Phenotypic and Functional Characterization of Müller Glia Isolated from Induced Pluripotent Stem Cell-Derived Retinal Organoids: Improvement of Retinal Ganglion Cell Function upon Transplantation

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ABSTRACT

Glaucoma is one of the leading causes of blindness, and there is an ongoing need for new therapies. Recent studies indicate that cell transplantation using Müller glia may be beneficial, but there is a need for novel sources of cells to provide therapeutic benefit. In this study, we have isolated Müller glia from retinal organoids formed by human induced pluripotent stem cells (hiPSCs) in vitro and have shown their ability to partially restore visual function in rats depleted of retinal ganglion cells by NMDA. Based on the present results, we suggest that Müller glia derived from retinal organoids formed by hiPSC may provide an attractive source of cells for human retinal therapies, to prevent and treat vision loss caused by retinal degenerative conditions.

INTRODUCTION

Glaucoma is one of the leading causes of blindness throughout the world [1]. It is characterized by high intraocular pressure, gradual loss of retinal ganglion cells (RGCs), and optic nerve damage [2, 3]. Current strategies to treat glaucoma only slow progression of the disease, and not all patients respond well to treatment, leading to severe sight loss and visual disability. Recent studies indicate that cell transplantation therapies may be developed with the aim to provide neurotrophic support to maintain the viability and function of remaining neurons and to potentially repair axonal damage.

Müller glia with stem cell characteristics were first identified in the zebrafish [4], in which they are responsible for the complete regeneration of the adult retina after injury [5, 6]. In this species, Müller glia re-enter the cell cycle to generate multipotent progenitors that proliferate, migrate, and differentiate into most neural cell types [7], that also restore retina function [8]. Although complete retinal regeneration has not been observed in other species, limited regenerative potential of Müller glia has been observed in chick [9] and rodent [10, 11] retinas. In rodent retina in vivo, it is reported that Müller glia can re-enter the mitotic cycle to generate amacrine cells in response to growth factors [10] or photoreceptors in response to N-methyl-D-aspartate (NMDA) [11]. A population of Müller glia isolated from the adult human retina has also been shown to have stem cell

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characteristics (human Müller stem cells [hMSC]) in vitro. These cells, can be isolated from cadaveric donors, become spontaneously immortalized in vitro, and acquire markers and function of retinal neurons after culture with various growth and differentiation factors [12–14]. However, there is no evidence of regeneration occurring after disease or injury in humans. That Müller glia may have potential for therapeutic application in glaucoma derives from experimental studies showing that hMSCs have the ability to partially restore visual function in rodent and feline models of NDMA-induced RGC damage [15, 16]. In addition, when directed toward a photoreceptor fate, these cells were shown to improve rod function in the P2H3 rat (a model of retinitis pigmentosa) after subretinal transplantation [17]. Müller glia derived from cadaveric donors present major difficulties for clinical application because of the risks of disease transmission caused by prions and nonidentified pathogens, as well as limitations because of the histocompatibility issues. Pluripotent stem cells, however, have the potential to overcome these issues, and recent studies have shown that retinal organoids that exhibit the characteristics of a whole laminated neural retina can be generated by both human induced pluripotent stem cells (hiPSC) and embryonic stem cells (hESC) in vitro [18, 19]. Various sources of hESCs and hiPSCs that comply with regulatory requirements for human therapies have become available and this will potentially facilitate the derivation of cells for retinal therapies in the near future.

This study aimed to isolate Müller glia from retinal organoids formed by hiPSCs in vitro and to assess their ability to partially restore visual function in rats depleted of RGC by NDMA. We examined the purity and nature of the cells isolated from these organoids by determining the gene and protein expression of well-known markers of Müller glia. We also investigated the ability of these cells to partially restore visual function upon transplantation into the vitreous of rats depleted of RGC as previously shown with Müller glia isolated from adult human retina [15]. The outcome of the transplantation was evaluated by assessing the negative scotopic threshold response (nSTR) of the electroretinogram (ERG) and by immunohistochemical analysis of the transplanted retina in vitro. Based on the present results, we propose that Müller glia derived from retinal organoids formed by hiPSCs may provide an attractive source of cells for human retinal therapies, to prevent and repair vision loss caused by retinal degenerative conditions such as glaucoma.

