JCI The Journal of Clinical Investigation

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J Clin Invest. 2016;126(8):3104-3116. https://doi.org/10.1172/JCI85193.

Research Article

Vision impairments and blindness caused by retinitis pigmentosa result from severe neurodegeneration that leads to a loss of photoreceptors, the specialized light-sensitive neurons that enable vision. Although the mammalian nervous system is unable to replace neurons lost due to degeneration, therapeutic approaches to reprogram resident glial cells to replace retinal neurons have been proposed. Here, we demonstrate that retinal Müller glia can be reprogrammed in vivo into retinal precursors that then differentiate into photoreceptors. We transplanted hematopoietic stem and progenitor cells (HSPCs) into retinas affected by photoreceptor degeneration and observed spontaneous cell fusion events between Müller glia and the transplanted cells. Activation of Wnt signaling in the transplanted HSPCs enhanced survival and proliferation of Müller-HSPC hybrids as well as their reprogramming into intermediate photoreceptor precursors. This suggests that Wnt signaling drives the reprogrammed cells toward a photoreceptor progenitor fate. Finally, Müller-HSPC hybrids differentiated into photoreceptors. Transplantation of HSPCs with activated Wnt functionally rescued the retinal degeneration phenotype in *rd10* mice, a model for inherited retinitis pigmentosa. Together, these results suggest that photoreceptors can be generated by reprogramming Müller glia and that this approach may have potential as a strategy for reversing retinal degeneration.

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Reprogramming Müller glia via in vivo cell fusion regenerates murine photoreceptors

Daniela Sanges,¹ Giacoma Simonte,¹ Umberto Di Vicino,¹ Neus Romo,¹ Isabel Pinilla,² Marta Nicolás,³ and Maria Pia Cosma¹,⁴

¹Center for Genomic Regulation (CRG), The Barcelona Institute of Science and Technology, Barcelona, Spain. ²Aragon Health Science Institute and University of Zaragoza, Zaragoza, Spain. ³Ferrer International SA, Barcelona, Spain. ⁴Institució Catalana de Recerca i Estudis Avançats (ICREA), Barcelona, Spain.

Vision impairments and blindness caused by retinitis pigmentosa result from severe neurodegeneration that leads to a loss of photoreceptors, the specialized light-sensitive neurons that enable vision. Although the mammalian nervous system is unable to replace neurons lost due to degeneration, therapeutic approaches to reprogram resident glial cells to replace retinal neurons have been proposed. Here, we demonstrate that retinal Müller glia can be reprogrammed in vivo into retinal precursors that then differentiate into photoreceptors. We transplanted hematopoietic stem and progenitor cells (HSPCs) into retinas affected by photoreceptor degeneration and observed spontaneous cell fusion events between Müller glia and the transplanted cells. Activation of Wnt signaling in the transplanted HSPCs enhanced survival and proliferation of Müller-HSPC hybrids as well as their reprogramming into intermediate photoreceptor precursors. This suggests that Wnt signaling drives the reprogrammed cells toward a photoreceptor progenitor fate. Finally, Müller-HSPC hybrids differentiated into photoreceptors. Transplantation of HSPCs with activated Wnt functionally rescued the retinal degeneration phenotype in rd10 mice, a model for inherited retinitis pigmentosa. Together, these results suggest that photoreceptors can be generated by reprogramming Müller glia and that this approach may have potential as a strategy for reversing retinal degeneration.

Introduction

Despite long-held dogma on the impossibility of changing the identity of differentiated cells, the expression of specific transcription factors, cell-cell fusion, and nuclear transfer experiments has shown the feasibility of reprogramming cellular identity toward both pluripotency and unrelated cell fates (1-5). Neurons are still considered among the most immutable cell types; nevertheless, recent studies have demonstrated that resident nonneuronal cells can be reprogrammed into induced neurons in vivo (6). This suggests that self-repair of nervous tissue can be promoted by inducing the fate conversion of resident cells into the desired cell type. Glial cells show the potential to divide and are very abundant in nervous tissue; therefore, they have been proposed as the ideal candidate cell type to generate new neurons. Glial cells can be turned into morphologically identifiable neurons in adult striatum upon expression of a cocktail of transcription factors (7) or just by the overexpression of Sox2, passing through an intermediate neuroblast stage (8).

