
lysolecithin-induced demyelination in vivo (Natarajan et al., 2016). It is likely that the amount of subcutaneously applied poly(I:C) reaching the lesion may not have sufficed to induce Fn aggregation, thus allowing remyelination to proceed. Following the addition of GD1a under Fn-aggregating conditions, the percentage of myelinated axons, a readout for remyelination, was significantly increased compared with vehicle (ethanol, ctrl) or with the indicated gangliosides (10 μM) and analyzed 4 d after treatment. Representative images of MBP immunocytochemistry of OLGs cultured on PLL and Fn 4 d after treatment are shown in B. Scale bar, 20 μm. The number of MBP-positive cells (differentiation, C, E), the number of MBP-positive cells bearing myelin membranes (myelination; D, E), the effects on cell toxicity (LDH, white bars; F), and cell viability (MTT, black bars; F) were assessed 4 d after treatment. Each bar represents the mean ± SD of three to four independent experiments. In each experiment, the data of vehicle-treated cells cultured on the respective substrate were set at 100% (horizontal line). The percentage of MBP-positive cells in vehicle-treated cells cultured on PLL, Fn, and aggregated Fn was 40.7 ± 5.7% (C), 38.4 ± 5.6% (C), and 39.3 ± 16.5% (E), respectively. The percentage of MBP-positive cells bearing myelin membranes in vehicle-treated cells cultured on PLL, Fn, and aggregated Fn was 38.3 ± 5.3% (D), 15.1 ± 2.3% (D), and 29.1 ± 7.2% (E), respectively. Statistical differences with vehicle-treated cells cultured on their respective substrate, as assessed with a one-sample t test, are indicated (*p < 0.05, n = 3). Note that ganglioside GD1a overcomes Fn-mediated inhibition of myelin membrane formation, while exposure to gangliosides has no effect on cell toxicity and viability.

Ganglioside GD1a stimulates OPC proliferation and overcomes inhibition of early OPC differentiation by aggregated fibronectin in vivo

To verify whether the stimulatory effect of GD1a on OPC maturation in the in vitro model systems also applies to in vivo conditions, the effect of GD1a on OPC proliferation and differentiation was investigated in the cuprizone animal model of toxin-induced demyelination–remyelination. Importantly, in contrast to events in OLG monocultures, in vivo Fn aggregates block early OPC differentiation in demyelinated white matter, as evidenced by decreased PLP mRNA levels (Stoffels et al., 2013). The cuprizone model is a highly reproducible toxin-induced animal model of demyelination in the CNS, particularly at the corpus callosum (Gudi et al., 2014). The copper chelator cuprizone induces selective OLG cell death, which results in a maximal demyelination after 5 weeks. Subsequent cuprizone withdrawal leads to recruit-
ment of resident OPCs and almost complete remyelination within 2 weeks (Gudi et al., 2014). Of interest, Fn is transiently upregulated upon cuprizone-mediated demyelination but is not aggregated (Hibbits et al., 2012; Espitia Pinzon et al., 2017). Therefore, aggregated Fn was injected into the demyelinated corpus callosum to create a local MS lesion-like environment (Fig. 4A). It should be noted that in this in vivo model, in contrast to the organotypic cerebellar slice cultures and MS lesions, the inhibitory effect of the injected Fn aggregates on OPC maturation can be partly overcome in time because of, among other reasons, aggregate clearance. Hence, since a transient Fn aggregate microenvironment is created, only the effect on early OPC differentiation, but not remyelination, can be examined. In this model, vehicle, GD1a, or GM1 was intraperitoneally injected at days 3 and 4 after Fn aggregate injection (Fig. 4A). Analyses at day 5 revealed that the number of Olig2-positive cells, representing OLG lineage cells, were increased more than twofold upon GD1a treatment (Fig. 4B,C; \( p = 0.019, n = 6 \)), but were slightly reduced (−20%) in GM1-treated lesions compared with vehicle-treated controls. To assess whether the enhanced Olig2-positive cell numbers were a reflection of increased proliferation, we analyzed the number of proliferating cells using a double-labeling approach with PDGF-Rα, an OPC marker, and Ki67, a marker for proliferation (Gerdes et al., 1983). GD1a-treated lesions showed a significant increase in the number of proliferating (i.e., Ki67-positive, PDGF-Rα-positive cells; Fig. 4D,E; \( p < 0.0001, n = 6 \)). These findings indicate that the increase in Olig2-positive cells is, at least partly, likely to be the result of an increased proliferation. Also, the number of PDGF-Rα-positive cells was increased in GD1a-treated lesions compared with vehicle- and GM1-treated lesions (Fig. 4D,E; \( p = 0.004, n = 6 \)). Analysis of OPC differentiation by in situ hybridization for PLP mRNA, the major myelin protein, showed a remarkable three times increased expression of PLP mRNA-positive cells at the lesion site upon GD1a treatment (Fig. 4D,E; \( p = 0.0001; n \geq 5 \)). In contrast, PLP mRNA-positive cells in both vehicle- and GM1-treated lesions were considerably less prominent. These data thus indicate that GD1a, but not GM1, stimulates OPC proliferation and concomitantly overcomes inhibition of early OPC differentiation in a Fn aggregate-containing cuprizone-demyelinated corpus callosum.

