A Conditional Immortalized Mouse Müller Glial Cell Line Expressing Glial and Retinal Stem Cell Genes

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PURPOSE. Müller glia have multiple functions in the retina, including synthesis of neurotrophic factors, uptake and metabolism of neurotransmitters, spatial buffering of ions, maintenance of the blood-retinal barrier, and response to injury. A population of Müller glia has some stem cell-like characteristics both in vivo and in vitro. The purpose of this study was to generate and characterize novel Müller glial cell lines from the postnatal mouse retina.

METHODS. Cells were cultured from postnatal day (P) 10 double heterozygous transgenic (H-2Kb-tsA58+/−; HRhoGFP/+ or C57BL/6 mice after papain dissociation. Interferon gamma (IFNγ) induction of the SV40 T-antigen (Tag) was assayed by immunohistochemistry and Western blot analysis. Proliferation was assayed by BrdU uptake and cell counts of calcein AM/ethidium bromide-stained cells. Gene expression was analyzed by RT-PCR and immunohistochemistry.

RESULTS. Conditionally immortalized (ImM10 [Immortomouse Müller P10]) and spontaneously immortalized (C57M10 [C57BL/6 Müller P10]) Müller glial cell lines were selected by differential adherence to laminin; both consisted of adherent flat cells with large, diffusely staining nuclei and an epithelial morphology. Tag induction stimulated BrdU uptake by Müller glia in mixed retinal cultures from H-2Kb-tsA58+/−; HRhoGFP/+ mice and increased the proliferation of ImM10 cells. ImM10 and C57M10 cells expressed genes characteristic of Müller glia but not genes characteristic of differentiated retinal neurons. ImM10 cells also expressed retinal stem cell genes.

CONCLUSIONS. The ImM10 cell line is a novel, conditionally immortalized Müller glial cell line isolated from the P10 mouse retina that expresses genes characteristic of Müller glial and retinal stem cells. (Invest Ophthalmol Vis Sci. 2010;51: 5991–6000) DOI:10.1167/iovs.10-5395

Müller glia, the radial glia of the neural retina, function to maintain retinal homeostasis through synthesis of neurotrophic factors, uptake and metabolism of neurotransmitters, spatial buffering of ions during retinal activity, and maintenance of the blood-retinal barrier.1,2 In the mature retina in mammals, Müller glia proliferate in the context of retinal injury or disease, contributing to gliosis and the formation of glial scars.3 In addition, a subset of Müller glia show some stem cell characteristics and upregulate stem cell-associated genes, although their inherent regenerative ability is extremely limited in vivo.4–5 In contrast, in teleost fish Müller glia are the source of retinal stem cells in the inner retina that generate rod photoreceptors during normal growth and regenerate all types of retinal neurons after injury.6–8 The robust neurogenic capacity of Müller glia in fish and, to a limited extent, in birds9 has sparked research to understand the stem cell properties and neurogenic potential of Müller glia in the mammalian retina both in vitro and in vivo.9–15

Cell culture is useful for studying basic cellular and molecular processes including cell-cell interactions, cell movements, and regulation of gene expression. The ability to generate large numbers of cells in vitro has utility for high-throughput assays for drug development and tissue engineering applications. For studies of differentiation and regeneration, in vitro culture systems enable precise control of the cellular environment and remove cells from inhibitory constraints that may limit their regenerative capacity in vivo. The use of primary cells for studying basic cellular and molecular processes is hampered by the low number of cells that can be isolated, difficulty in obtaining pure populations of cells, and lack of inherent proliferative capacity of adult mammalian retinal cells. To overcome these problems, immortalized cell lines have been generated from tumors (e.g., Y79 and Weri-Rb1 retinoblastoma cell lines16,17), by constitutive overexpression of exogenous oncogenes in primary cells18–20 or by spontaneous immortalization.21 The increased proliferative capacity of transformed cells enables the generation of large numbers of cells. However, permanent immortalization alters the innate characteristics of primary cells; if used for in vivo transplantation, there are concerns that unregulated proliferation could generate tumors.

Conditional immortalization using inducible oncogenes enables precise control over the timing of immortalization.22 H-2Kb-tsA58 transgenic mice (Immortomouse; Jackson Laboratories, Wilmington, MA; hereafter designated tsA58) carry a transgene encoding an inducible, thermostable simian virus 40-large T-antigen (Tag) under control of the mouse major histocompatibility complex H-2kb class 1 promoter.23 Induction by interferon-gamma (IFNγ) upregulates TAg expression and immortalizes cells cultured at the permissive temperature (33°C). Removal of IFNγ represses transgene expression, and incubation at physiological temperatures (37°C–39°C) inactivates residual TAg, permitting differentiation. Conditionally immortalized cell lines have been generated from both neuronal and nonneuronal cells isolated from tsA58 mice and rats. Conditionally immortalized retinal cell lines described to date include rat Müller glia,24 retinal capillary endothelial cells25
and pericytes, and mouse retinal endothelial cells and astrocytes.