**MATERIALS AND METHODS**

**Stem Cell Maintenance and Retinal Organoid Differentiation**

The hiPSC line used in this study was created from BJ fibroblasts (Stemgent, Glasgow, U.K. Cat. No. 08-0027) as previously described by Carter et al. [20]. BJ iPSC cells were maintained in TeSR-E8 (StemCell Technologies, Cambridge, U.K.) and grown on Matrigel (Corning, St David’s Park, U.K.)-coated six-well plates. Differentiation of human pluripotent stem cells into retinal organoids was based on previous protocols developed by Nakano et al. [18] and Parfitt et al. [21]. Upon confluence, cells were washed with ×1 phosphate-buffered saline (PBS) and dissociated with TrypLE (ThermoFisher, U.K.) containing 10 μM ROCKi (Y-27632, Millipore, Watford, U.K.) and 0.5 mg/ml DNase (Sigma-Aldrich, U.K.) and pelleted. Cells were seeded to a density of 9,000 cells per well in a v-bottomed 96-well plate (PrimeSurface Sumilon low adhesion, Alpha Laboratories, U.K.) in Glasgow Minimum Essential Medium (GMEM) with L-glutamine containing 20% KOSR (ThermoFisher), 1% ×100 sodium pyruvate (ThermoFisher, U.K.), 1% ×100 nonessential amino acids (ThermoFisher), 1% ×100 penicillin/streptomycin (ThermoFisher, U.K.), and 50 μM β-mercaptoethanol (ThermoFisher, U.K.) containing 20 μM ROCKi and 3 μM Wnt antagonist (Millipore, Watford, U.K.). After 2 days in culture, wells were topped up with 100 μl of GMEM media containing 20% Knockout serum replacement (KOSR), 1% ×100 sodium pyruvate, 1% ×100 nonessential amino acids, 1% ×100 penicillin/streptomycin, 50 μM β-mercaptoethanol, 20 μM ROCKi, 3 μM Wnt antagonist, and 2% Matrigel. Half media change was performed twice weekly. On day 12, embryoid bodies (EBs) formed were transferred into separate wells of a 25-well squared, low-adhesion plate, and incubated in GMEM with L-glutamine, 20% KOSR, 10% fetal calf serum (FCS) 1% ×100 sodium pyruvate, 1% ×100 nonessential amino acids, 1% ×100 penicillin/streptomycin, and 50 μM β-mercaptoethanol containing 1% Matrigel and 100 nM smeothened agonist (SAG; Millipore, Watford, U.K.). On day 15, the same media was replaced containing 1% Matrigel and 100 nM SAG. On day 18, medium was replaced with Dulbecco’s modified Eagle’s medium/F12-glutamax containing 10% FCS, 1% ×100 N2 supplement (ThermoFisher, U.K.; 1% penicillin/streptomycin/amphotericin and 0.5 μM retinoic acid [Sigma-Aldrich, U.K.]). EBs, from which retinal organoids become visible between days 18 and 30, were then fed twice weekly. From days 30 to 50, retinal organoids were cut from the EBs under the microscope using microblades and placed into new 25 square low-adhesion plates. This was to ensure that only neural retinal tissue remained and matured in culture. Before dissection, any pigmented cell, occasionally forming with the organoids, was removed to prevent any possible retinal pigment epithelial contamination in the growing cultures. For long term, culture medium was replaced twice weekly (illustrated in Fig. 1A). All cells were maintained at 37°C, 5% CO2, and atmospheric O2.