**GD1a becomes effective in overcoming fibronectin-mediated inhibition of myelin membrane formation at a relatively late stage of OLG maturation**

To clarify the underlying mechanism of GD1a, we first assessed whether GD1a may interfere with the binding between Fn and its cell surface receptors (integrins) on OLGs. Preincubation of Fn with GD1a reduced the adhesion of OPCs to Fn, but not PLL, by 50% (Fig. 5A; \( p = 0.034, n = 3 \)), while the preincubation of OPCs with GD1a reduced the adhesion of the cells on both Fn (Fig. 5B; \( p = 0.010, n = 3 \)) and PLL (Fig. 5B; \( p = 0.004, n = 3 \)). However, the effect of GD1a appeared to be nonspecific, since GM1 blocked the adhesion of OPCs to Fn in a very similar way (Fig. 5A, \( p = 0.034, B, p = 0.003; n = 3 \)). Accordingly, given the nonspecificity, these data clearly indicate that interference between Fn and its receptors likely does not contribute to the ability of GD1a to overcome OPC maturation in a Fn environment. As an alternative, GD1a may activate intracellular signaling pathways. Hence, we examined whether such an effect of GD1a might depend on the developmental stage of the OLGs (Fig. 5C). Yet, GD1a is not
Figure 3. Exogenous addition of GD1a overcomes inhibition of (re)myelination by aggregated DRGN-OLG cocultures and in organotypic cerebellar slice cultures. 

A, B, OPCs were added to in vitro DRGNs in the absence or presence of aggregated Fn (aFn). DRGN-OLG cocultures were treated 1 h after plating OPCs with vehicle (veh; ethanol, ctrl) in the absence of aFn, or the indicated gangliosides (2 μM) for 14 d. Myelin was visualized with MBP (green) and DRGNs with NF-H (red). Representative images at 14 d in coculture of three independent experiments are shown in A, and quantification in B. Scale bar, 50 μm. The percentages of myelinated axons in vehicle-treated cocultures in the absence of aFn (ctrl) was set at 100% (horizontal line). The percentage of myelinated axons in vehicle-treated cells was 21.1 ± 10.6%. Data are represented as the mean ± SD, and statistical differences with vehicle-treated cocultures, as assessed with a one sample t test, are indicated (**p < 0.01; ***p < 0.001, n = 3). Note that ganglioside GD1a overcomes aggregated Fn-mediated inhibition of myelination.

