Dendritic cells (DCs) release exosomes with different characteristics based on stimulus. Here, we showed that DC cultures stimulated with low-level IFNγ released exosomes (IFNγ-DC-Exos) that contained microRNA species that can increase baseline myelination, reduce oxidative stress, and improve remyelination following acute lyssolecithin-induced demyelination. Furthermore, nasally administered IFNγ-DC-Exos increased CNS myelination in vivo. IFNγ-DC-Exos were preferentially taken up by oligodendrocytes, suggesting that they directly impact oligodendrocytes to increase myelination. Thus, our results show great potential for use of these IFNγ-DC-Exos as a therapeutic to promote remyelination in multiple sclerosis and dysmyelinating syndromes.

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activate T cells (Viaud et al., 2011), we focus on miRNA contents. Previous studies performed in the lab have shown that exosomes found in the periphery can impact brain myelination, possibly via delivery of miR-219 (Pusic and Kraig, in press). Here, we found that IFNγ-stimulated DCs exported exosomes, from here on referred to as IFNγ-DC-Exos, that were able to increase myelination and oxidative tolerance from control levels in vitro. In addition, application of IFNγ-DC-Exos improved recovery from MS-like demyelination modeled in slice culture. In vivo work confirmed that nasally administered IFNγ-DC-Exos can effectively increase brain myelination. In vitro tracking assays revealed preferential uptake of IFNγ-DC-Exos by oligodendrocytes, and to a lesser extent, by microglia. This is an important first step in elucidating the mechanisms of exosome-mediated increase in myelin, and will aid in the development of these exosomes as a therapy for remyelination. This work has appeared in preliminary form (Pusic et al., 2013).

2. Material and methods

2.1. Animal use

Wistar rats were obtained from Charles River Laboratory (Wilmington, MA) and were used in accordance with the University of Chicago Animal Care and Use Committee. Untimed pregnant Wistar female rats were single-housed with Enviro-dri® paper bedding (Shepherd, Watertown, TN) and Nestlets (Ancare, Bellmore, NY) and pups (cuddled to ten at birth) were used for hippocampal slice cultures. Male Wistar rats (10–12 weeks old) were double-housed and used for bone marrow isolations. Sprague Dawley rats were obtained from Harlan Laboratories (Madison, WI) and used for oligodendrocyte progenitor isolations.

2.2. Isolation of dendritic cells

Immature bone marrow cells were isolated from Wistar rats, as previously described (Powell et al., 2003). Briefly, animals were anesthetized with progressive exposure to 100% carbon dioxide and then immediately decapitated. Using aseptic techniques, bone marrow was aspirated out of the femurs and tibias and stromal cells were purified through the passage of bone and debris through a strainer. Red blood cell lysis buffer consisting of 0.15 M NH₄Cl, 10 mM KHCO₃, and 0.1 mM EDTA (Sigma, St. Louis, MO) was used to remove the red blood cells. Cells were then washed and plated in 6-well plates (BD Falcon, San Jose, CA) at a density of 10⁶ cells/mL in RPMI 1640 (Invitrogen, Carlsbad, CA) containing 20 ng/mL of GM-CSF (Peprotech Inc., Rocky Hill, NJ) and used for oligodendrocyte progenitor isolations.

2.3. Generation of dendritic cell-derived exosomes and isolation

Day seven BMDCs were plated at a concentration of 10⁶ cells/mL and were either placed in media alone or stimulated with media containing 500 U of IFNγ (R&D Systems, Minneapolis, MN). Three days later culture media was removed and spun down to remove any cells and debris. Exosomes were then isolated using ExoQuick (SBI, Mountain View, CA). ExoQuick was added to culture media at a ratio of 1:5, incubated at 4 °C overnight, and exosomes were precipitated by centrifugation at 2000 × g for 30 min. The exosome pellet was then resuspended in 100 µL of sterile phosphate buffered saline at a pH of 7.3. Isolation of exosomes was confirmed via immunoblot for two exosomal protein markers, CD63 and Alix (AbD Serotec, Raleigh, NC) (Schroyer and Bhatnagar, 2008) and electron microscopy (Thery et al., 2006). Quantification of exosomes was performed by BCA assay (ThermoFisher Scientific, Waltham, MA) of protein content.

2.4. Slice culture preparation and use

Hippocampal slice cultures were used to examine grey matter demyelination in vitro. Hippocampal slice cultures allow for the treatment of brain tissue in a controlled environment and are advantageous in that they maintain the following: the neurovascular unit (Kovacs et al., 2011), multi-synaptic electrical activity (Kunkler and Kraig, 1997), pyramidal neuron vitality (Hulse et al., 2008), responsive astrocytes and microglia (Ransohoff and Perry, 2009; Grinberg et al., 2011), and neural immune signaling including the expression of cytokines (Kunkler et al., 2004) as seen in vivo.

Untimed pregnant Wistar female rats (Charles River, Wilmington, MA) were single-housed with Enviro-dri® paper bedding (Shepherd, Watertown, TN) and Nestlets (Ancare, Bellmore, NY). The addition of Nestlets provided enrichment for the pregnant rats thus leading to the improvement of slice culture health; from 80–85% viable (n = 104 litters) to 95–100% viable (n = 156 litters). For culturing, P9–P10 rat pups were used to make hippocampal slice cultures (350 µm thick) as previously described (Kunkler and Kraig, 1997; Mitchell et al., 2010). After 18 days in vitro (DIV) cultures were transferred to a serum-free media, which does not activate microglia and does not contain horse serum-derived exosomes, thus allowing for accurate assessment of the impact of exosome treatments (Pusic and Kraig, in press). Media composition is further described in the Supplementary materials. Cultures were used between 21 and 35 DIV, as this time frame is sufficient to allow measurements of neural tissue responses to experimentally induced disease states. All cultures were screened for viability via Sytox™ (Invitrogen) staining, a fluorescent maker of cell death prior to experimental use (Mitchell et al., 2010). Cultures with evidence of CA3 or CA1 pyramidal neuron loss were not used for experiments.