To generate novel cell lines from the postnatal mouse retina, we isolated retinal cells from transgenic mice heterozygous for the tsA58 and HRhoGFP transgenes and from C57BL/6 mice. The HRhoGFP transgene encodes a human rhodopsin-GFP fusion protein under the control of the endogenous rhodopsin promoter and is expressed specifically in postmitotic rod photoreceptors. This enables longitudinal imaging of living photoreceptors in culture and could prove useful in future studies of the neurogenic potential of conditionally immortalized Müller glia. We report the isolation and characterization of two novel Müller glia cell lines from the postnatal day (P) 10 mouse retina: conditionally immortalized ImM10 (Immortal mouse Müller postnatal day 10) and spontaneously immortalized C57M10 (C57BL/6 Müller postnatal day 10).

METHODS

Mice

All mice were handled and euthanatized according to the animal care and use policies of the University of Houston and Johns Hopkins University School of Medicine and the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. The following strains were used: C57BL/6 (Jackson Laboratories), and H-2 Kb-tsA58 transgenic (tsA58; Immortomouse; Charles River Laboratories). Homozygous tsA58 males were bred to homozygous HRhoGFP females to generate double heterozygous mice (tsA58/+; HRhoGFP/+) used for retinal dissociation and cell culture.

Retinal Dissociation

P10 pups were euthanatized by halothane or CO2 inhalation, eyeballs were enucleated, and retinas were dissected free of the retinal pigment epithelium and dissociated by incubation in phosphate-buffered saline (PBS) containing 16.5 U/ml activated papain (Worthington, Lakewood, NJ) and 124 U/ml DNase (Sigma-Aldrich, St. Louis, MO) for 30 minutes at 37°C with gentle trituration. Digestion was stopped by the addition of ovomucoid (Worthington, Lakewood, NJ); cells were pelleted by centrifugation (800 rpm, 8 minutes) and resuspended in plating medium (Neurobasal containing B27 supplement, 20 mM L-glutamine, and penicillin/streptomycin antibiotics (Pen/Strep)). Cell culture media, supplements, and reagents were obtained from Gibco/Invitrogen (Carlsbad, CA) unless otherwise indicated.

Cell Culture

Dissociated primary retinal cells were plated on poly-lysine/laminin-coated multiwell plates or 100-mm culture dishes and maintained in cell culture incubators at 5.5% CO2. To select Müller glia cells from coated multiwell plates or 100-mm culture dishes and maintained in antiformaldehyde conditions, GM was supplemented with mouse recombinant IFN (generous gift of Theodore Wessel), and H-2 Kb-tsA58 transgenic (tsA58; Immortomouse; Charles River Laboratories). Homozygous tsA58 males were bred to homozygous HRhoGFP females to generate double heterozygous mice (tsA58/+; HRhoGFP/+) used for retinal dissociation and cell culture.

Cell Culture

Dissociated primary retinal cells were plated on poly-lysine/laminin-coated multiwell plates or 100-mm culture dishes and maintained in cell culture incubators at 5.5% CO2. To select Müller glial cells from mixed cultures, papain-dissociated retinal cells were allowed to settle on 12-well dishes coated with poly-lysine/laminin for 5 hours; subsequently, nonadherent cells were transferred to empty poly-lysine/laminin coated wells and, after overnight incubation, remaining nonadherent cells were transferred to empty wells. After plating, adherent cells were maintained for 7 days at 37°C in plating medium under nonimmortalizing conditions and subsequently were transferred to immortalizing conditions in growth medium (GM; Neurobasal with 2% fetal bovine serum (FBS); Gibco/Invitrogen) and incubated at 33°C with 100% media changes every 3- to 5-day intervals. For growth under immortalizing conditions, GM was supplemented with mouse recombinant IFN (PeproTech; Rocky Hill, NJ) at 50 U/ml, unless otherwise indicated. To eliminate tsA58 transgene expression (nonimmortalizing conditions), primary cultures of mixed retinal cells and ImM10 Müller cells were cultured in GM without IFN and incubated from 37°C to 39°C. Primary cells from C57BL/6 were isolated according to the same method but were maintained at 37°C in DMEM/F12 medium containing 10% FBS and antibiotics (Pen/Strep; Gibco/Invitrogen). During the initial isolation of both ImM10 and C57M10 Müller cells, we used poly-lysine/laminin–coated plates. For maintenance and subsequent analyses, cells were grown on uncoated tissue culture dishes.