**Isolation of Müller Glia from Retinal Organoids**

Müller cells were harvested from retinal organoids between 30 and 281 days after initiation of retinal differentiation for characterization analysis. Cells used for intravitreal transplantsations were derived from organoids at day 281. Organoids were dissociated using a papain dissociation kit protocol supplied by the manufacturer (Worthington Biochemical, Lakewood, NJ, USA). Briefly, retinal organoids were incubated with papain for approximately 10-15 minutes at 37°C with agitation by pipetting every 5 minutes. Pelleted cell suspensions were then plated as described previously for Müller glia isolated from human cadaveric retina (illustrated in Fig. 1B).

**Immunohistochemistry of Retinal Organoids and Isolated Cells**

Cells were grown to a confluent monolayer on fibronectin-coated glass coverslips and washed in PBS before fixation in 4% paraformaldehyde for 5 minutes. After fixation, cells were cryoprotected in 30% sucrose for 30 minutes and allowed to dry before freezing. For use, cells were defrosted and 500 μl of tris-buffered saline (TBS) + 0.3% triton X was added to the cells. Cells were blocked for 1 hour in TBS + 0.3% triton + 5% donkey serum before the addition of the primary antibody (diluted in blocking buffer). Primary antibodies (see Supporting Information Table S1)
were incubated overnight at 4°C. Cells were then washed with TBS three times for 5 minutes. Secondary antibodies (Alexa flour, Invitrogen, U.K. 1:500 in TBS + 0.3% triton) were incubated for 3 hours at room temperature in the dark. Slides were then washed in TBS and coverslips mounted with Fluoroshield Mounting Medium containing 4′,6-diamidino-2-phenylindole (DAPI; Abcam, Cambridge, U.K.), sealing with nail varnish.

**Flow Cytometry Analysis**

Cells were detached from tissue culture flasks to obtain a single cell suspension. These were then washed in PBS to remove any remaining tissue culture medium. Cells were then incubated with primary antibodies to CD29, CD44 (positive controls), SSEA4, and cytokeratin-18 (negative control markers) for 30 minutes in the dark. The cells were then washed three times by centrifugation and resuspended in sterile PBS. Corresponding isotype controls and a no primary antibody cell suspension were used as negative controls (see Supporting Information Table S1). Fluorescence-activated cell sorting (FACS) analysis was performed using a The BD LSRFortessa X-20 cell analyzer (BD Biosciences, Berkshire, U.K.). Supporting Information Fig. S1 shows data for negative markers SSEA4 and cytokeratin-18, and Supporting Information Fig. S2 shows flow cytometry data for all isotype controls.

**Gene Expression Analysis by Reverse Transcription-Polymerase Chain Reaction**

Total cellular RNA was isolated from cell pellets using the RNeasy system (Qiagen, Hilden, Germany; http://www.qiagen.com). For each reverse transcription reaction, 1 μg of total RNA was reverse transcribed in 20-μl reactions by adding 1 μl of oligo d(T)12-18 primers (Invitrogen) and 1 μl of dNTP (Promega) to each sample and incubated for 5 minutes at 65°C in a thermal cycler before adding 4 μl of first strand buffer (Invitrogen), 1 μl 0.1 M DTT, 0.5 μl Rnase inhibitor (RNasin Plus, N2611, Promega, U.K.), and 1 μl of SuperScript IV (Invitrogen) and incubating in the thermal cycler for a further 10 minutes at 55°C and 10 minutes at 80°C. Polymerase chain reaction (PCR) amplification was performed in 20-μl volume by addition of 10 μl of GoTaq Green master mix (Cat. No. M712; Promega, U.K.), which contains a mix Taq DNA polymerase, dNTPs, MgCl2, and reaction buffers and was used along with 7 μl RNase free water, 1 μl of each forward and reverse primers, and 1 μl cDNA. The mixture was incubated at 94°C for 5 minutes, followed by an appropriate number of cycles as follows: 94°C for 1 minute, annealing temperature °C for 1 minute, 72°C for 1 minute; and one cycle of 72°C for 5 minutes. PCR products were analyzed by agarose gel electrophoresis (2%) containing gel red (Supporting Information Table S2) for primer sequences.