Exosome treatments were applied to the media of slice cultures and incubated for three days unless otherwise noted. Treatments consisted of 100 µg of exosomes in 50 µL of sterile phosphate buffered saline (PBS). All experimental measurements were compared to age-matched control slice culture levels.

For lyssolecithin-induced demyelination, mature (21–24 DIV) hippocampal slice cultures were incubated in serum-free media containing lyssolecithin (0.5 mg/mL) for 17 h (Birgbauer et al., 2004). Cultures were then rinsed three times in Neurobasal (Invitrogen), and returned to standard incubation conditions with or without exosome treatment (Pusic and Kraig, in press).

Oxidative stress [(OS), i.e., excess production of oxidants over that of anti-oxidants] was measured using CellRiX™ Deep Red Reagent (Invitrogen), a cell-permeant fluorogenic probe was used as previously described (Grinberg et al., 2012). Membrane preservation (8.6 µg/mL; Supelco Analytical, St. Louis, MO) was used to generate reactive oxygen species. Exosomes were applied to naïve slices for 3 h then co-incubated with CellRiX™ and membrane for 2 h. CellRiX™ fluorescence intensity was quantified via digital imaging strategies as described below. A standardized area of interest at the CA3 pyramidal neuron layer was used for all quantifications.

Regional and microglial glutathione levels were assessed using ThioTracker™ (Invitrogen) a fluorescent dye that reacts with reduced thiols in intact cells, which predominantly reflect glutathione (Mandavilli and Janes, 2010). Procedures for staining followed manufacturer’s protocol, modified for use with hippocampal slice cultures. ThioTracker™ was dissolved in dimethyl sulfoxide (2 mM, Sigma) and used at 20 µM in a thiol-free solution. Briefly, culture inserts were dipped into three separate 60 mm culture dishes containing 10 mL each of Gey’s balanced salt solution (Sigma) supplemented with 7.25 mL 45% p-glucose (Sigma) to remove extracellular thiols. Cultures were placed for 30 min in glucose supplemented Gey’s solution containing ThioTracker™ that had been pre-equilibrated to incubator conditions. Inserts were then washed again as described above and fixed in 10% phosphate buffered formalin at
4 °C overnight, stained with isoelectin GS-IB4 AlexaFluor® 594 conjugate (1:20; Invitrogen), mounted, and cover slipped.

Glutathione was quantified using a self-calibrating sensitive CCD digital imaging system consisting of a QuantEM-512SC camera (Photometrics, Tucson, AZ), electronic shutter (Lambda SC Smart Shutter; Sutter Instruments, Novato, CA), and a standard 100 W Hg light on a DMIIRE2 inverted microscope (Leica, Heidelberg, Germany) at 20 × gain. A standardized area of interest at the CA3 pyramidal neuron layer was used for all quantifications. Digital images were thresholded to a uniform range and the average optical intensity registered using MetaMorph software (ver. 7.5.4.0; Molecular Devices, Sunnyvale, CA). Confocal images of microglial Thioflavine™ staining were acquired using a Leica TCS S Phelps I A0BS laser scanning confocal microscope (University of Chicago Integrated Microscopy Core Facility).

2.5. Isolation and use of oligodendrocyte progenitor cells

Oligodendrocyte progenitor cells (OPCs) were isolated from Sprague Dawley rats (Harlan, Madison, WI) using the sequential immunopanning protocol previously described (Dugas et al., 2006). Briefly, cerebral cortices were dissected from P6 rats and enzymatically dissociated with papain at 37 °C and 5% O2 and 5% CO2, following enzymatic dissociation and gentle titration the single cell suspension was sequentially incubated on dishes coated with antibodies against Ran-2, galactocerebroside (GalC), and O4. Ran-2 and GalC removed contaminating astrocytes and mature oligodendrocytes respectively, while the O4 coated plate captured OPCs. Ran-2, GalC, O4" OPCs were then trypsinized from the O4 panning plate and seeded onto poly-o-lysine coated T150 flasks (Fisher Scientific, Hampton, NH) in 20 mL of proliferation media (defined below) at 2500 cells/mL.

Cells were cultured at 37 °C in 10% CO2 in high glucose DMEM (Invitrogen), supplemented with human transferrin (100 μg/mL), bovine serum albumin (100 μg/mL), putrescine (16 μg/mL), progesterone (60 ng/mL), sodium selenite (40 ng/mL), N-acetyl-l-cysteine (5 μg/mL), o-biotin (10 ng/mL), insulin (5 μg/mL), glutamine (2 mM), sodium pyruvate (1 mM), penicillin-streptomycin (100 U each), Trace Elements (1×; Mediatech, Herndon, VA), and the following growth factors for proliferation media: Forskolin (42 μg/mL), NT-3 (1 ng/mL), IGF-1 (10 μg/mL), Peprotech, Rocky Hill, NJ) and the OPC mitogens CNTF (10 ng/mL, Peprotech), and PDGF-AA (10 ng/mL, Peprotech). All reagents for OPC isolation were purchased from Sigma unless otherwise noted.