BrdU Assay

Cells were plated in GM+IFN by containing 10 μM BrdU and incubated at 33°C for 17 hours. Cells were washed in PBS, fixed in 10% formalin for 10 minutes, washed in PBS, and incubated in 2 N HCl in PBS for 30 minutes before immunostaining.

Immunohistochemistry

For immunostaining, tissues or cells were fixed in 4% paraformaldehyde or 10% formalin (for BrdU immunostaining) for 30 minutes. Samples for PAX2 immunostaining were fixed for 5 minutes in 4% paraformaldehyde. Samples were washed in PBS, incubated in 1% sodium borohydride (2 minutes), and blocked with either PBS containing 20% normal goat serum (0.5% Triton X-100 or PBS containing 10% normal goat serum/0.5% Triton X-100/1% fish gelatin/5% bovine serum albumin for 2 hours. Primary antibodies (Supplementary Table S1, http://www.iovs.org/cgi/content/full/51/11/5991/DC1) were applied overnight at 4°C. Secondary antibodies conjugated to AlexaFluor488, AlexaFluor543, AlexaFluor633 (Molecular Probes, Eugene, OR) or TRITC (Sigma, St. Louis, MO) were diluted 1:200 and incubated for 1 to 2 hours at room temperature. For BrdU labeling, a Cy3-conjugated primary antibody was used. Cells were counterstained with Hoechst 33542 or 4‘,6-diamidino-2-phenylindole (DAPI). Specificity of labeling was confirmed by omitting primary antibody or by substituting normal serum for the species used to generate the primary antibody. Immunostained cells were imaged with an inverted microscope (IX71; Olympus, Tokyo, Japan) with a monochromed, cooled CCD digital camera (Roiwa-VR; Q-Imaging, Surrey, BC, Canada). Images were subsequently pseudocolored and adjusted for contrast using image editing software (Photoshop; Adobe, San Jose, CA). Images used for comparison of staining intensity (BrdU, SV40Tag, PAX2 immunostaining) were photographed at a uniform exposure setting and prepared as a montage with brightness and color adjusted simultaneously for all images.

Western Blot Analysis

ImM10 cells in GM containing 0 to 200 U/ml IFN were grown on 100-mm plates for 6 days at 33°C, with 100% media changes every 2 days. Cells were harvested in PBS and sonicated. Cell lysates were quantified by bicinchoninic acid assay (BCA-1; Sigma-Aldrich), diluted in Laemmli buffer (2× final concentration) and were loaded at 15 μg/lane onto duplicate polyacrylamide gels (Bio-Rad, Hercules, CA). After electrophoresis at 100 V, proteins were transferred onto nitrocellulose (NT8017; Sigma-Aldrich) by electroblotting and processed for immunodetection using anti-TAG antibodies (RDI; Fitzgerald Industries, Concord, MA) at 1:250 and enhanced chemiluminescence (Amersham Biosciences, Pittsburgh, PA). Duplicate gels run in parallel and stained (GelCode Blue; Pierce, Rockford, IL) were used to compare protein loading.

Cell Counts

Proliferation assays used on-chip flow cytometry (Bioanalyzer 2100; Agilent, Foster City, CA). ImM10 cells were plated in 12-well dishes at 1 x 10^5 cells per well on duplicate plates and were cultured at 33°C in GM+IFN and at 39°C in GM without IFN. For cell counts, three individual wells for each condition were stained with 2.5 μM calcein AM and 2.5 μM ethidium bromide homodimer (Live-Dead Stain; Invitrogen) in PBS for 30 minutes. Cells were pelleted and resuspended in 50 μl cell buffer (cell kit #5067-1519; Agilent), and 10 μl resuspended cells from each sample was loaded into individual wells of a cell chip. Fluorescence on the two channels (calcein AM: excitation 470 nm/emission 535 nm; ethidium: excitation 630 nm/emission 680 nm) was read for 6 minutes per well. Wells that gave low counts in the
calcine channel were examined microscopically, and those with bubbles or blockage in the microfluidics channels were repeated on a second chip using the rest of the sample. The threshold for calcine fluorescence intensity for a “live cell” (>0.7 × 10^5) was determined empirically using calcine staining of methanol-killed cells and unstained cells. For passages 4 to 7, the entire experiment was repeated twice; for passages 14 to 17, the entire experiment was repeated three times.