**Animal Husbandry, Tolerization, Immunosuppression, and Anesthesia**

Wild-type Lister hooded rats were maintained according to the U.K. Home Office regulations for the care and use of laboratory animals (Scientific Procedures Act 1986). The use of the animal species for the study was approved by the local ethics committee at University College London, Institute of Ophthalmology.
and the U.K. Home Office. The animals were given access to food and water ad libitum and kept under 12-hour light/12-hour dark cycles. All animals used in the study were tolerated within 24 hours of birth by an intraperitoneal injection of $1 \times 10^5$ Müller glia derived from iPSC retinal organoids using protocols described previously [15]. Anesthetic for intravitreal transplantation procedures consisted of ketamine (60 mg/kg) and xylazine (7.5 mg/kg). Immunosuppression, started 2 days before cell transplantations, was administered daily in drinking water, and consisted of 25 mg azathioprine, 5 mg prednisolone, and 210 mg cyclosporin per liter.

**Induction of RGC Damage by NMDA and Intravitreal Müller Cell Transplantation**

Four-week-old animals were anesthetized and pupils treated with tropicamide (1% w/v, Minims; Bausch and Lomb, Kingston-upon-Thames, Surrey, U.K.), phenylephrine hydrochloride (2.5% w/v, Minims; Bausch and Lomb, Kingston-upon-Thames, Surrey, U.K.), and oxybuprocaine hydrochloride (0.4% w/v, Minims Bausch and Lomb, Kingston-upon-Thames, Surrey, U.K.) drops. Viscotears (Bausch and Lomb, Kingston-upon-Thames, Surrey, U.K.) were used to make sure the fellow eye did not dry out during the procedure. Injections were performed using a 5 μl Hamilton syringe and 32g needle. RGC damage was induced by injection of 2 μl of a mixture of NMDA (80 μM) and triamcinolone (80 mg/ml) into the intravitreal space of the left eye. To allow for assessment of RGC damage at 1 week after NMDA injection (Supporting Information Fig. S3), we performed ERG analysis at this time point. To comply with U.K. Home Office regulations and to allow for anesthetic recovery from the ERG test, cells were transplanted a week after the first ERG test. Animals were injected with 2 μl of a cell suspension consisting of 1 μl chondroitinase ABC and 1 μl of cells into the same eye (illustrated in Fig. 1C). Because Müller cells were frozen in small numbers upon isolation from the organoids, to obtain sufficient number of Müller glia for transplantation and characterization, we expanded these cells over a period of 2 weeks before injection. Three different injections were given to each of three experimental groups (n = 16 for each group) as follows—group A: medium-no cells (sham); group B: $4 \times 10^4$ Müller glial cells derived from hiPSC retinal organoids; and group C: $1 \times 10^5$ cells Müller glial cells derived from hiPSC retinal organoids. Four weeks after transplantation, animals were assessed for their RGC function by dark adapted ERGs and subsequently euthanized by terminal anesthesia, followed by paraformaldehyde (4%) perfusion.

**Scotopic ERG Recordings**

Animals were dark adapted overnight before ERGs. Rats were anesthetized as described above, and both eyes treated with tropicamide (1% w/v) and phenylephrine hydrochloride (2.5% w/v) to dilate the pupils and oxybuprocaine hydrochloride (0.4% w/v) to anesthetize. Viscotears were used to ensure the cornea remained hydrated. Animals were housed in a Faraday cage for ERG recordings and placed on a heated table to control the body temperature, in a Ganzfeld stimulator (Color dome; Diagnosys, Cambridge, U.K.; http://diagnosysllc.com). Flash stimuli (4-millisecond duration, repetition rate of 0.13 Hz) were presented in the Ganzfeld color dome by light-emitting diode stimulator at intensities of $-6.5$ to $-2$ log cd second/m². The flash stimulus was repeated 30 times for each light intensity. The responses were measured using the Espion Diagnosys V6 software. Measurements for nSTR analysis were analyzed by determination and recording of the maximal negative response in the range of 160–230 milliseconds (for nSTR) at each light intensity.