C, D, Organotypic cerebellar slice cultures obtained from newborn rats were cultured for 3 weeks (wks) to allow for myelination. Slices were left untreated (ctrl) or treated with lysolecithin for 17 h to induce myelination (day 0). In D, at day 2, demyelinated organotypic cerebellar slice cultures were treated with vehicle (ethanol, ctrl) or with 10 μM GD1a for 12 or 19 d (n = 2). The percentage of myelinated axons was assessed 5 d (C), 14 d (D), or 21 d (C, D) after lysolecithin treatment. Data are represented as the mean ± SD, and for C, statistical differences with control cultures at day 5 (C; one-way ANOVA, Newman–Keuls post-test, *p < 0.05, n = 3) are shown. E–H, A schematic representation of the treatments is indicated in F. Organotypic cerebellar slice cultures obtained from newborn rats were cultured for 3 weeks (wks) to allow for myelination. At day 0, demyelination was induced by lysolecithin. At day 2, demyelinated slices were untreated (—) or treated with the TLR3 agonist poly(I:C) to induce Fn aggregation (E). Western blot analysis of Fn aggregates (aFn) at day 21, MBP is a mature OLG marker, actin is used as a loading control. Upon Fn aggregation at day 5, slices were treated with vehicle (ethanol, veh) or the indicated gangliosides (10 μM) for 16 d. Myelin was visualized with immunostaining for MBP (green) and neurons with NF-H (red) 21 d after initiating demyelination. Representative images of three independent experiments are shown in G; quantification is shown in H. Scale bar, 50 μm. The percentages of myelinated axons in control nondemyelinated cultures at day 5 were set in each independent experiment at 100% (horizontal line). The mean percentage of myelinated axons at day 5 was 19.5 ± 5.4%. Data are represented as the mean ± SD, and statistical differences with control nondemyelinated cultures at day 5 (one sample t test, ***p < 0.001, n = 3) and vehicle-treated aFn-containing demyelinated cultures at day 21 (one-way ANOVA, Dunnett’s post-test, **p < 0.01, n = 3). Note that ganglioside GD1a overcomes aFn-mediated inhibition of remyelination in organotypic cerebellar slice cultures.
effective when OPCs or immature OLGs, which are grown on Fn, are transiently exposed to the ganglioside (Fig. 5E; OPCs, first (F) 72 h (F72h); immature OLGs, F24h, F48h), indicating that the effects of GD1a on proliferation and maturation are likely separate events. Indeed, a 24 h exposure to GD1a of the postmitotic MBP-expressing OLGs, just before maturation, suffices to overcome the myelination-inhibiting effect of Fn in vitro [Fig. 5E; last (L) 24 h (L24h), p = 0.0005, n = 7]. OLG differentiation was unaffected at all conditions (Fig. 5D). Notably, exposure at the L24h resulted in the same effect as that obtained after a 96 h exposure at an earlier differentiation stage (Fig. 5E, 96 h; p = 0.0001, n = 7), suggesting a prolonged effectivity and implying that GD1a is likely not (rapidly) degraded, which is consistent with the known long turnover rates of gangliosides (days to weeks).

Figure 4. Exogenous addition of GD1a overcomes the inhibition of early OPC differentiation by aggregated fibronectin in vivo. A, Schematic representation of the treatments as indicated in B–G. Animals were fed with cuprizone for 5 weeks (wks) to allow for demyelination. At day 0, Fn aggregates (aFn; 1.5 μg in 3 μl) were injected into the demyelinated corpus callosum, followed by treatment with vehicle (veh, ethanol) and the indicated gangliosides (10 μM in 3 μl) at day 2 and day 3. B–G, 5 d after aFn injection, the density of Olig2-positive (“oligodendrocyte lineage”; B, C), the amount of Ki67-positive and PDGFαR-positive (“OPC proliferation”; D, E), and PLP mRNA-positive (“differentiated oligodendrocytes”; F, G) cells were determined by immunohistochemistry (B–E) or in situ hybridization (F, G). Images in B, D, and F (in F the place of cannula is indicated with the dashed line) are representative images in the core of the Fn aggregate–injected demyelinated areas in the corpus callosum of five to six different animals. Scale bars: B, D, 10 μm; F, 100 μm. Data in C, E, and G are represented as the mean ± SD, and statistical differences with vehicle-treated lesions, as assessed with a one-way ANOVA are indicated (Newman–Keuls post-test, *p < 0.05; n ≥ 5). Note the increase in Olig2-positive, Ki67-positive/PDGFαR-positive and PLP mRNA-positive cells after treatment with ganglioside GD1a, but not ganglioside GM1, in aFn-injected demyelinated areas of the corpus callosum.
5–15% increase in the level of GM1 (Fig. 5F; \( p = 0.019, n = 3 \)), which is ineffective in reversing Fn inhibition (Figs. 1, 2, 3). To assess whether and which signaling pathways is/are potentially the target for GD1a in relieving the Fn myelination block (i.e., of relevance to OPC maturation), we performed an STK array.