**RNA Isolation and RT-PCR**

RNA was harvested from ImM10 and C57M10 cells and whole mouse retina using affinity columns (RNasea, Qiagen, Valencia, CA) according to the manufacturer’s instructions. Briefly, cells were washed in RNase/DNase-free PBS, lysed in a guanidinium thiocyanate containing buffer, and sheared using a 20-gauge needle before loading onto an RNA affinity spin column. Total RNA was eluted in RNase-free water and quantified by spectrophotometry (Nanodrop; Thermo Scientific, Wilmington, DE), and quality was assessed (RNA nano Chip on a Bioanalyzer 2100; Agilent). RNA samples with quality scores >9.5 were used for analysis.

For analysis of gene expression by quantitative RT-PCR, total RNA (200 ng) was reverse transcribed using oligo-dT primers (AffinityScript; Stratagene, La Jolla, CA). Primers (IDT, Coralville, IA) were designed to target the 3’ end of genes and to span introns when possible (Supplementary Table S2, http://www.iovs.org/cgi/content/full/51/11/5991/DC1). Primers were optimized using total retinal RNA and were analyzed by melting curve analysis and gel electrophoresis to verify amplification of a single product of the predicted size. Standard curves were generated to determine amplification efficiency and the linear range of amplification. PCR amplification (Brilliant II SYBR Green; MX3005p instrument; Stratagene) consisted of a 10-minute incubation at 95°C (denature), 30 seconds 60°C (anneal), and 30 seconds 72°C (extension). Reactions were conducted in triplicate on three independent RNA samples per condition. Fold-differences in gene expression were calculated using the ΔΔCt method, corrected for amplification efficiency. Statistical analysis was performed with pairwise fixed reallocation randomization using Relative Expression Software Tool [http://www.gene-quantification.de/rest.html]). Additional samples amplified without the melting curve analysis were separated by gel electrophoresis (1.0% agarose, 0.5% TBE), stained with ethidium bromide, and photographed.

**RESULTS**

**Mixed Retinal Cultures from tsA58/Rho-GFP Transgenic Mice**

Under immortalizing conditions (plating medium at 33°C, with 50 U/mL IFNγ), cultures of total retinal cells from PN10 retina of tsA58/++; HRhoGFP/+ mice consisted of multiple, morphologically distinct populations. One population consisted of large, flat, adherent cells that were vimentin positive and contained large nuclei containing diffuse, lightly stained chromat and prominent nucleoli (Figs. 1A, 1C). The second population consisted of abundant small, spherical cells with intensely staining nuclei that, by 4 days in culture, were almost exclusively clustered on the surfaces of the adherent cells (Figs. 1A-D). The spherical cells were vimentin negative, and most were HRhoGFP+ rod photoreceptors, as evidenced by the fluorescence of the GFP-tagged rhodopsin in their cell membranes and the polarized accumulation of HRhoGFP at the cell surface (Fig. 1B, small arrows). A subpopulation of the HRhoGFP-negative spherical cells was immunostained by antibodies against PKC (Figs. 1E-H, large arrowhead), consistent with a rod bipolar cell identity. In mixed cultures, rare pigmented epithelial cells were present at the initial plating and...
were identifiable as adherent cells containing distinct melanin granules. Melanin-containing cells were never associated with the large flat cells and were not observed in long-term cultures (data not shown).

The adherent flat cells and small spherical cells survived up to 50 days in mixed cultures. Over time, the flat cells covered increasing areas of the substrate, whereas the spherical cells did not appear to increase in number. This was confirmed using BrdU incorporation to assay DNA synthesis. After 17 hours of BrdU exposure in mixed retinal cultures (tsA58/+/HRhoGFP/+) grown in the presence of IFNγ at 33°C, BrdU immunoreactivity was detected exclusively in the large nuclei of the adherent cells (Figs. 2A–C, arrowheads).

To examine TAg expression, mixed retinal cells were cultured at 33°C for 4 days with or without 50 U/mL mouse recombinant IFNγ and analyzed by immunohistochemistry (Figs. 2G–L). TAg immunoreactivity was detected in cells cultured in the presence of IFNγ in both GFP-negative cells (non-rod photoreceptors; Figs. 2G–I, small arrows) and HRhoGFP-expressing rod photoreceptors (Figs. 2G–I, large arrowheads), although some rod photoreceptors lacked distinct TAg immunoreactivity (Figs. 2G–I, small arrowheads).