All ERG analyses were performed on R version 3.4.0 (The R Foundation for Statistical Computing) using the R program “Analyse summary statistics R.” An analysis of variance model was fitted to each intensity in turn. That model investigated effects of day and treatment group. The equivalent ERG summary post-NMDA was considered as a potential covariate but no relationship was observed ($p > 0.10$), so the summary was not included in the model. Least square (LS) means were obtained for each treatment group, and differences in mean response to the control group were presented together with corresponding 95% confidence intervals (CI). As an additional exploration, a mixed effects model was fitted to data for intensities $-4.5$ to $-3$ combined. The fixed effects included in the model were day, intensity, treatment, and treatment by intensity interaction. Animal was fitted as a random effect. The LS means were obtained as an averaged for each treatment across the intensities used in the test, and differences in average response to the control group were presented, together with 95% CI. The assumptions of all models were assessed visually using diagnostic plots.

**Immunocytochemistry and Confocal Microscopy for Visualizing Transplanted Cells**

After perfusion fixation, eyes were enucleated and placed in 4% paraformaldehyde overnight before replacing with 30% sucrose for cryopreservation. The lens was then removed from each eye and the remaining orbit embedded in OCT before sectioning at 16 μM using a Leica cryostat and mounting on Superfrost plus slides (VWR, U.K.). Sections were washed with TBS and blocked at room temperature with TBS containing 0.3% Triton and 5% donkey serum for 1 hour. To identify the transplanted human Müller glia in the rat eyes, mouse monoclonal antibodies to human CD29 (1:300; Santa Cruz Biotechnology, U.S.A) and human Nestin (1:300; Millipore, Watford, U.K.) were used in conjunction. Primary antibodies were diluted in blocking buffer and incubated overnight at 4°C. Slides were then washed 3 times in TBS, and both primary antibodies were probed with the secondary antibody Alexa fluor 488 (Invitrogen, U.K.) (1:500; 3 hours; Thermo Fischer, U.K.). Slides were washed in TBS 3 times before addition of a mounting media containing DAPI, covered with a glass coverslip, and sealed with nail varnish for microscope examination. Immunofluorescence staining was captured using a Leica 710 confocal microscope.

**RESULTS**

**Müller Glia Development Within Retinal Organoids**

Two days after initiation of the differentiation process, hiPSCs formed cellular spheroids characteristic of EBs. These EBs continued to enlarge, and by day 12, small cell protrusions were seen emerging from these structures. After 18 days, these protrusions were observed expanding on the periphery of the EBs to form nearly transparent cell structures, which denote retinal development. Between days 20 and 35, these retinal structures were sectioned from the EB and transferred to new plates to allow maturation for up to 300 days. During this period, retinal laminations could be increasingly observed by phase contrast microscopy (Fig. 2A). Immunocytochemical
staining of these organoids showed that after 90 days from initiation of the retinal differentiation process, Müller glia markers including Nestin, vimentin, glutamine synthetase, Chx10, and Cellular retinaldehyde binding protein (CRALBP) were expressed in cells that extend across the whole width of the retinal structure, which is one of the main features of Müller glia in the mature mammalian retina in vivo (Fig. 2B).

Figure 2. Retinal organoid development and Müller glia identification. (A): Representative phase microscopy images of retinal organoid development from induced pluripotent stem cells from days 2 to 300. (B): Confocal microscopy images of a retinal organoid at day 99 after initiation of differentiation, showing expression of the Müller glia-specific markers CRALBP, nestin, vimentin, Chx10, and glutamine synthetase. Scale bar = 20 μm.