GD1a overcomes inhibition of myelin membrane formation by fibronectin via activation of a PKA–signaling pathway

STK activity profiles in OLGs, induced by GD1a treatment, were generated with the STK PamChip peptide microarray system (Eriksson et al., 2014; Shi et al., 2016). The array contains 144 peptides, corresponding to activity-modulating phosphorylation sites of key signal transduction mediators, thus generating a network view on signal transduction activity. Ninety of the 144 peptides showed either an increase or a decrease of phosphorylation upon a 20 min exposure to GD1a (Fig. 6A). Peptides that differed significantly in the level of phosphorylation in vehicle-treated versus GD1a-treated OLGs are depicted in Table 1, and in Table 2, the annotated upstream kinases, which potentially are responsible for the altered phosphorylation levels, are shown. Upon a 20 min exposure to GD1a, a set of 10 of 144 peptides wassignificantly phosphorylated, of which most (i.e., four peptide sequences) are annotated to the PKA signaling pathway (Fig. 6A, *).

Figure 5. GD1a becomes effective in overcoming fibronectin-mediated inhibition of myelin membrane formation at a relatively late stage of OLG maturation. A, OPCs were plated on plasma Fn- or PLL-coated 96-well plates that were pretreated with vehicle (ethanol, ctrl) or the indicated gangliosides (10 \( \mu M \)). OPCs were allowed to adhere for 1 h, after which the relative percentage of adherent cells was determined using a colorimetric assay. The horizontal line represents adherence to vehicle-treated substrates, which was set at 100% in each independent experiment. The percentage adherence of OPCs on vehicle-treated PLL and Fn was 66.5 ± 7.3% and 50.8 ± 4.4%, respectively. Statistical differences with OPCs cultured on the respective vehicle-treated substrate, as assessed with a one-sample t test, are indicated (Fn, \( * \ p < 0.05, n = 3 \)). Note that GD1a and GM1 pretreatment of Fn, but not of PLL, decreases the adhesion of compared with adhesion on vehicle-treated substrate. B, Freshly isolated OPCs were pretreated for 30 min with vehicle (ethanol, ctrl) or the indicated gangliosides (10 \( \mu M \)), plated on Fn- or PLL-coated 96-well plates, and subjected to an adhesion assay (A, \( * \ p < 0.05, ** \ p < 0.01, n = 3 \)). C, Schematic representation of the treatments indicated in D and E. Upon differentiation of OPCs cultured on Fn, cells were continuously or transiently exposed to GD1a at the indicated differentiation stage and duration. MBP immunocytochemistry was performed at the mature OLG stage for all conditions. h = h.

D, E, The number of MBP-positive cells (differentiation, D) and the number of MBP-positive cells bearing myelin membranes (“myelination,” E) were assessed. Each bar represents the mean ± SD of three independent experiments. In each experiment, the data of vehicle-treated cells was set at 100% (horizontal line). The mean percentages of MBP-positive cells and MBP-positive cells bearing myelin membranes in vehicle-treated cells were 44.6 ± 6.9% and 20.2 ± 6.5%, respectively. Statistical differences with vehicle-treated cells cultured as assessed with a one-sample t test, are indicated (\( * \ p < 0.05, ** \ p < 0.01, *** \ p < 0.001, n = 3 \)). Note that ganglioside GD1a overcomes Fn-mediated inhibition when present at late differentiation. F, Immature OLGs and OLGs cultured on Fn at 6 d after initiating differentiation were treated with \( [3H] \)-GD1a for 96 and 24 h, respectively. Total gangliosides were extracted and separated by TLC, and the percentage of total activity versus GD1a-treated OLGs are depicted in Table 1, and in Table 2, the annotated upstream kinases, which potentially are responsible for the altered phosphorylation levels, are shown. Upon a 20 min exposure to GD1a, a set of 10 of 144 peptides was significantly phosphorylated, of which most (i.e., four peptide sequences) are annotated to the PKA signaling pathway (Fig. 6A, *).