For differentiation experiments, OPCs were passaged from a T150 flask using 0.05% Trypsin-EDTA diluted 1:10 in Earle’s Basic Salt Solution and plated on poly-o-lysine coated 12 mm glass coverslips in 2 mL of proliferation media minus PDGF-AA at 2500 cells/mL. OPCs were allowed to recover overnight before treatment. For exosome treatment, IFNy-DC-Exos (50 μg) were added directly to media. Transfection with mirVana hsa-miR-219-5p mimic (20 nM, Invitrogen) was performed using Lipofectamine RNAiMAX transfection reagent (Invitrogen) as per the manufacturer’s protocol. Control cultures were maintained in proliferation media without PDGF-AA. Positive controls were supplemented with triiodothyronine (T3; 40 ng/mL) to stimulate differentiation. Three days post-treatment, cultures were collected and fixed in cold 4% paraformaldehyde and immunostained for O4 and O1 (hybridoma-derived antibodies provided by the Popko lab), with DAPI (Invitrogen) counterstaining. Percent of O4 and O1 positive cells were calculated as positive cell count/total cell count.

2.6. Immunoblotting

Samples were collected by dissecting out the CA3 region of the hippocampal slice cultures with a diamond knife (Fine Scientific, Foster City, CA) and protein was extracted according to standard protocol (Cook et al., 2011; Rountree et al., 2011). Isolated protein concentration was then measured via BCA assay and run on SDS-PAGE gels, transferred onto PVDF membranes (Bio-Rad, Hercules, CA), and probed with appropriate primary antibody. Antibodies used for probing included CD63, Aixa (AbD Serotec, Raleigh, NC) and myelin basic protein (MBP; Novus Biologicals, Littleton, CO), all at a concentration of 1:1000. For assessment of mir-219 target protein levels, PDGF-Rα (Santa Cruz) was used at 1:200, and ELOVL7 (Sigma) was used at 1:500. Secondary antibody, horseradish peroxidase conjugated anti-mouse or anti-rabbit (Sigma) was used at a concentration of 1:1000 and blots were then visualized via chemiluminescence. Analysis was performed on Quantity One® 1-D Analysis Software (Bio-Rad). All protein was normalized to β-actin (Sigma) with both primary and secondary antibody concentrations of 1:1000.

2.7. Immunohistochemistry and confocal microscopy

Hippocampal slice cultures were fixed with periodate-lysine-paraformaldehyde fixative or methanol for 30 min at 4 °C (depending on the cell surface stain) and stained for CNPase, Iba1, GFAP, and NeuN. Primary antibodies used in this study are: monoclonal mouse anti-CNPase IgG (1:1000, Millipore, Billerica, MA), polyclonal rabbit anti-Iba1 IgG (1:1000, Wako Pure Chemical Industries, Osaka, Japan), monoclonal mouse anti-GFAP IgG (1:1000, Santa Cruz Biotechnology, Santa Cruz), monoclonal mouse anti-NeuN IgG (1:1000, Millipore, Billerica, MA), polyclonal rabbit anti-Musashi-1 (Msi1) and Musashi-2 (Msi2) (1:50, Cell Signaling Technology, Danvers, MA), monoclonal mouse anti-NG2 (2×50, Life Technologies, Grand Island, NY), and Ki-67 (1:500, Novus). Slices were then incubated with AlexaFluor® 488 or AlexaFluor® 594 labeled secondary antibody specific to the appropriate animal species (1:1000, Millipore, Billerica, MA) and mounted using 1:1 glycerol/PBS (see below). Fluorescently stained slices were then visualized using a confocal laser microscope at the Light Microscopy Core Facility at the University of Chicago. All images were acquired at 100 × gain.

2.8. miRNA extraction and profiling

miRNAs were extracted from exosomes using the mirVana™ miRNA Isolation Kit (Life Technologies, Grand Island, NY) according to manufacturer’s instructions. Purity and miRNA yield were assessed by NanoDrop2000 (ThermoFisher Scientific). miRNA (100 ng) was then reverse transcribed into cDNA using the TaqMan® MicroRNA Reverse Transcriptase Kit (Life Technologies) and further amplified with Megaplex™ PreAmp Primers (Rodent Pool set) with TaqMan® PreAmp Master Mix Kit (Life Technologies). Samples were then run on TaqMan® Array Rodent MicroRNA A + B Cards Set v3.0 (Life Technologies) on the 7900HT Real-Time PCR System (Life Technologies). All procedures above were performed according to manufacturer’s instructions.

2.9. DC-derived exosome coupling to Quantum Dots and uptake by slice culture

DC-derived exosomes were coupled to Quantum Dots (QDs), which are 24 nm in size and fluoresce at 620 nm. QDs were first conjugated to monoclonal CD63 antibody (AbD Serotec, Raleigh, NC) according to manufacturer’s protocol (Ocean Nanotech, Springfield, AR). Briefly, using covalent coupling chemistry, carboxyl groups on the surface of the QDs (1 nmol) were activated and incubated for 2 h at room temperature with 0.05 mg of CD63. After incubation, the reaction was quenched by adding Ocean’s quenching buffer (Ocean Nanotech) and incubating for 15 min. Conjugated QD-CD63 were run on a 1.5% agarose gel to separate conjugated QD-CD63 from unconjugated QDs. The conjugated QD-CD63 band was excised out of the gel and placed into a GeBAlpha-tube (GeBa Gene Bio-Application, Yavne, Israel). GeBAlpha-tube was used according to manufacturer’s protocol to elute out QD-CD63 from the gel. Isolated QD-CD63 was then used to tag IFNy-DC-Exos by gently rocking for 2 h at room temperature, referred to from here as QD-Exos. QD-Exos were introduced to 21 DIV hippocampal slice cultures.
and incubated for 3 h, washed with PBS, fixed in methanol, and immuno-
nostained for specific cell types to visualize uptake/location of QD-Exos. We found that fixation with methanol preserved QD fluorescence for 
over two weeks compared to PLP, 4% paraformaldehyde, 10% buffered
formaldehyde which lead to deterioration of fluorescence in 
days. Mounting with Anti-fade (Invitrogen) also progressively reduced 
QD fluorescence intensity over days. Accordingly tissue sections tagged
with QDs were cleared with glycerol/PBS and cover slipped.