After dissociation of retinal organoids, cells were plated on fibronectin-coated culture plates. Initially they formed small cell clusters (Fig. 3Ai) and after a few days in culture, many of these cell groups form neurosphere-like structures, which is often observed with Müller glia isolated from the adult human retina [13] (Fig. 3Ai). Cells were expanded in the presence of Epidermal growth factor (EGF) and Basic fibroblast growth factor (bFGF), and after 4–6 weeks culture, they were examined

Figure 3. Morphology and expression profile of Müller glia isolated from induced pluripotent stem cell-derived retinal organoids. (A): (i) Appearance of Müller glia after 1 day in culture (ii) characteristic formation of sphere-like structures at early passages. (iii) Müller glia grown in the presence of EGF alone display typical bipolar morphology and adhesion to culture plates. (iv) Müller glia cultured in the presence of EGF and FGF show fast growth and division as indicated by the presence of phase bright nuclei, characteristic of dividing cells. Scale bar = 100 μm. (B): Gel bands represent the expression of mRNA coding for Müller glia and progenitor factors in cells isolated from retinal organoids at different times after initiation of retinal differentiation. Duplicate bands represent two consecutive passages of the same cell preparation.

Characterization of Müller Glia Isolated from Retinal Organoids Formed by iPSC

After dissociation of retinal organoids, cells were plated on fibronectin-coated culture plates. Initially they formed small cell clusters (Fig. 3Ai) and after a few days in culture, many of these cell groups form neurosphere-like structures, which is often observed with Müller glia isolated from the adult human retina [13] (Fig. 3Ai). Cells were expanded in the presence of Epidermal growth factor (EGF) and Basic fibroblast growth factor (bFGF), and after 4–6 weeks culture, they were examined
for gene and protein expression of Müller glial cell markers. We observed that bFGF greatly increased expansion rate of the cell colonies but was not necessary for continued expansion after the first three passages (Fig. 3Aiii–iv). Proliferating cells showed a characteristic Müller glia bipolar-like morphology with membrane protrusions, characteristic of Müller glia. This morphology was maintained throughout several passages. If cells were maintained in the presence of FGF for a long term, a morphology characteristic of neurosphere-like forming cells with shorter processes and phase bright nuclei was often observed (Fig. 3Aiv).

Cells isolated from retinal organoids at 34, 70, 90, 219, and 281 days after initiation of organoid differentiation were examined for the expression of genes coding for Müller glia and neural progenitor markers, the characteristics of adult human Müller glia [13]. Expression of mRNA coding for the Müller glia markers CRALBP, glutamine synthetase, nestin, CD44, and Sox9, characteristic markers of Müller glia. Scale bar = 50 μm. Abbreviation: DAPI, 4’,6-diamidino-2-phenylindole.
Purity of the Müller Cell Preparation Obtained from Retinal Organoids

Cells isolated from retinal organoids were expanded on fibronectin-coated culture plates to obtain sufficient number for characterization studies. As determined by flow cytometry (FACS) analysis, the proportion of cells coexpressing the Müller cell surface markers CD44 and CD29 was above 99.7% (Fig. 4A). To confirm specificity, the cell preparations were also examined for the expression of negative markers including SSEA4, a known stem cell marker and cytokeratin-18, an epithelial cell marker. Both markers were shown to be under 1% positive for our Müller glia cell preparation (Supporting Information Fig. S1).

Immuno-fluorescence staining confirmed the expression of progenitor and Müller glia markers by the isolated retinal cells. Costaining for the cell surface marker CD29 and the intermediate filament protein vimentin showed that all cells were expressing both markers. Well-known Müller glia markers including CRALBP and glutamine synthetase also showed prominent cytoplasmic staining in the majority of the cells (Fig. 4B). Immuno-staining for the Müller glia cell surface marker CD44 was also observed in all the cells, with a large proportion also staining for nestin (Fig. 4B). In addition, nuclear staining for the Muller progenitor marker Sox9 was observed in all cells (Fig. 4B).