In vivo conversion of glial cells into neurons represents a self-healing mechanism to regenerate nervous tissue in some species. Müller glia (MG) is the major glial cell type in the retina,

Conflict of interest: D. Sanges and M.P. Cosma hold a patent (PCT/EP2012/065327) related to the technology described in this manuscript that was licensed to Ferrer International SA. M.P. Cosma received licensing fees from Ferrer International SA for a different patent (PCT/EP2009/051512). D. Sanges and M.P. Cosma received research grants from Ferrer International. M.N. Farrés is directly employed at Ferrer International, but has no other financial interest.

Submitted: October 22, 2015; Accepted: May 24, 2016.

Reference information: *J Clin Invest*. 2016;126(8):3104–3116. doi:10.1172/JCI85193.

is common to all vertebrates, and normally contributes to retinal structure and homeostasis (9). After an injury, zebrafish MG dedifferentiates and generates proliferating multipotent neural progenitors, which in turn exit the cell cycle and redifferentiate, replacing all lost retinal cell types with sight restoration (10-12). In mammals, MG normally responds to damage by becoming reactive and undergoing gliosis (13, 14). However, some studies have indicated that a limited numbers of mouse and rat MG can reenter the cell cycle and be converted into retinal neurons after retinal injury and upon stimulus induction or expression of specific factors (15-18). These early studies made some pioneering observations; however, direct conversion of MG into neurons can be definitely concluded only with lineage-tracing experiments (19). Overall, the efficiency of the reported conversion was low or was not quantified, and importantly, electrophysiological functionality of the neurons obtained was not investigated. Nevertheless, these results suggest that the barrier that blocks MG reprogramming can also potentially be overcome in mammals.

Interestingly, the activation of Wnt/ β -catenin signaling can mediate proliferation of MG in the injured mammalian retina (16). Wnt/ β -catenin signaling strongly enhances the efficiency of reprogramming of somatic cells upon cell fusion with pluripotent stem cells (20, 21). Transplanted BM-derived cells (BMDCs) can fuse with somatic cells and generate hybrids, which in turn can differentiate in myocytes (22), hepatocytes (23), neurons (24, 25), and other cell types in vivo (26, 27). We recently showed that, upon drug-induced ganglion cell (GC) damage, transplanted Wnt-activated hematopoietic stem and progenitor cells (HSPCs) can fuse with retinal cells. The newly formed hybrids undergo reprogramming and differentiate into ganglion and amacrine neurons (28).