2.10. Intranasal administration of IFNγ-DC-Exos and assessment of 
myelin change

Wistar rats were nasally administered exosome preparations. Rats 
were placed in a fume hood with a heat lamp and thermo-regulator to 
maintain temperatures at 37 °C. Isoflurane (Baxter, Deerfield, IL) anes-
thesia was delivered via a nose cone (five percent induction and two-
three percent maintenance, delivered in oxygen). 50 μL of exosomes
(100 μg protein) were administered over a 20 minute period at a rate of 
5 μL every 2 min to alternating nostrils (Liu et al., 2001a,b). Shams 
animals were administered 50 μL of PBS alone, following the protocol 
above.

Three days later, animals were anesthetized with progressive expos-
ture to 100% carbon dioxide and decapitated. Brains were rapidly 
removed, flash frozen in isopentane, and stored at −80 °C until further 
use. Brains were sectioned (20 μm) using a cryostat (Leica), fixed in 
10% buffered formalin phosphate (ThermoFisher Scientific) for 15 min, 
and incubated with FluoroMyelin™ (1:300; Invitrogen) for 40 min to 
stain myelin (Dugas et al., 2010). FluoroMyelin™ intensity was quantified
as described above using computer-based digital imaging strategies to 
assess integrated optical density at defined areas of interest.

2.11. Electron microscopy

Hippocampal slice cultures were sectioned perpendicularly to the long 
axis of CA3 in order to capture Schaffer collateral axons and images 
were obtained through the Electron Microscopy Core Facility at the 
University of Chicago. Myelin thickness was measured via g ratio.
ImagePro software (v.4.1; Media Cybernetics, Silver Springs, MA) was 
used to measure inner axonal diameters and total outer diameters for 
g ratio calculation (inner axonal diameter/total outer diameter) (Guy 
et al., 1991). Exosomes were also imaged using the Electron Microscopy 
Core Facility.

2.12. Data handling and statistics

Data were analyzed using SigmaStat software (v.3.5; Systat Soft-
ware, Chicago, IL). All data were subject to normality testing (p-value to 
reject: 0.05), equal variance testing (p-value to reject: 0.05), and
power (1-β: >0.8). Mean control data in each experiment was scaled to
1.00 with all subsequent parameters scaled proportionally to better 
allow inter-experiment comparisons. Molecular biological data
(miRNA RT-PCR and Immunoblot of protein) analyses included two 
technical replicates per experimental (i.e., biological) measurement.
All experimental groups consisted of biological replicates of n ≥ 3.
Pairwise comparisons were made with the Student’s t-test and multi-
ple comparisons done via ANOVA plus post hoc Holm–Sidak testing.

3. Results

3.1. Confirmation of IFNγ-DC-Exo isolation

Isolation of exosomes released from IFNγ-stimulated DCs was first 
confirmed via immunoblot for CD63 (50 kDa) and Alix (96 kDa), two
well-known markers present on exosomes (Fig. 1A) (Schoery and
Bhatnagar, 2008). The presence of exosomes was further determined
by electron microscopy (EM), which revealed the presence of vesicles
with a diameter of around 50 nm (Fig. 1B). IFNγ-DC-Exos were estab-
lished to be non-toxic, as hippocampal slice cultures treated with IFNγ-
DC-Exos did not show any signs of cell death when stained with a fluores-
cent marker of cell death (Mitchell et al., 2010) at 3, 6, 9, and 12 days
post-application (Fig. 1C–D). Additionally, toxicity of UV treated IFNγ-
DC-Exos were also assessed, and likewise showed no signs of cell death in 
hippocampal slice cultures (data not shown). Thus, we proceeded
with subsequent experiments involving the use of hippocampal slice 
cultures.

3.2. IFNγ-DC-Exos increased compact myelin and myelin levels in 
slice cultures

Different exosome treatments were applied to hippocampal slice 
cultures and EM imaging was performed three days later to determine 
changes in compact myelin. EM images demonstrated intact and tightly 
laminated myelin whose thickness was increased with the application of
IFNγ-DC-Exos (Fig. 2A). Subsequent calculations of g ratios revealed
significant (p = 0.008) improvement of laminated myelin with IFNγ-
DC-Exo treatment compared to control (Fig. 2B).

When applied to naïve 24 DIV hippocampal slice cultures, IFNγ-DC-
Exos significantly (p = 0.02) increased production of myelin basic pro-
tein (MBP) as measured via immunoblot (Fig. 2C). UV–treatment of IFNγ-DC-Exos (545 nm, 45 min 100 μWatts/cm²) prior to application
abrogated this effect, indicating involvement of RNA species in the ob-
served increase in myelin production (Eldh et al., 2010). Additionally, a
significant (p < 0.001) decrease of MBP levels was seen with UV–
treatment of IFNγ-DC-Exos compared to control (Fig. 2C). This is likely
due to the delivery of exosomal proteins damaged through UV treat-
ment (Eldh et al., 2010).