**Purification of ImM10 Cell line**

We used preferential adhesion to laminin-coated substrates to enrich for the adherent cell population in primary cultures (Supplementary Fig. S1, http://www.iovs.org/cgi/content/full/51/11/5991/DC1). After 26 days in culture, the adherent cells reached confluence. Only low numbers of HRhoGFP-positive

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cells were present after the initial enrichment and were rarely detected after the first passage. By the third passage, all cells in the culture were GFP-negative, adherent cells with large nuclei containing prominent nucleoli typical of Müller glia (Figs. 3A–H). This new cell line, named ImM10, has been maintained for up to 50 passages under immortalizing conditions.

Using the same approach, we also generated a cell line from nontransgenic C57BL/6 mice (P10) for comparison of gene expression between immortalizing and nonimmortalizing conditions (Fig. 4C). We also analyzed the expression of additional genes characteristic of Müller glial or other retinal cell types in ImM10 cells grown under immortalizing conditions, cultures of spontaneously immortalized Müller glia from C57BL/6 mice, cultured under nonimmortalizing conditions. 

Gene Expression in ImM10 Cells

Quantitative RT-PCR using a panel of primer pairs for eight genes expressed in Müller cells showed that ImM10 cells expressed vimentin, glutamine synthetase, clusterin, Dkk3, nestin, Pax6, Six3, and Hes1, with no significant differences in gene expression between immortalizing and nonimmortalizing conditions (Fig. 4C). We also analyzed the expression of additional genes characteristic of Müller glial or other retinal cell types in ImM10 cells grown under immortalizing conditions, cultures of spontaneously immortalized Müller glia from C57BL/6 mice, cultured under nonimmortalizing conditions.

Figure 3. Photomicrographs of representative cultures of ImM10 and C57M10 cell lines in culture. (A) Calcein-stained ImM10 cells, passage 14, cultured under immortalizing conditions (GM with 50 U/mL IFNγ, 33°C). (B) Calcein-stained ImM10 cells, passage 14, plated in parallel cultured under nonimmortalizing conditions (GM without IFNγ, 39°C). (C, D) DIC images of ImM10 cells, passage 5, cultured under immortalizing conditions. (E, F) ImM10 cells, passage 5, cultured under nonimmortalizing conditions. (G) ImM10 cells, passage 37, cultured under immortalizing conditions. (H) ImM10 cells, passage 37, plated in parallel cultured under nonimmortalizing conditions. (I) Spontaneously immortalized Müller cells from retinas of PN10 C57Bl/6 mice, passage 47, cultured in DMEM with 10% FBS. Scale bars: 500 μm (A, B); 200 μm (C–I).

Figure 4. Comparison of proliferation of Tag-induced and noninduced ImM10 cells in culture. (A) Passages 4 to 7 and (B) passages 16 to 18 cultured under immortalizing (open squares; 33°C, GM + IFNγ) or non-immortalizing (closed squares; 39°C, GM without IFNγ) conditions showing mean fold-change in live cells (calcein AM positive, ethidium bromide homodimer negative). Error bars show SEM. Early-passage cells were not counted on days 1 to 6 because visual observations and pilot studies showed minimal changes in cell number before 7 days in culture. (C) Quantitative RT-PCR comparing mRNA expression between cells cultured under immortalizing conditions (33°C) and nonimmortalizing conditions (39°C). Image shows ethidium bromide-stained agarose gels of amplicons. Table shows fold-change and P-values.
mice (DMEM/F12; 10% FBS, 37°C) using RT-PCR (Fig. 5P). Both ImM10 and C57M10 adherent cell cultures expressed the Müller cell genes vimentin, glutamine synthetase, retinaldehyde binding protein 1 (Rlbp1/CRALBP), GLAST, and cyclin D3. There were some differences in gene expression patterns between the ImM10 and C57M10 Müller cell lines. Primers for glial fibrillary acidic protein (Gfap) amplified a faint band from reverse-transcribed RNA from ImM10 cells, but no visible product was amplified from C57M10 cells. Although transcripts for carbonic anhydrase 14 were detected in C57M10 cells (passage 5), no expression was detected in ImM10 cells or in later passages of C57M10 cells (passage 20).