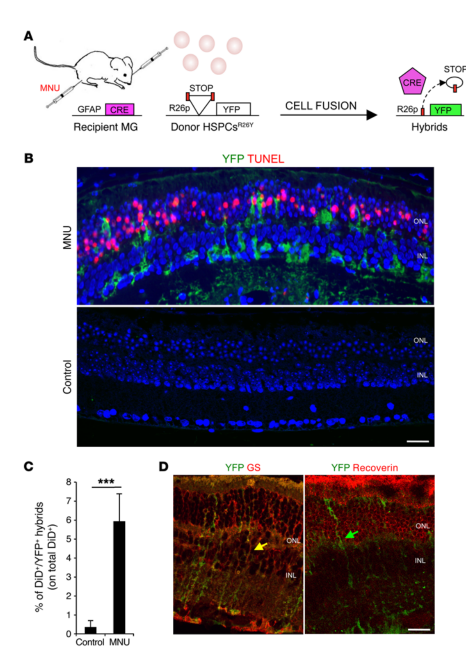


Figure 1. Transplanted HSPCs fuse with MG upon photoreceptor damage. (A) Schematic representation of the experimental plan. Cell fusion between HSPCs isolated from LoxP-STOP-LoxP-YFP donor mice (R26Y) and recipient Gfap-Cre MG cells leads to excision of the floxed stop codon and, in turn, to the expression of YFP. (B) Representative coimmunostaining of YFP+ hybrids (green) and TUNEL+ (red) apoptotic photoreceptors on retinal sections harvested from MNU-damaged or healthy (control) Gfap-Cre eyes 12 hours after subretinal transplantation of HSPCsR26Y. YFP+ hybrids (green) derived from cell fusion are detected in MNU-damaged retinas, which show TUNEL* photoreceptors in the ONL, but not in the undamaged eyes (control). Nuclei were counterstained with DAPI (blue). Scale bar: 20 μ m. n = 3. (C) Statistical analysis of the percentage of DiD+YFP+ hybrids evaluated on the total amount of DiD-labeled HSPCsR26Y detected by FACS analysis in MNU-damaged or healthy (control) Gfap-Cre retinas 24 hours after transplantation. Data are represented as mean ± SD of 3 independent experiments. n = 3. ***P < 0.0001by unpaired Student's t test. (D) Representative immunostaining of YFP+ hybrids (green) also positive for the MG marker (GS, red; yellow arrows) but not for the photoreceptor marker recoverin (red; green arrow) detected 24 hours after transplantation of HSPCsR26Y in MNU-damaged *Gfap-Cre* retinas. Scale bar: 20 μ m. n = 3.

By taking into account all of these previous observations, we here hypothesize that MG can be reprogrammed in vivo via cell fusion and become a source of new photoreceptors. In the present study, we demonstrate that Wnt-activated HSPCs transplanted in the retina of mouse models of genetic and drug-induced photoreceptor degeneration can fuse with MG with high efficiency. We found cell fusion to be the functional way through which MG reenters the cell cycle, proliferates, is reprogrammed back to a pluripotent/neural progenitor-like state, and finally differentiates into photoreceptors. Electrophysiological rescue and retinal regeneration are observed, which suggest that MG reprogramming and the subsequent differentiation into photoreceptors via cell fusion is a possible strategy for treatment of retinitis pigmentosa.

Results

MG cells fuse with transplanted HSPCs upon photoreceptor degeneration. We aimed to determine whether transplanted HSPCs can

fuse with MG in a drug-induced mouse model of photoreceptor degeneration. We used Gfap-Cre recipient mice (expressing Cre recombinase under the long form of the Gfap promoter), which restrict expression of Cre to MG (19, 29). Indeed, by using the lineage-tracing model that expresses yellow fluorescent protein (YFP) to track MG, we confirmed the exclusive colocalization of YFP with the MG marker glutamine synthase (GS) and not with PKC- α (a marker of bipolar cells), recoverin (a marker of photoreceptors), calbindin (a marker of horizontal cells), calretinin (a marker of amacrine and GCs), or neurofilament, heavy polypeptide (NEFH) and β-tubulin III (markers of GCs) (Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/ JCI85193DS1). We treated Gfap-Cre recipient mice with N-methyl-N-nitroso urea (MNU), which induces progressive photoreceptor cell death (TUNEL+) within 24 hours (Figure 1, A and B, and Supplemental Figure 1, B and C; compare WT with MNU/PBS) and complete degeneration of the outer nuclear layer (ONL), where