3.3. IFNγ-DC-Exo treatment does not cause progenitor depletion

To determine whether exosome-mediated increase of OPC differen-
tiation has a deleterious effect on progenitor populations, the presence
of neural progenitor cells and OPCs was assessed in hippocampal slice
cultures treated with IFNγ-DC-Exos and unstimulated-DC-Exos com-
pared to untreated control. Staining with Musashi (Msi1/Msi2)
(Fig. 3A) for neural stem cells revealed no signif-
cant difference in the number of positive cells
among slice cultures. Administration of these exosomes 3 h prior to
microglia, as seen by isolectin-GS-IB4 double staining (Fig. 4C).
Glutathione was found local-
ized to microglia, as seen by isolectin-GS-IB4 double staining (Fig. 4C).

3.4. IFNγ-DC-Exos increased oxidative tolerance in hippocampal 
slice culture

IFNγ-DC-Exo treatment also significantly increased oxidative toler-
ance of slice cultures. Administration of these exosomes 3 h prior to
menadione exposure significantly (p < 0.001) reduced oxidative stress,
as seen by CellROX™ staining (Fig. 4A–B). ThioflorTracker™ staining
revealed significantly (p = 0.001) increased levels of reduced gluta-
thane in cultures treated with both types of exosomes compared to un-
treated controls (Fig. 4C–D). However, treatment with IFNγ-DC-Exos
triggered a significantly greater rise in reduced glutathione than that
seen with unstimulated-DC-Exos alone. Glutathione was found local-
ized to microglia, as seen by isolectin-GS-IB4 double staining (Fig. 4C).

3.5. Specific miRNA involved in oligodendrocyte differentiation and 
anti-inflammatory pathways are highly expressed in IFNγ-DC-Exos

Screening of exosomal miRNA revealed significant differences be-
tween the contents of IFNγ-DC-Exos and unstimulated-DC-Exos. IFNγ-
DC-Exos were enriched in a number of miRNA species known to be
involved in either oligodendrocyte differentiation or anti-inflammatory
responses.
IFNγ treatment of DC cells increased expression and packaging of miRNAs involved in oligodendrocyte differentiation and myelin production pathways, listed in Fig. 5A. Notably, miR-219 was very highly enriched in IFNγ-DC-Exos and undetectable in unstimulated-DC-Exos. Enriched anti-inflammatory response miRNAs are shown in Fig. 5B. miR-181a, miR-451, miR-532-5p, and miR-665 were especially highly enriched (>10 fold) in IFNγ-DC-Exos versus unstimulated-DC-Exos. Increased presence of these miRNA species indicates the possibility that

IFNγ treatment of DC cells produced non-toxic exosomes. Exosome isolation confirmed by (A) Immunoblot for surface markers CD63 and Alix, and by (B) electron microscopy. Scale bar, 50 nm. (C) Exosome application to slice cultures showed no toxic effects. Slices were stained with Sytox™ at 3, 6, 9, and 12 days post-treatment. NeuN immunostaining image (left) shows normal neuronal architecture. Dotted lines indicate pyramidal (large arc) and dentate gyrus (small arc) principal neuronal layers. Sytox™ positive image (center) shows a control with principal neuronal injury induced by 24 hour exposure to 20 μM N-methyl-D-aspartate. Sytox™ negative image (right) of exosome treated culture showed no injury. Images were inverted to enhance visualization. Scale bar, 250 μm. (D) Quantification of Sytox™ fluorescence intensity (n = 9 slices/group) confirmed no significant change (ANOVA plus post hoc Holm–Sidak testing).

Fig. 1. IFNγ-stimulated dendritic cells produced non-toxic exosomes. Exosome isolation confirmed by (A) Immunoblot for surface markers CD63 and Alix, and by (B) electron microscopy. Scale bar, 50 nm. (C) Exosome application to slice cultures showed no toxic effects. Slices were stained with Sytox™ at 3, 6, 9, and 12 days post-treatment. NeuN immunostaining image (left) shows normal neuronal architecture. Dotted lines indicate pyramidal (large arc) and dentate gyrus (small arc) principal neuronal layers. Sytox™ positive image (center) shows a control with principal neuronal injury induced by 24 hour exposure to 20 μM N-methyl-D-aspartate. Sytox™ negative image (right) of exosome treated culture showed no injury. Images were inverted to enhance visualization. Scale bar, 250 μm. (D) Quantification of Sytox™ fluorescence intensity (n = 9 slices/group) confirmed no significant change (ANOVA plus post hoc Holm–Sidak testing).

Fig. 2. IFNγ-stimulated-DC-Exos increased myelination in slice culture. (A) Representative electron microscopy images illustrate increased compact myelin in IFNγ-DC-Exo treated slice cultures. Treatments from left to right: control (no treatment); IFNγ-DC-Exo; UV-IFNγ-DC-Exo; and unstimulated-DC-Exo. Scale bar, 200 nm. (B) Quantification of myelin g ratios from electron microscopy images (n = 3 slices/group; and 10 cells/slice) showed a significant (*) , p = 0.008 increase in compact myelin thickness with IFNγ-DC-Exo treatment. (C) Immunoblot confirmation of myelin basic protein (MBP) showed bands at 18, 23, and 45 kDa representing three transcripts of MBP. A quantification of all three transcripts of MBP showed a significant (*, p = 0.02) increase in MBP levels in slice cultures treated with IFNγ-DC-Exos and a significant (#, p = <0.001; n = 9, 15, 11, 9 slices/group, respectively) decrease in MBP in slice cultures treated with UV-IFNγ-DC-Exo. Significance was determined by ANOVA plus post hoc Holm–Sidak testing and ANOVA testing.
IFN-DC-Exos may reduce inflammation and oxidative stress. Detailed tabulation of miRNA screening is listed in the Supplementary materials.