Intravitreal Transplantation of hiPSC Derived Müller Glia Partially Restores RGC Function in a Rat Model of RGC Depletion

Intravitreal cell transplantation was performed as previously described [15] following 2 weeks after intravitreal injection of...
NMAD to cause RGC damage. Sixteen animals were used in each of three experimental groups, receiving either a sham injection (medium alone), 4 × 10⁴ cells, or 1 × 10⁵ cells. To assess RGC function, scotopic ERGs were performed 4 weeks after cell transplantation to examine the nSTR, which denotes RGC function. Using 95% CI and compared with control treatment, statistical analysis of the effects of human Müller glia cell injection showed an increase in the amplitude of the nSTR across a range of relevant light intensities, with significant difference (p < .05) being demonstrated with 1 × 10⁵ human Müller glia cells at a light intensity of −3.5 log cd second/m² (Fig. 5A). Furthermore, statistical analysis of the data obtained from eyes injected with both 4 × 10⁴ and 1 × 10⁵ iPSC-derived human Müller glial cells when compared with control treatment demonstrated a dose response and significant improvement (p = .038) in the nSTR across light intensities ranging from −3.0 to −4.5 log cd second/m² (Fig. 5B). Control animals receiving no NMAD treatment but no cell transplantation exhibited reduced nSTR and larger b-waves for luminance −4 through to −3.0 log cd second/m² to −7 in comparison to animals receiving cell transplantations (Fig. 5C, arrows).

Antibodies to human CD29 and Nestin were used to locate the transplanted human cells in the rodent eye. Immunohistochemical staining of retinal sections from eyes transplanted with 1 × 10⁵ cells showed that cells were present in 13 out of the 16 eyes transplanted. Cells with healthy appearance were observed mostly in the vitreous (11 of 13), and although no integration of the transplanted Müller glia into the retina was observed, cells could be clearly seen strongly attached to the RGC layer in three cases (Fig. 6A, white arrow). In addition, transplanted cells were observed attached to the lens in two cases (not shown). Overall, fewer cells were observed in histological sections form eyes transplanted with 4 × 10⁴ cells as compared with those receiving the higher dose (illustrated on left image of Fig. 6B). In some cases, cell aggregates could be seen in the vitreous (illustrated on right image of Fig. 6B).

**DISCUSSION**

In this study, we have used human iPSC to produce retinal organoids for isolation and propagation of Müller glia in vitro. Müller cells isolated from the organoids were then used for transplantation into a rat model of RGC depletion by NMAD. Results presented in this study indicate that Müller glia that express CRALBP emerge at around day 90 after induction of differentiation of iPSC into retinal organoids. These observations are comparable to other studies that have reported expression of CRALBP in retinal organoids at similar stages of organoid formation [19]. Upon transplantation, these cells partially restored RGC function in the NMAD-damaged rat retina. These observations suggest that retinal organoids derived from iPSCs constitute an attractive source of Müller glia for cell transplantation therapies to treat retinal degenerative conditions. Therefore, we propose that Müller cells derived from organoids produced by pluripotent cells in vitro may be potentially used as a reliable and traceable source of cells for use in retinal therapies.

The data showed that Müller glia isolated from retinal organoids can be expanded over several passages and that their profile is comparable to published Müller cell lines derived from human cadaveric donors, including the established human MIO-M1 Müller cell line [12, 13]. Based on our previous studies, we have characterized these cells by their morphology as well as their gene and protein expression profiles which are distinctive of Müller glia. Isolated cells exhibited bipolar morphology and showed the ability to form sphere-like structures that have been previously observed in the establishment of other Müller glia cell lines when cultured in the presence of FGF2 [13]. Cells isolated from the organoids expressed mRNA and protein coding for well-known Müller glia markers including glutamine synthetase, CRALBP, nestin, and vimentin. The method of isolation is simple and exploited on the expression of CD29, a β1 integrin, by Müller glia, whose main ligand is fibronectin [22]. Hence, adherence of Müller glia to fibronectin-coated plates selectively promoted the emergence of highly pure populations of Müller glia (>97%) as determined by FACS analysis of the CD29 positive cell population, avoiding the need for further purification. This constitutes a simple method for selection and purification of Müller glia for regulatory compliance in the event that these cells can be used for cell therapies. It is also well documented that retinal neurons do not survive well in culture without specific on-going support from extracellular signals [23], suggesting that only glia are likely to survive and proliferate in culture under the conditions used.