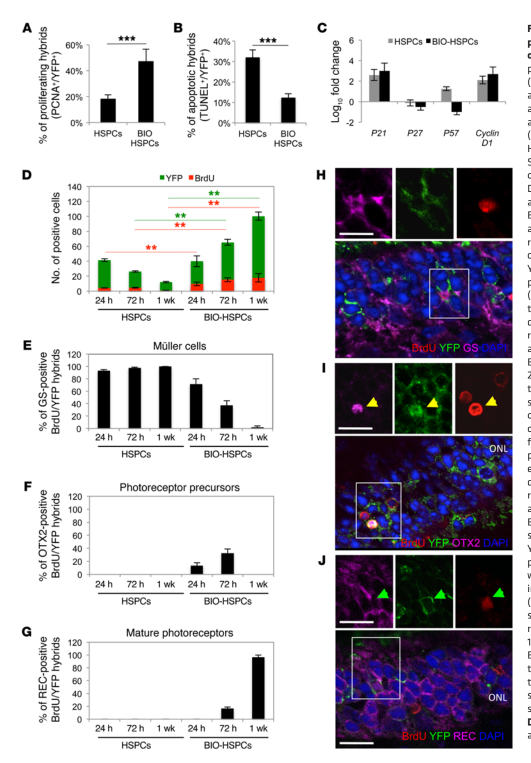
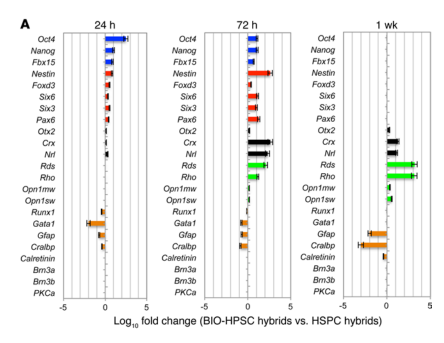


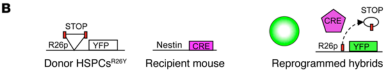
Figure 2. Activation of Wnt signaling promotes proliferation and survival of hybrids. (A and B) Percentages of proliferating (PCNA+, A) or apoptotic (TUNEL*, B) YFP* hybrids on the total amount of YFP+ hybrids in MNU-damaged Gfap-Cre retinas 24 hours after transplantation of untreated (HSPCs) or BIO-treated (BIO-HSPCs) $\mathsf{HSPCs}^\mathsf{R26Y}$. ***P < 0.0001, unpaired Student's t test. (C) qPCR analysis of cell-cycle genes on FACS-sorted DiD+YFP+ hybrids harvested 24 hours after transplantation of untreated or BIO-treated HSPCsR26Y in MNU-damaged Gfap-Cre retinas. Data are represented as mean ± SD of log₁₀ fold changes of gene expression in DiD+ YFP+ hybrids with respect to DiD+YFP+ population-depleted retinas. n = 3. (D) Total YFP+ hybrids (green bars) that also incorporated BrdU (red bars) detected in MNU-damaged Gfap-Cre retinas 24 hours, 72 hours, and 1 week after transplantation of untreated or BIO-treated HSPCsR26Y. **P < 0.001, 2-way ANOVA and Bonferroni's posttest. Red and green lines show the statistical significance of green (YFP) or red (BrdU) bars. (E-G) Percentages of YFP*BrdU* hybrids also positive for GS (E, Müller cells), OTX2 (F, photoreceptor progenitors), or recoverin (G, REC, mature photoreceptors) detected in MNU-damaged Gfap-Cre retinas 24 hours, 72 hours, and 1 week after transplantation of untreated or BIO-treated HSPCsR26Y. (H-J) Representative immunostainings of double YFP+ (green)/BrdU+ (red) hybrids also positive for either GS (magenta in H, white arrowheads), OTX2 (magenta in I, yellow arrowheads), or recoverin (magenta in I, green arrowheads) stainings in MNU-damaged Gfap-Cre retinas 24 hours (H), 72 hours (I), and 1 week (J) after transplantation of BIO-treated HSPCs. Nuclei were counterstained with DAPI (blue). Images on the top show higher magnification and single channels of areas in the white squares. Scale bars: 20 μm. (A, B, and D-G) Values in graphs are represented as mean \pm SD (n = 9).