3.6. IFN-DC-Exos deliver functional miR-219, resulting in decreased protein levels from miR-219 target mRNAs

To confirm the functional effect of miR-219 from IFN-DC-Exos, exosomes were applied to slice cultures and immunoblots for known targets were performed. Specific targets were: PDGFRα, the receptor for a mitogen that promotes proliferation and inhibits differentiation, and ELOVL7, which regulates lipid metabolism and redox homeostasis (Dugas et al., 2010; Zhao et al., 2010). Three days after IFN-DC-Exo application, slice cultures contained significantly (* p < 0.001) lower levels of both PDGFRα and ELOVL7 compared to untreated control slice cultures (Fig. 6A and B).

3.7. IFN-DC-Exos restores myelin levels post lyssolecithin-induced demyelination

Lysolecithin was used as a means to induce demyelination, as a model of MS in hippocampal slice cultures (Birgbauer et al., 2004). Treatment with IFN-DC-Exos post lyssolecithin exposure significantly (* p < 0.001) increased remyelination toward normal levels, measured at five days post injury, compared to cultures treated with lyssolecithin alone or given UV–IFN-DC-Exos (Fig. 7).

3.8. Nasal administration of IFN-DC-Exos increased levels of myelin in the brain

To assess the ability of IFN-DC-Exos to increase myelin in vivo, IFN-DC-Exos were nasally administered to rats and their ability to...
increase myelin in vivo was determined. Three days post-nasal administration, brains were harvested and improved myelination was observed in the motor cortex by FluoroMyelin™ staining (Fig. 8A) and MBP immunostaining (Fig. 8B). FluoroMyelin™ staining intensity was significantly \((p < 0.001)\) higher in IFNγ-DC-Exo treated animals than PBS treated (sham) animals (Fig. 8C). Likewise, MBP immunostaining intensity was significantly \((p = 0.001)\) higher in IFNγ-DC-Exo treated animals than shams (Fig. 8C). Immunoblot analysis similarly showed significantly \((p = 0.019)\) increased MBP levels in the cortex of animals treated with IFNγ-DC-Exos compared to sham (Fig. 8C).

### 3.9. QD tagged IFNγ-DC-Exos are preferentially taken up by oligodendrocytes

To determine whether QD nanoparticles were successfully conjugated to anti-CD63 antibody, unconjugated QD nanoparticles and conjugated QD-CD63 were analyzed on a 1.5% agarose gel. Conjugated QD-CD63 (Fig. 9A, lane 1) migrated at a higher molecular weight in comparison to unconjugated QD (Fig. 9A, lane 2) indicating the successful conjugation of the coupling of QD-CD63, seen as a circular structure with an electron dense core (Fig. 9B, arrowhead), to exosomes (Fig. 9B, arrow) was visualized by EM imaging.

QD-IFNγ-DC-Exos (Fig. 10A) and QD-unstimulated-DC-Exos (Fig. 10B) were applied to hippocampal slice cultures and immunostained for stage-specific cell types. Tracking of both types of QD-Exos resulted in co-localization with oligodendrocytes, microglia, and astrocytes at different rates; no uptake in neurons was observed.

QD positive cells are listed as a percentage of cells measured (Fig. 10C). The 700 nm thick sections allowed for accurate visualization of cell-specific staining surrounding an unstained nucleus. In this fashion 60 cells per cell-specific staining group were counted and the number of QD-positive cells noted. QD-IFNγ-DC-Exos in slice showed that they were preferentially taken up by oligodendrocytes (72%) and to a lesser extent microglia (34%) and astrocytes (12%). In comparison, QD-unstimulated-DC-Exos were found to also co-localize with oligodendrocytes but to a lesser extent (7%), with uptake by microglia being similar (38%) to QD-IFNγ-DC-Exos, and astrocytes having the highest uptake at 63%. This suggests a difference in surface composition, where IFNγ-DC-Exos were significantly \((p < 0.001)\) targeted to oligodendrocytes and unstimulated-DC-Exos were significantly \((p < 0.001)\) targeted to astrocytes.

### 3.10. Transfection of primary OPCs with miR-219 mimic showed effects on differentiation and proliferation similar to treatment with IFNγ-DC-Exos

To determine if IFNγ-DC-Exos increase OPC differentiation through miR-219, a miR-219 mimic was applied to primary OPC cultures. Primary OPC cultures were grown at low density on glass coverslips, and either treated with IFNγ-DC-Exos or transfected with a miR-219 mimic. Supplementation with T3, which induces OPC differentiation, was used as a positive control. Three days after treatment, IFNγ-DC-Exo treated OPCs showed increased differentiation compared to control cultures, as determined by increased staining for stage-specific markers, O4 (Fig. 11A) and O1 (Fig. 11B). OPC cultures transfected with the miR-219 mimic likewise showed increased differentiation (Fig. 10A and B). Quantification of the percent O4 and O1 positive cells per treatment group revealed that both IFNγ-DC-Exos and the miR-219 mimic promoted OPC differentiation to the same extent as treatment with T3 (positive control), and were significantly \((p = 0.002\) and \(p < 0.001\), respectively) increased from control (Fig. 11C and D).
Additionally, the extent of OPC proliferation was also determined by Ki-67 staining after the above treatments. Compared to control, cultures treated with IFNγ-stimulated-DC-Exos and those transfected with the miR-219 mimic showed significant (p < 0.001; n = 3 animals/group) increases in O4 positive OPC proliferation, and proliferated to the same extent as cultures supplemented with T3 (positive control) growth factor (Fig. 11E).