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**FIGURE 5.** Immunostaining and quantitative RT-PCR analysis of gene expression in ImM10 cells cultured in GM with 50 U/ml IFNγ at 33°C. In each row, the first panel shows antibody staining, the second panel shows 5′, 6-diamidino-2-phenylindole (DAPI) staining of nuclei, and the third panel shows the overlay of the first two images. (A–C) Gfap antibody staining, (D–F) Vimentin, (G–I) Glutamine synthetase (GS), (J–L) Cellular retinaldehyde binding protein (CRALBP), (M–O) Cyclin D3. Scale bar, 50 μm for all panels. (P) Ethidium bromide-stained agarose gels showing amplicons from quantitative RT-PCR analysis of (I) ImM10 cells, C57M10 cells at (M₂) passage 5 and (M₉) passage 20, and [R] whole adult C57BL/6 mouse retina. (Vim); glutamine synthetase (Glu1); glial fibrillary acidic protein (Gfap); cellular retinaldehyde binding protein (Rlbp1); calbindin (Calb1); medium wavelength cone opsins (Opn1mw); cellular retinaldehyde receptor mGluR6 (Grm6). POU domain, class III homeobox transcription factor 3 (Pou4f3/Brn3c); melanopsin (Opn4); calcium-binding protein 5 (Cab5); carbonic anhydrase 14 (Car14); rhodopsin (Rho); inflammatory factor-1 (Iba1); cone-rod homeobox (Crx); acidic ribosomal phosphoprotein P0 (Rplp0) was used as normalizing gene.32

Immunostaining of ImM10 cells confirmed the RT-PCR results and revealed uniform expression of Müller glial proteins in all cells. Antibodies against glial glutamate aspartate transporter (GLAST) revealed a typical punctate pattern throughout the cytoplasm, with the most intense staining in the perinuclear region (Figs. 5A–C). Antibodies against the intermediate filament protein vimentin stained a characteristic network of filaments throughout the cytoplasm with strong perinuclear staining (Figs. 5D–F). Antibodies against glutamine synthetase labeled the entire cell, including processes and nuclei (Figs. 5G–I), similar to what is observed in histologic sections of the retina. Antibodies against cellular retinaldehyde binding protein (CRALBP) showed relatively diffuse cytoplasmic staining, with some perinuclear enrichment (Figs. 5J–L) as typical for Müller glia. Antibodies against cyclin D3 showed predominantly nuclear staining (Figs. 5M–O), characteristic of Müller glia in the adult retina.33

In addition to expressing genes characteristic of Müller glia, neither ImM10 nor C57M10 cell lines expressed genes characteristic of other differentiated retinal cell types (Fig. 5P). By RT-PCR, both cell lines were negative for genes expressed in photoreceptors (rhodopsin/Rho, medium wavelength cone opsins/Opn1mv, cone-rod homeobox/Crx), horizontal cells (calbindin/Calb1), bipolar cells (metabotropic glutamate receptor mGluR6/Grm6, calcium binding protein 5, Cab5), ganglion cells (Pou4f3/Brn3c), and photosensitive ganglion cells (melanopsin/Opn4). Both cell lines were also negative for the microglial marker Iba1. Antibodies against PAX2 showed no nuclear staining in ImM10 cells (Figs. 6A–F), though some cytoplasmic immunoreactivity was present. In contrast, strong nuclear staining was present in retinal astrocytes in the inner

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**FIGURE 6.** PAX2 immunostaining in ImM10 cells and mouse retina. In each row, the first panel shows PAX2 antibody staining, the second panel shows 5′, 6-diamidino-2-phenylindole (DAPI) staining of nuclei, and the third panel shows the overlay of the first two images. (A–C) PAX2 immunostaining of ImM10 cells cultured at 33°C shows cytoplasmic, but not nuclear, staining. (D–F) Higher magnification of boxed area in (A). Arrowheads: nuclei. (G, I, J) PAX2 immunostaining of adult C57BL/6 mouse retina shows nuclear staining in retinal astrocytes in the inner retina (arrowheads). (K–M) Higher magnification of boxed area in (G). Scale bars, 50 μm (A–F), 100 μm (G–M).
Expression of Retinal Stem Cell Genes in ImM10 Cell Line

Using RT-PCR, we showed that ImM10 cells expressed mRNA for Nestin and the transcription factors Pax6, Six3, and Hes1 (Fig. 7). We confirmed the expression of Nestin and PAX6 using immunohistochemistry (Figs. 7A–F). By immunostaining, ImM10 cells also expressed SOX2 (Figs. 7G–I). As expected, Pax6 and SOX2 were localized primarily in the nucleus. By RT-PCR, ImM10 cells expressed other genes characteristic of retinal progenitors, including the RNA-binding protein musashi-1 (Msi1), nucleostemin (Gnl3), and cyclin D1 (Cnd1) (Fig. 7). In contrast, secreted frizzled related protein 2 (Sfrp2), fibroblast growth factor 15 (Fgf15), and retinal and anterior neural fold homeobox (Rax) are expressed in retinal progenitors during development, but not in mature Müller glia.23,24 and expression of these three genes was not detected by RT-PCR in RNA isolated from ImM10 cells.