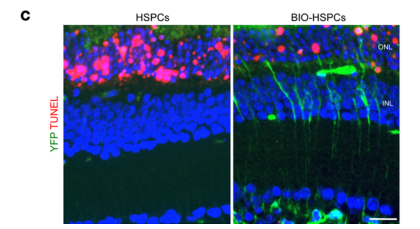
photoreceptors are localized, after 7 days (30, 31). Twenty-four hours after MNU damage, we injected HSPCs isolated from LoxP-STOP-LoxP-YFP (R26Y) transgenic donor mice into the subretinal space of *Gfap-Cre* recipient eyes, close to the layer of damaged photoreceptors. *Gfap-Cre* mice not treated with MNU were also transplanted as controls. Twelve hours after HSPCs^{R26Y} transplantation, we detected many YFP⁺ cells close to the site of the injection; this was due to the stop codon excision by Cre after fusion of HSPCs^{R26Y} with Müller cells from *Gfap-Cre* mice (MG^{Gfap-Cre}) (Figure 1, A

and B, and Supplemental Figure 1D). The YFP⁺ hybrids localized radially across the thickness of the retina, which is typical of MG morphology. YFP⁺ hybrids were detected upon transplantation in MNU-damaged retinas, where many photoreceptors were apoptotic (Figure 1B, TUNEL⁺), but they were not appreciable in sections where HSPCs^{R26Y} were transplanted into nondamaged *Gfap-Cre* eyes, which suggests that photoreceptor cell death is necessary to induce cell fusion (Figure 1B). A mean of 5.9% ± 1.5% of the total transplanted DiD-labeled HSPCs^{R26Y} was also YFP⁺ when inject-

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ed into MNU-damaged eyes (DiD+YFP+ hybrids; Figure 1C and Supplemental Figure 1, E and H; *Gfap-Cre*), which corresponds to the frequency of cell-fusion events. However, a few YFP+DiD-hybrids (not included in the analysis) were also found; thus, we cannot exclude an underestimation of fusion frequency. In contrast, we observed very few hybrids (detectable only by FACS analysis and likely due to the mechanical damage induced by the injection) when DiD-labeled HSPCsR26Y was transplanted into nondamaged (control) *Gfap-Cre* recipient eyes (Figure 1C and Supplemental Figure 1H; *Gfap-Cre*, see below), thus confirming that cell fusion between HSPCs and MG can occur in the presence of tissue damage.

Cell fusion was also confirmed by transplanting DiD-labeled HSPCs expressing *Cre* under the BM-specific promoter *Vav* (referred to here as HSPCs *Vav-Cre*) into MNU-damaged and nondamaged R26Y

Figure 3. Activation of Wnt signaling promotes reprogramming of hybrids. (A) Gene expression analysis of pluripotent (blue bars), neural progenitor (red bars), photoreceptor progenitor (black bars), mature photoreceptor (green bars), and differentiated cell (orange bars) markers in DiD+YFP+ hybrids. The hybrids were FACS sorted 24 hours, 72 hours, and 1 week after transplantation of untreated or BIO-treated HSPCsR26Y in MNU-damaged Gfap-Cre retinas. Data represent the mean of log, fold changes ± SEM of gene expression detected in hybrids obtained upon fusion with BIO-treated DiD-HSPCs with respect to hybrids obtained upon fusion with untreated DiD-HSPCs. n = 3. (B) Schematic representation of the experimental plan to detect dedifferentiation of retinal cells upon fusion. Reactivation of the nestin promoter in retinal cells fused with transplanted HSPCsR26Y leads to Cre expression and the consequent formation of YFP+ hybrids. (C) Representative immunostaining of YFP+ hybrids (green) and TUNEL+ apoptotic photoreceptors (red) on retinal sections obtained from MNU-damaged nestin-Cre retinas 24 hours after transplantation of untreated or BIO-treated HSPCsR26Y. Nuclei were counterstained with DAPI (blue). Scale bar: 20 µm.

eyes, where a mean of $7.5\% \pm 1.2\%$ DiD*YFP* hybrids was found 24 hours after transplantation, but only in the damaged group (Supplemental Figure 1, F-H; HSPCs/R26Y).