4. Discussion

Current MS therapeutics are largely designed to reduce inflammation that contributes to initial demyelination, but do little to promote remyelination. In this study we focused on restoring remyelination, which is lost in MS and significantly contributes to disease progression (Hogancamp et al., 1997). Previously (Pusic and Kraig, in press), we examined the role of environmental enrichment (EE; volitionally increased physical, intellectual, and social activity) in improving outcomes of neurodegenerative disorders, including demyelinating diseases such as MS (Laviola et al., 2008). EE enhances memory and increases production of myelin at all ages (Fields, 2008). We discovered that exosomes found in the serum of EE animals may play a role in the increase in myelination. These serum-derived exosomes contained increased miR-219 relative to exosomes isolated from non-enriched animals (Pusic and Kraig, in press). In this study, we promote the use of stimulated DC exosomes as an effective treatment for remyelination, and highlight a number of their positive attributes.

We explored the use of DCs as a scalable, exogenous source of pro-mytelinating exosomes and found that stimulation with IFNγ induced production of exosomes containing high levels of miR-219. The contained miR-219 is functional when delivered to slice cultures via exosomes, as evidenced by decreased levels of two targets, PDGFRα and ELOVL7 in treated cultures versus controls (Fig. 6). Application of these exosomes to slice cultures increased levels of MBP, which was reflected in the production of structurally normal and thicker myelin sheaths (Fig. 2). IFNγ-DC-Exos also effectively increased MBP in MS modeled by acute demyelination of slice culture, further indicating their ability to promote recovery (Fig. 7). As miRNA-mediated regulation of various stages of oligodendrocyte maturation and myelin production plays an important role in CNS myelination, we screened exosomal miRNA content with a focus to these species (Fig. 5) (Dugas et al., 2010; Zhao et al., 2010). miR-219 plays an essential role in the formation and maintenance of compact myelin (Dugas et al., 2010). It has been shown to be necessary for OPC differentiation (Shin et al., 2009), and is deficient in MS lesions (Junker et al., 2009). The high miR-219 content in IFNγ-DC-Exos suggests that these DC exosomes can modulate oligodendrocyte maturation by targeting multiple steps in the differentiation pathway, as previously observed with EE derived exosomes (Pusic and Kraig, 2013). Other miRNAs also involved in the oligodendrocyte differentiation pathway were found in IFNγ-DC-Exos. miR-9, which is induced during normal oligodendrocyte maturation and enhances differentiation (Lau et al., 2008; Lin and Fu, 2009), was significantly enriched in these exosomes. Additionally, mature species of the miR-17–92 cluster, known to induce OPC proliferation both in vivo and in vitro (Budde et al., 2010), were also present and/or enriched. Thus, IFNγ-DC-Exos contain high levels of several miRNA species that are involved in oligodendrocyte development pathways (Fig. 5A), which may account for the observed increase in myelin.
Furthermore, the presence of the miR-17–92 cluster indicates that improved recovery from lysolecithin-induced demyelination via differentiation of OPCs can be accomplished without the depletion of progenitors. This was validated by equivalent counts of progenitors prior to and after exosome treatments in hippocampal slice cultures (Fig. 3). Treatment with IFN-γ-DC-Exos was also shown to increase proliferation of OPCs, thus it is likely that IFN-γ-DC-Exos can replenish and stabilize the population of OPCs.

Furthermore, we found that IFN-γ-DC-Exos also increased oxidative tolerance of slice cultures (Fig. 4A–B) and also increased antioxidant levels in microglia (Fig. 4C–D), an effect that potentially involves anti-inflammatory miRNAs. A number of miRNA species involved in anti-inflammatory responses were found to be highly enriched in IFN-γ-DC-Exos compared to unstimulated-DC-Exos (Fig. 5B). For example, miR-181a is involved in regulation monocyte/macrophage responses, where it acts to dampen pro-inflammatory signaling and

Fig. 10. IFNγ-stimulated-DC-Exos preferentially enter oligodendrocytes. (A) Merged images (top row) of QD tagged IFNγ-stimulated-DC-Exos (middle row, red) and cell-specific immunofluorescence (bottom row, green). (B) Merged images (top row) of QD tagged unstimulated-DC-Exos (middle row, red) and cell-specific immunofluorescence (bottom row, green). Left to right oligodendrocytes (anti-CNPase), microglia (anti-Iba-1), astrocytes (anti-GFAP), and neurons (anti-NeuN). Scale bars (A and B), 10 μm. (C) Uptake of QD tagged IFNγ-stimulated-DC-Exos and QD tagged unstimulated-DC-Exos for each cell type. Percent uptake calculated per 60 cells from n = 3 slices/group. These results indicate that DC exosomes can track to specific brain cell types. IFNγ-stimulated-DC-Exos were significantly (*, p < 0.001) localized to oligodendrocytes, while unstimulated-DC-Exos were significantly (*, p < 0.001) localized to astrocytes. Significance was determined by Student’s t-test.
reduces production of reactive oxygen species (Xie et al., 2013). Recent evidence suggests that miR-181a plays a similar role in regulating inflammation in the CNS (Hutchison et al., 2013). Another enriched miRNA, miR-124, promotes anti-inflammatory signaling and downregulates M1-associated pro-inflammatory IL-6, TNFα and iNOS (Ponomarev et al., 2013). Importantly, high levels of miR-27a were found in IFNγ-DC-Exos. miR-27a is expressed at significantly higher levels throughout the lifespan of the Ames dwarf mouse versus wildtype mice. Ames dwarf mice are used to study healthy aging due to their longevity, and miR-27a may contribute to their increased oxidative tolerance via regulation of glutathione synthesis (Bates et al., 2010). Thus, miRNA species may be responsible for the reduced OS and increased glutathione observed in IFNγ-DC-Exo treated slice cultures. The potential for immune modulation via this exosomal delivery of specific miRNAs should be further explored. It is possible that IFNγ-DC-Exos can also be utilized for immune suppression, as seen with traditional MS drugs, while simultaneously serving to promote myelin repair. Also, IFNγ-DC-Exos might be utilized for immune suppression, as seen with traditional MS drugs, while simultaneously serving to promote myelin repair.