Given that PKA has been implicated in the regulation of OPC differentiation (Raible and McMorris, 1990, 1993; Shiga et al., 2005; Syed et al., 2013), we next analyzed whether GD1a can overcome the inhibition of OPC maturation on Fn via a PKA–dependent mechanism. A PKA activity assay revealed that GD1a,
Figure 6. Exogenous addition of GD1a overcomes inhibition of myelin membrane formation by fibronectin via activation of a PKA-dependent pathway. A, OLGs cultured on Fn at 6 d after initiating differentiation were treated with ganglioside GD1a (10 μM) for 10 min, 20 min, or 16 h and subjected to a STK PamChip array. Clustering of normalized peptide phosphorylation signals. The phosphorylation signals were averaged (n = 4 for all conditions except 16 h (n = 2)). The heatmap shows the 90 of 144 peptides that were phosphorylated in at least one of the conditions compared with baseline phosphorylation (T = 0). Red and blue spots represent increased and decreased phosphorylation compared with the average phosphorylation. The peptide labels refer to the origin (protein and sequence) of the peptide sequences that are spotted on the PamChip array. A red asterisk (*) indicates peptides that are annotated to the upstream kinase PKA and that showed significant increased phosphorylation upon 20 min GD1a treatment compared with vehicle treatment. Other peptides that were significantly different phosphorylated upon GD1a treatment of OLGs (6 d in differentiation) compared with vehicle treatment are depicted in Table 1.

B, OLGs cultured on Fn at 6 d after initiating differentiation were treated with gangliosides GD1a or GM1 (10 μM) for the indicated time points and subjected to PKA activation assay. Each bar represents the mean ± SD of three to five independent experiments. In each experiment, the data of T = 0 min was set at 100% (horizontal line). Statistical differences with T = 0 min as assessed with a one-sample t-test are indicated (*p < 0.05, n ≥ 3). Note that GD1a activated PKA upon 20–30 min exposure, while PKA activity decreased to basal levels after 60 min.

C, D, OLGs cultured on Fn at 6 d after initiating differentiation were treated with ganglioside GD1a (10 μM) for the indicated time points and subjected to Western blot analysis (20 μg) using the CREB-S133 antibody. The CREB-S133 levels were quantified relative to the expression of the housekeeping protein actin (D). Each bar represents the mean ± SD of four independent experiments. In each experiment, the data of T = 0 min was set at 100% (horizontal line). Statistical differences with T = 0 min as assessed with a one-sample t test, are indicated (nearly significant, 60 min, p = 0.053, n ≥ 3). E, F, OLGs cultured on Fn at 6 d after initiating differentiation were treated with vehicle (ethanol, ctrl), H89 (20 μM), dBcAMP (1 mM), and/or ganglioside GD1a (10 μM) for 24 h. H89 was added 1 h before GD1a. The number of MBP-positive cells (differentiation, E) and the number of MBP-positive cells bearing myelin-like membranes (myelination; F) were determined by MBP immunocytochemistry. Each bar represents the mean ± SD of three to four independent experiments. In each experiment, the data from vehicle-treated cells were set at 100% (horizontal line). The percentage of MBP-positive cells in vehicle-treated cells was 38.5 ± 4.8%, and the percentage of MBP-positive cells bearing myelin-like membranes was 17.9 ± 3.6%. Statistical differences with vehicle-treated cells cultured (F; one-sample t test, *p < 0.05, **p < 0.01, n ≥ 3) and GD1a-treated cells (F; one-way ANOVA, Dunnett’s post-test, ###p < 0.0001, n = 3) are indicated. Note that GD1a treatment of OLGs cultured on Fn activates PKA, phosphorylates CREB (S133) and that the PKA inhibitor H89 overcomes Fn-mediated inhibition of myelin membrane formation, while the activation of a PKA signaling pathway mimicked the effect of GD1a.
but not GM1, activated PKA following a 20 min exposure, while after 60 min the PKA activity had decreased to basal levels (Fig. 6B; 20 min, p = 0.015; n = 6; 30 min, p = 0.025; n = 5). To further explore the GD1a-mediated signaling via PKA, we next analyzed its effect on the phosphorylation of the cellular transcription factor cAMP response element-binding protein (CREB), which is a downstream target of PKA. GD1a increased the S133 phosphorylation of CREB after a 60–120 min exposure to OLGs (Fig. 6C; p = 0.053, n = 3). Furthermore, H89, a PKA inhibitor, counteracted the effect of GD1a (Fig. 6F, ###p < 0.0001; n = 3), while dBcAMP, a PKA activator, mimicked the effect of GD1a on myelin formation in the presence of Fn (Fig. 6F, *p = 0.020; n = 4). Differentiation remained unaltered (Fig. 6E). Together, these findings indicate that GD1a overcomes the inhibition of myelin membrane formation by Fn via a mechanism that relies on a PKA signaling pathway.