Given that we obtained approximately the same percentages of hybrids after transplantation into R26Y or into Gfap-Cre recipient mice (Supplemental Figure 1H), we hypothesized that HSPCsVav-Cre fused mainly with MG upon photoreceptor damage. Furthermore, YFP+ hybrids were immunoreactive to the MG marker GS, but not to the photoreceptor marker recoverin (Figure 1D), which further suggests fusion of HSPCs with MG. Finally, almost no YFP+ hybrids were identified in both damaged and undamaged retinas harvested 24 hours after transplantation of DiD-labeled HPSCs from R26Y mice (HSPCsR26Y) either in recipient mice carrying Cre under the photoreceptor-specific rhodopsin promoter (Rho-Cre; ref. 32) or in recipient mice car-

rying Cre under the ganglion- and amacrine-specific calretinin promoter (*Calr-Cre*; ref. 33), thus excluding major fusion of transplanted HSPCs with other retinal cell types (Supplemental Figure 1, H-J). Taken together, these results indicate that HSPCs can fuse almost exclusively with MG when transplanted in the subretinal space of an MNU-induced mouse model of photoreceptor degeneration.

MG reenters the cell cycle upon fusion with HSPCs through a Wnt-dependent process. MG can reenter the cell cycle after retinal damage (14). Furthermore, we previously demonstrated that Wnt signaling enhances the proliferation of hybrids formed in vitro and in vivo (20, 28). Thus, we investigated whether MG-derived hybrids can reenter the cell cycle upon Wnt activation. We transplanted into the eyes of MNU-damaged *Gfap-Cre* mice either HSPCs^{R26Y}, where the Wnt signaling pathway was preactivated by the GSK-3 inhibitor 6-bromoindirubin-3'-oxime (BIO) (Sup-

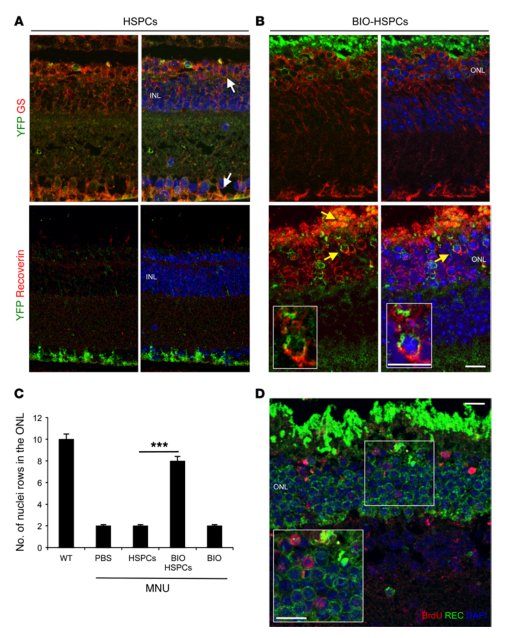


Figure 4. MG reprogrammed upon fusion with Wnt-activated HSPCs generates new photoreceptors. (A and B) Representative immunostainings of YFP+ (green) hybrids also immunoreactive for either GS (red, white arrows) or recoverin (red, yellow arrows) detected in MNU-damaged Gfap-Cre retinas 1 week after transplantation of untreated (A) or BIO-treated (B) HSPCsR26Y. Nuclei were counterstained with DAPI (blue). n = 3. (C) Statistical analysis of the number of photoreceptor nuclear rows in the ONL of healthy (WT) or MNU-damaged eyes 1 week after treatment with either PBS or untreated (HSPCs) or BIO-treated HSPCs (BIO-HSPCs) or BIO alone. Data are represented as mean ± SD counted in 3 different sections spanning the site of the injection for each mouse. n = 9. ***P < 0.0001, unpaired Student's t test. (D) Representative immunodetection of BrdU-positive cells (red) also positive for the photoreceptor marker recoverin (green) in the ONL of MNU-damaged Gfap-Cre retinas 1 week after transplantation of BIO-treated HSPCsR26Y. Nuclei were counterstained with DAPI (blue). Insert represents a higher magnification of the region in the white square. (n = 3). Scale bars: 20 µm.