Though we chose to focus on miR-219 for its well-documented involvement in oligodendrocyte differentiation, it will be important to further explore the relative contributions of other highly enriched miRNA species to the observed increase in myelination. It is likely that synergistic activity of multiple miRNA species is necessary to produce conditions supportive of myelination/remyelination.

Nasal administration of drugs to treat MS has recently been shown to be an effective means to lessen the effects of this malady (Duchi et al., 2013). We extend our proof-of-principle work using slice cultures to suggest that IFNγ-DC-Exos may also be effectively delivered. Administration of IFNγ-DC-Exos nasally resulted in a significant increase in

![Fig. 11. miR-219 mimic and IFNγ-stimulated-DC-Exos similarly promote OPC differentiation. (A) Representative images of O4 positive staining with DAPI counterstain used for quantification. Scale bars (A and B), 20 μm. (C) Representative high-power image of O4 staining to illustrate morphology, and quantification of percent of O4 positive cells (O4+ cells/total DAPI+ cells) per field (3 images per coverslip, n = 3 coverslips/group). Scale bar, 10 μm. Treatment with miR-219 mimic and IFNγ-stimulated-DC-Exos stimulated differentiation of OPCs into O4 expressing cells similar to T3 supplementation, and all groups were significantly (*, p < 0.001) increased from control. (D) Representative high-powered image of O1 positive staining and quantification of percent of O1 positive cells (O1+ cells/total DAPI+ cells) per field (3 images per coverslip, n = 3 coverslips/group). Scale bar, 10 μm. Treatment with miR-219 mimic and IFNγ-stimulated-DC-Exos stimulated differentiation of OPCs into mature O1 expressing cells similar to T3 supplementation, and all groups were significantly (*, p = 0.002) increased from control. (E) Representative images of O4 and Ki-67 positive staining of OPCs given different treatments. From left to right: control, T3, miR-219 mimic, and IFNγ-stimulated-DC-Exos treatments. Quantification of percent O4/Ki-67 double positive cells per field (3 images per coverslip, n = 3 coverslips/group) shows that treatment with miR-219 mimic and IFNγ-stimulated-DC-Exos increased OPC proliferation similar to T3 supplementation, all groups were significantly (*, p < 0.001) increased compared to control. Scale bar, 10 μm.](image-url)
myelin (Fig. 8) within normal brain. Having confirmed that IFN-γ-DC-Exos are effective in vivo to promote myelin, we began in vitro tracking studies to determine the cellular targets. Though both QD-tagged exosomes were found in oligodendrocytes, astrocytes, and microglia, the QD-IFN-γ-DC-Exos were preferentially taken up by oligodendrocytes (72%), whereas QD-unstimulated-DC-Exos showed no oligodendrocyte specificity (7%) (Fig. 10).

If exosome uptake is governed by endocytosis alone, we would have expected highest uptake by microglia, as they have the greatest endocytic activity of neural cell types. However, though there was a comparable and considerable uptake of both types of exosomes by microglia, there was a striking differential increase in uptake by other cell types. This suggests that exosome uptake can be receptor mediated, and indicates a difference in surface composition of each exosome type. Thus, IFN-γ-DC-Exos may affect myelin by directly delivering specific microRNA species, such as miR-219, to oligodendrocytes. A better understanding of the mechanism by which IFN-γ-DC-Exos interact with oligodendrocytes will be an important next step to further improve efficacy. For example, this could aid in devising strategies to specifically target IFN-γ-DC-Exos to oligodendrocytes in order to maximize remyelination. However, the contribution of microglia to the observed increase in remyelination should also be explored. The second-highest uptake of IFN-γ-DC-Exos was seen by microglia, and it is possible that they may have indirect effects on myelination and/or may be necessary to create a microenvironment that supports remyelination. Further characterization of the effects of IFN-γ-DC-Exos on microglia will be a necessary next step. Additionally, though astrocyte endocytosis is expected to increase in response to inflammatory stimuli, application of unstimulated-DC-Exos produced no significant increase in slice culture oxidative stress, an indirect read-out of increased inflammation, relative to control slices. The significance of preferential uptake of unstimulated-DC-Exos by astrocytes should therefore be further explored.

Since tracking studies suggest preferential uptake of IFN-γ-DC-Exos by oligodendrocytes, and IFN-γ-DC-Exos contained high levels of miR-219, we assessed their ability to affect OPC differentiation via miR-219 using primary OPC cultures. Primary OPCs treated with IFN-γ-DC-Exos or transfected with a miR-219 mimic differentiated to a similar degree (Fig. 11). This provides evidence for the role of IFN-γ-DC-Exo delivery of miR-219 in promoting remyelination.

Recent evidence has indicated DC involvement in the pathogenesis of MS and that the function of these cells may be altered to improve MS (Nuyts et al., 2013). Here we suggest for the first time the use of these stimulated cells in vitro to generate nutritive exosomes that can remyelinate normal and damaged brain. Our study provides strong evidence that exosomes can be an effective therapeutic for remyelination. Furthermore, our approach indicates that, dependant on the stimulation mechanism by which IFN-γ-DC-Exos was seen to affect remyelination within normal brain. Having con

development of DC exosomes as a form of gene therapy. Exosomes could aid in devising strategies to specifically target DC exosomes to certain cell types to promote the repair of damaged myelin.

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Appendix A. Supplementary data

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References