**Discussion**

When considering a remyelination-based therapy, both the cellular and molecular environment in MS lesions should be taken into account. Among others, perturbed degradation of ECM molecules contributes to a lack of OPC maturation in MS lesions (Back et al., 2005; Lau et al., 2012; Stoffels et al., 2013). Indeed, we previously demonstrated that aggregates of Fn are persistently present in MS lesions, which frustrates the differentiation of OPCs in demyelinated lesions and thereby impedes remyelination (Stoffels et al., 2013). To trigger OPC maturation and subsequent remyelination within the lesion, a minimal therapeutic requirement appears to be either a lesional clearance of the aggregates as such or a circumvention or suppression of their impeding effects on myelin assembly. Our present findings demonstrate that the exogenous addition of ganglioside GD1a overruled the devastating effect of the presence of Fn aggregates on OPC maturation. The underlying mechanism appears related to a specific GD1a-dependent activation of a PKA signaling pathway. Thus, exogenously added GD1a, but not other gangliosides tested, antagonizes the inhibitory effect of (aggregated) Fn on myelin membrane formation in primary OLG monocultures and DRGN–OLG cocultures and remyelination in demyelinated cerebral slice cultures. The significance of the in vitro data was corroborated in the cuprizone demyelination–remyelination model. In this model, we managed to mimic a transient local Fn aggregate-containing lesion in the demyelinated corpus callosum that allows study of the effect of GD1a on OPC proliferation and early differentiation. Injection of GD1a into this Fn aggregate-containing lesion increased the number of OLG lineage cells, which reflects an increased OPC proliferation and, based on our observations obtained in OLG monocultures, presumably a decreased migration rather than an increased survival. In addition, the number of PLP mRNA-containing cells in GD1a–treated lesions, but not GM1–treated lesions, which is consistent with an enhanced OPC differentiation. Evidently, the present data support the notion that targeted delivery of GD1a to Fn (aggregate)-containing MS lesions may act as a potential novel molecular and therapeutic tool to induce OPC proliferation and maturation and to overcome remyelination failure.