plemental Figure 2A), or nonactivated HSPCsR26Y. We found YFP+ hybrids that were also positive for the proliferation marker proliferating cell nuclear antigen (PCNA) in sections of retinas harvested 24 hours after transplantation of HSPCsR26Y (Figure 2A and Supplemental Figure 2B, HSPCs). However, the majority of these embarked upon apoptosis, as more than 35% of YFP+ cells were also positive for TUNEL staining (Figure 2B and Supplemental Figure 2, C and D, HSPCs). In contrast, more PCNA+ hybrids were detected 24 hours after transplantation of BIO-treated HSPCs (Figure 2A and Supplemental Figure 2B, BIO-HSPCs) and only a few YFP+ hybrids underwent apoptosis (Figure 2B and Supplemental Figure 2, C and D, BIO-HSPCs). These results suggest that preactivation of Wnt signaling in HSPCs enhances the survival and proliferation of hybrids formed upon fusion with MG. Of note, Wnt activation did not increase the fusion efficiency (Supplemental Figure 2E; compare HSPCs with BIO-HSPCs).

We then investigated the expression levels of different cell-cycle regulators in hybrids that were FACS sorted 24 hours after transplantation of BIO-treated or untreated HSPCsR26Y in MNU-damaged Gfap-Cre eyes. Cyclin D1, which is expressed in mitotic retinal progenitor cells and is required for progenitor cell proliferation (34, 35), was strongly upregulated in hybrids formed after transplantation of both BIO-treated and untreated HSPCs (Figure 2C). We also found upregulation of p21^(Cip1/WAF1), which is correlated with neural stem cell maintenance and tissue regeneration (36). Interestingly, the cyclin kinase inhibitors $p27^{kip1}$, which is downregulated in activated MG (37), and $p57^{kip2}$, which is associated with cell-cycle exit of retinal progenitor cells (14), were both downregulated in hybrids obtained after transplantation of BIO-treated HSPCs (Figure 2C), which suggests the prolonged proliferative potential of the BIO hybrids. These observations indicate that some cell-cycle genes are

perturbed in both types of hybrids; however, the activation of HSPCs with BIO provides a proliferative advantage.

To further analyze the fate of the proliferating hybrids, we injected BrdU twice every 12 hours in MNU-damaged Gfap-Cre mice that were transplanted with untreated or BIO-treated HSPCs^{R26Y}. Thus, we analyzed YFP+ hybrids that incorporated BrdU 24 hours, 72 hours, and 1 week after transplantation. Although the total number of YFP+ hybrids detected in each retinal section was similar 24 hours after transplantation of either untreated or BIO-treated HPSCs, Wnt signaling activation resulted in more hybrids that incorporated BrdU (Figure 2D), which indicates their proliferative advantage. Interestingly, 72 hours and 1 week after transplantation of untreated HSPCs, the total number of YFP+ hybrids was reduced, which suggests that hybrids that did not proliferate (i.e., YFP+BrdU- cells) possibly died (Figure 2D). In contrast, the number of double-positive BrdU+YFP+ cells as well as the number of total YFP+ hybrids increased 72 hours and 1 week after transplantation of BIO-treated HSPCs, which suggests that upon activation of Wnt signaling, some proliferating BrdU+YFP+ hybrids had moved into and through a new M phase and survived. Moreover, we also detected a higher number of YFP+BrdUcells 72 hours and 1 week after transplantation of BIO-HSPCs, with respect to those counted 24 hours after transplantation (Figure 2D), which suggests that either some proliferating hybrids did not incorporate BrdU or that they underwent multiple divisions and thus lost the BrdU. Together, these results suggest that activation of Wnt signaling in transplanted HSPCs promotes proliferation and survival of hybrids formed after fusion with MG.