DISCUSSION

We report the isolation and characterization of two novel cell lines isolated from the P10 mouse retina: ImM10, a conditionally immortalized Müller glial cell line generated from cx458/+; HrboGFp/+ mice, and C57M10, a spontaneously immortalized cell line from C57BL/6 mice. Several mammalian Müller glial cell lines have been previously described, including a conditionally immortalized Müller glia from rat retina (TR-MUL5),25 nonconditional, SV40 Tag immortalized cell lines from rat (rMC-1),19 and spontaneously immortalized Müller glia from postmortem human retina.11,21 Untransformed mouse Müller glia have been reported to survive in vitro for at least 7 passages.36 However, this is the first report of a conditionally immortalized Müller glial cell line from mouse that maintains Müller glial characteristics for more than 50 passages in vitro. Moreover, consistent with recent evidence that Müller cells in vivo express retinal stem cell genes,35,37,38 the ImM10 cell line is the first with demonstrated expression of multiple retinal stem cell genes when cultured as adherent monolayers.

ImM10 and C57M10 cell lines show the typical morphology of cultured Müller glia: large adherent cells with large, diffusely staining nuclei containing prominent nucleoli and an epithelioid morphology with multiple broad processes with distinct lamellipodia.31 This is consistent with the morphology of primary and immortalized Müller glia from other species, including rat,12,19-39 mouse,36 and human.21 ImM10 and C57M10 cells express a combination of genes typical of mature Müller glia, including Vim, Rlbp1, Dkk3, Clu, and Glua1, and we found no significant differences in gene expression between ImM10 cells cultured under immortalizing or nonimmortalizing conditions. Carbonic anhydrase 14 (Car14), a gene typically expressed in Müller glia in vivo, was not detected in mRNA from ImM10 cells, whereas C57M10 cells expressed low levels of Car14 mRNA at early passages but not at later passages. Downregulation of proteins involved in glial physiological functions has previously been reported in cultured Müller glia and is thought to reflect adaptation to in vitro conditions.40,41 Thus, the absence of Car14 expression in ImM10 cells is likely associated with adaptation to in vitro culture rather than a consequence of conditional immortalization.

The ImM10 and C57M10 cell lines were not clonally derived, and, although we cannot completely exclude the possibility that they consist of mixed populations, no heterogeneity was observed in the patterns of immunostaining. However, both Müller glial and retinal astrocytes express many of the same genes and show similar morphologies in culture. CRALBP expression is characteristic of Müller glia and has been used in by other groups to identify cultured Müller glia.11,12,21,36,40-42 Although developing retinal astrocytes transiently express CRALBP as they migrate from the optic nerve head, they downregulate expression in the postnatal retina with the protein no longer detectable by P7,43 prior to the age when we isolated these cell lines. Consistent with a Müller glial identity,
$\textit{Rbp4}$ mRNA was expressed in both ImM10 and C57M10 cell lines, and immunostaining revealed uniform and robust expression of CRALBP protein in 100% of the ImM10 cells. Furthermore, ImM10 cells also uniformly expressed cyclin D3, which is specifically expressed by Müller glia in the mature retina.5

PAX2 is a marker of retinal astrocytes, and previous reports describing conditionally immortalized retinal astrocytes from the transgenic mice show distinct nuclear staining for PAX2.28 In ImM10 cells, PAX2 immunoreactivity was present but consistently showed cytoplasmic rather than nuclear localization (Fig. 6). Although cytoplasmic accumulation of PAX2 protein has been previously reported in mammalian gammad,34 the significance of cytoplasmic PAX2 is unclear. This may reflect non-specific staining because we have previously observed nonspecific staining of nerve fibers in mouse embryos using these antibodies.45 However, multiple dilutions of the PAX2 antibodies all resulted in similar staining patterns. Similarly, omission of the primary antibodies revealed no nonspecific staining by the secondary antibodies. Because ImM10 cells expressed low levels of Pax2 mRNA by qRT-PCR, we cannot exclude the possibility that the cytoplasmic staining may reflect the presence of PAX2 protein. PAX2 expression is not detected in Müller glia in the intact retinas of mammals, including mice, though a few PAX2-positive glial cells have been observed in the peripheral retinas in older dogs.46 In contrast, in the chick retina, FGF2 treatment followed by excitotoxic injury upregulates PAX2 expression in Müller glia that reenter the cell cycle, suggesting a potential role in the glial-stem cell switch in Müller glia.40 To our knowledge, expression patterns of PAX2 have not been reported for other Müller glial cell lines; therefore, it is not apparent whether a low level of cytoplasmic PAX2 is unique to this cell line or is a more general property of Müller glia in culture.