We also investigated the ability of Müller glia isolated from retinal organoids formed by iPSC to restore RGC function in rats depleted of RGC by NMAD as previously shown in our laboratory [15]. This model mimics the pathological features of RGC damage observed in glaucoma and is a reliable model of RGC damage which can be easily assessed by measuring the nSTR response of the ERG [15]. Following NMAD damage, we observed a reduction in the nSTR and a slight elevation in the b-wave of the ERG, which is characteristic of similar glaucoma models [24]. We observed a partial but significant restoration of the nSTR after transplantation of the highest dose of cells used (1 × 10⁵). Although injection of a smaller number of cells (4 × 10⁴) also resulted in an improvement in the nSTR, this response was not significant, indicating the need for a suitable number of cells for transplantation to improve retinal function. As with previous experimental models, we observed cell aggregates in a small number of animals (Fig. 6). However, we do not know whether this effect could interfere with visual function. We could speculate that because floaters present in human vitreous do not obstruct visual acuity, small cell aggregates may not interfere with vision. However, further studies are needed to assess the impact of these cell aggregates before any translation into the clinic.

**CONCLUSION**

Results shown in this study are in agreement with previous findings from our laboratory, where we showed therapeutic benefit of the transplanted human Müller glia cell line MIO-M1 in this rat model of glaucoma-like damage [15]. As previously observed in the Lister hooded rat and feline models of RGC depletion by NMAD [15, 16], cell integration into the retina was not observed, yet the RGC function was partially restored. Several studies have shown that RGC depletion occurs after NMAD injection, which at present requires
ex vivo examination of the retinal tissue and prevents the assessment of transplantation in the same animal. Therefore, we undertook functional analysis by means of ERGs to assess RGC damage and the actual recovery of function in the NMDA-treated animals. This also minimized the number of animals used for the study. Studies by Goldberg and collaborators have shown that if the soma of the RGC is still intact, the cell recovers function in response to neurotrophic factors, but if it is damaged, they will not be repaired by these factors [25]. This is the mechanism by which we are proposing that Müller cells improve RGC function upon intravitreal transplantation.

Together the results show that Müller glia derived from pluripotent stem cells possess similar phenotypic characteristics and ability to partially restore visual function in NMDA depleted retina as Müller derived from the adult human retina. On this basis, it can be suggested that iPSC-derived Müller glia may constitute a valuable resource for transplantation studies that could be potentially translated into the clinic.

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AUTHOR CONTRIBUTIONS

G.A.L.: conception/design, financial support, manuscript writing, final approval of manuscript; K.E.: conception/design, collection and/or assembly of data, data analysis and interpretation, manuscript writing; W.W., C.M.-D., A.V., M.S.: collection and/or assembly of data; H.J., P.C.: provision of study materials; A.J.F.C., C.M.R.: administrative support, provision of study materials; K.G.: data analysis and interpretation; N.C.: conception/design, financial support, administrative support, provision of study materials final approval of manuscript; P.T.K.: conception/design, final approval of manuscript.

DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST

H.J. declared consultant/advisory role with Allergan and honoraria from Santen, Allergan, and Laboratories Thea. K.G. declared consultant/advisory role with Apollo Therapeutics. N.C. declare employment with Apollo Therapeutics that funded research in Prof. Limb’s lab. P.T.K. declare inventor or patent holder on Müller Cells and the use in retinal regeneration, consultant/advisory role and received honoraria from Santen, Belkin, Novartis, and Aerie, and received research funding from Apollo Therapeutics. G.A.L. declared inventor or patent holder on Müller cells and their use in retinal regeneration. The other authors indicated no potential conflicts of interest.
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