In the CNS, endogenous GD1a is prominently expressed in neurons (Schnaar and Lopez, 2009), while conflicting findings about its presence in OLG lineage cells have been reported (Satoh et al., 1996; Marconi et al., 2005; Silajdžić et al., 2009). Among others, gangliosides actively participate in the organization of membrane signaling microdomains (Sonnino and Prinetti, 2010), for example, by directing adhesion receptors to discrete areas for optimal interaction with the opposing membrane. Both spatial
and temporal formation of membrane microdomains play a key role in myelination (Baron et al., 2005; Aureli et al., 2015), while oligodendroglial membrane microdomains are perturbed by Fn (Maier et al., 2005; Baron et al., 2014). More specifically, in cells grown on an Fn substrate, the galactolipid sulfatide is excluded from membrane microdomains, which in turn precludes the propagation of myelination-promoting laminin-2 signaling (Baron et al., 2014). Moreover, in MS lesions, an altered membrane microdomain distribution of oligodendroglial N155 and sulfatide is observed, compared with the distribution of these components in rafts, and isolated from healthy white matter (Maier et al., 2005, 2007). It is therefore tempting to suggest that exogenously added GD1α modulates perturbed membrane microdomains, which were induced by Fn (aggregates) in OLGs. In fact, exogenously added gangliosides may alter the lateral membrane distribution (Simons et al., 1999), while exogenously added GD1α readily integrates into OLG membrane microdomains (Heffer-Lauc et al., 2003). The thus restored membrane microdomain integrity might re-establish the correct PKA-mediated signaling environment that is required for OPC maturation. Consistently on Fn PKA, activation mimics the (re)myelination stimulus of GD1α, while PKA inhibition blocks the effect of GD1α. Specifically, in OLGs cultured on Fn GD1α exposure activated PKA, causing phosphorylation of its downstream target CREB. Several lines of evidence indicate that the elevation of cAMP levels, the ensuing activation of PKA, and CREB phosphorylation are necessary events in regulating OLGA maturation and myelin protein expression in primary OLGs (Raible and McMorris, 1990, 1993; Sato-Bigbee and DeVries, 1996; Shiga et al., 2005). Furthermore, inhibitors of cAMP-hydrolyzing phosphodiesterases induce the expression of myelin-specific components (Medina-Rodriguez et al., 2013; Syed et al., 2013), while a regulatory control mechanism for PKA activity has been described for a timed progression of the OLG lineage (Baron et al., 1998; Simon et al., 2016). The current work thus leads to a scenario in which (1) Fn may hinder the activation of a PKA signaling pathway that is essential for OPC maturation, while (2) this impediment can be overcome by exogenous addition of GD1α. However, the coinvolvement of the other annotated serine/threonine kinases (Table 2), such as CaMKII (Waggener et al., 2013) or p70S6K (Baron et al., 2000; Michel et al., 2015) cannot be excluded. Strikingly, MAPK, a downstream target of PKA in OLGs and shown to be implicated in early OPC differentiation on inert substrates, among others, by the phosphorylation of CREB (Baron et al., 2000; Fragoso et al., 2007; Guardiola-Diaz et al., 2012; Syed et al., 2013), was not identified with the STK array as an immediate kinase, activated by GD1α on Fn. However, whether MAPK may be critical for the effect of GD1α on Fn remains to be determined. In addition, given the polarized nature of OLGs, it is of interest that PKA is also involved in the biogenesis of polarized membrane domains in epithelial cells; the activation of PKA facilitates exocytosis, reduces (apical) endocytosis, and modulates transcytosis (Wojtal et al., 2008). Given our previous studies showing that polarized transport in OLGs becomes severely impaired in cells cultured on Fn (Baron et al., 2014; Sisková et al., 2006), it seems reasonable to speculate that GD1α and the subsequent activation of PKA restore polarized transport and hence myelin biogenesis in OLGs cultured of Fn.

Without effective therapies in treating MS, it is appealing to consider GD1α as a novel tool in promoting OPC proliferation and maturation, and thereby remyelination, in MS lesions. GD1α acts in vitro during a late step in OLGA maturation to induce myelin membrane formation, but can be added independent of the developmental stage of OLG maturation. In addition, GD1α does not affect myelin formation on either inert PLL or laminin-2, suggesting that its effect is Fn dependent. These features may all bear therapeutic relevance to the treatment of MS. For example, in MS lesions, a significant fraction of OPCs remains quiescent (Lucchetti et al., 1999; Chang et al., 2002; Kuhlmann et al., 2008), among others due to the presence of Fn aggregates (Stoffels et al., 2013). Here, GD1α may act to expand OPCs and generate remyelinating OLGs. In fact, in postmortem MS lesions of patients with a relative short disease duration, mature OLGs that make contact with surrounding demyelinated axons are a prominent feature (Wolswijk, 2000; Chang et al., 2002; Kuhlmann et al., 2008). GD1α may act on these cells to boost OPC maturation. In chronic MS lesions, the presence of immature OLGs is limited while OPC proliferation is a rare event (Wolswijk, 1998; Kuhlmann et al., 2008). However, based on the present findings and the presence of Fn in chronic MS lesions, GD1α may enhance the proliferative capacity of OPCs in these lesions. In addition, since the presence of immature OLGs is increased in lesions with high OPC densities (Wolswijk, 2002), OLGA maturation may improve. Together, the robust effect of exogenous GD1α, evoked by the activation of a PKA signaling pathway, on myelin membrane formation in the presence of Fn, and proliferation, in conjunction with the presence of either OPCs and/or immature OLGs and Fn (aggregates) in MS lesions, points to a novel treatment option that may boost the proliferation and maturation of resident OPCs and thereby overcome remyelination failure in MS lesions.

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