The low expression level of GFAP expression in ImM10 and C57M10 cells is also consistent with identification as Müller glia rather than as retinal astrocytes. In the retina, Müller glia express low levels of Gfap in the absence of injury or disease, whereas retinal astrocytes constitutively express high levels of Gfap.47 Consistent with this, the SAGE tag ratio of vimentin to Gfap in two human Müller glial cell lines was 164:148, by immunohistochemistry, human Müller glial cell lines21 and primary rat Müller glial cultures51 also expressed very low levels of GFAP. In contrast to ImM10 and C57M10 cells, mouse Müller cells (mMCI and mMCII) cultured from P5 to 12 mice express high levels of Gfap and low levels vimentin by RT-PCR.56 The rat rMC-1 cell line52 and some primary human Müller cell cultures49 show robust immunoreactivity for the GFAP protein. Although the basis for these differences is unclear, upregulation of GFAP is characteristic of Müller glial activation after injury and in disease.50-51 The rMC-1 cell line was generated from light-damaged retinas; therefore, injury-induced upregulation of Gfap before cell isolation is one possibility. Because only some of the previously characterized human Müller cell cultures have elevated Gfap expression, preexisting injury or disease in human donor eyes might have contributed to higher than expected Gfap expression in these particular cultures.49

ImM10 cells also express multiple genes characteristic of both embryonic retinal stem cells and Müller glia in vivo, including the transcriptional regulators Pax6, Hes1, and Sox2.35,37,38 However, ImM10 cells did not express Sfrp2, Fgf15, or Rax, which are characteristically expressed in early retinal progenitors. These results are consistent with detailed gene expression profiling of single Müller glial cells that showed coexpression of Pax6, Hes1, and Sox2 in ImM10 cells but no significant expression of Sfrp2 or Rax.55 Although overexpression of Rax can promote differentiation of Müller glia,55 activity of the Rax promoter in Müller glia is downregulated as Müller glia differentiate53 and is absent in Müller glia in the mature retina.54 Sfrp2 and Fgf15 are also downregulated as differentiation proceeds and, by SAGE tag counts and in situ hybridization, both Sfrp2 and Fgf15 are undetectable in the adult retina.55 The absence of Sfrp2, Fgf15, and Rax expression in ImM10 cells appears to be consistent with their identification as Müller glia, despite the expression of multiple genes characteristic of retinal stem cells.

Expression of retinal stem cell-associated genes in ImM10 cells differs from the previously described mMCI and mMCII mouse Müller cells,56 which expressed only low levels of retinal stem cell genes when cultured as adherent cultures. Interestingly, mMCI cells upregulate nestin when induced to generate neurospheres by culturing in serum-free, FGF2/EGF-supplemented medium.56 Similarly, after growth factor stimulation, human MIO-M1 cells11 and primary Müller glia from rat15 and mouse retinas54 generate neurospheres that express stem cell genes, including Pax6, Sox2, and nestin. However, many published reports do not describe gene expression patterns in Müller glial cultures grown under nonneurosphere conditions. Thus, it is unclear whether the expression of retinal stem cell genes in neurospheres generated using other Müller cell line represents de novo upregulation or persistent expression of genes that were expressed before neurosphere induction.

Multiple immortalized retinal cell lines characterized as neuronal in origin have been generated from embryonic or fetal tissue.55-57 In contrast, few have been generated from postmitotic retinal neurons, suggesting that retinal progenitors are more amenable to immortalization. We observed that postmitotic neurons from the transgenic mice survived in mixed cultures. However, cells with neuronal morphologies did not incorporate BrdU and did not increase in number over time in culture, even with the induction of Tag expression. Müller glial cell lines from multiple species have been generated from a variety of ages by both spontaneous immortalization and overexpression of viral oncogenes.11,19,21,24-48 This likely reflects the inherent proliferative ability of Müller glia in vivo and potentially the limited stem cell-like properties of some Müller glia, both in vivo and in vitro.10

We conclude that the morphologic and gene expression profiles of the ImM10 and C57M10 cell lines are consistent with their identification as Müller glia. These mouse cell lines will facilitate research on the physiology and function of Müller glia and will complement the numerous genetic mouse models of retinal disease in use. In light of the growing evidence that some Müller glia may have stemlike characteristics both in vivo and in vitro, evaluation of the neurogenic capacity of ImM10 cells is under way.

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