MG is reprogrammed in vivo after fusion with Wnt-activated HSPCs, and the hybrids differentiate into photoreceptors. We aimed to determine whether MG is reprogrammed in vivo upon fusion with Wnt-activated HSPCs. Interestingly, we observed that, while the majority of the proliferating YFP+BrdU+ hybrids were positive for the MG marker GS 24 hours, 72 hours, and 1 week after transplantation of untreated HSPCs, the percentage of YFP+BrdU+ hybrids positive for GS was drastically reduced 72 hours and 1 week after transplantation of BIO-HSPCs (Figure 2, E and H, see below) and the latter were mainly positive for the expression of the photoreceptor progenitor marker orthodenticle homeobox 2 (OTX2) at 72 hours after transplantation (Figure 2, F and I, see below). Furthermore, we noted that the photoreceptor marker recoverin started to be expressed in the YFP+BrdU+ hybrids 72 hours after transplantation of BIO-HSPCs and that all the Brdu⁺ YFP+ cells expressed recoverin 1 week after transplantation (Figure 2, G and J, see below). Of note, the proliferating hybrids (i.e., double YFP+BrdU+ hybrids) were mainly localized in the ONL already at 24 hours (Figure 2H), when they expressed MG markers, and consequently, at late time points (Figure 2J, see below), when they differentiated toward photoreceptors, which indicates their movement toward the ONL. These results suggest that Wnt signaling promotes the reprogramming of the proliferating hybrids toward a photoreceptor progenitor fate.

To further strengthen these results, DiD+YFP+hybrids were FACS sorted 24 hours, 72 hours, and 1 week after transplantation of DiD-labeled BIO-HSPCs^{R26Y} or untreated DiD-HSPCs^{R26Y} in MNU-damaged *Gfap-Cre* eyes. The gene expression profiles were analyzed (Figure 3A). Twenty-four hours after transplantation, pluripotent markers (*Oct4*, *Nanog*, and *Fbx15*) were upregulated in the hybrids obtained

after fusion of BIO-HSPCs and progressively silenced 1 week after transplantation. The Wnt target Axin2 was also upregulated in the hybrids 24 hours after transplantation of BIO-HSPCs, which indicates activation of Wnt signaling (Supplemental Figure 3A). In contrast, neural progenitor markers (Nestin, Foxd3, Six6, Six3, and Pax6) were upregulated up to 72 hours after transplantation, while they were silenced in 1 week. The postmitotic photoreceptor progenitors (Otx2, Crx, and Nrl) were upregulated up to 1 week after transplantation. Finally, the mature rod photoreceptor markers peripherin (Rds) and rhodopsin (Rho) were upregulated already 72 hours after transplantation, but were very highly expressed within 1 week. Low upregulation of the cone photoreceptor markers Opn1mw and Opn1sw was also detected. However, no upregulation of calretinin (Calr, expressed in ganglion and amacrine cells) nor of Brn3a, Brn3b (markers of GCs) nor of *Pkca* (a marker of bipolar cells), was detected at any time point, which suggests that hybrids were not differentiated toward these retinal cell fates. The HSPC (Runx1, Gata1) and MG (Gfap, Cralbp) markers were downregulated already 24 hours after transplantation (Figure 3A). For the hybrids obtained after the fusion of untreated HSPCs, we did not observe upregulation of the above-mentioned markers; instead, HSPC (Runx1, Gata1) and MG (Gfap, Cralbp) genes were expressed (Supplemental Figure 3B). These data indicated that the hybrids formed after fusion of BIO-HSPCs underwent reprogramming to a pluripotent/neural progenitor-like fate and then committed toward photoreceptors in vivo.

To confirm this conclusion, we transplanted BIO-treated or untreated HSPCs^{R26Y} in the eyes of MNU-damaged nestin-*Cre* recipient mice (Figure 3B). We observed YFP+ hybrids only after transplantation of BIO-HSPCs (Figure 3C), which indicates that reactivation of nestin and reprogramming of the hybrids was dependent on Wnt activity. Furthermore, upon MNU treatment, in 24 hours, photoreceptors start to die (TUNEL+) in retinas transplanted with either untreated HSPCs, BIO alone, or vehicle alone (PBS). Instead, a much-reduced number of TUNEL+ photoreceptors was seen in BIO-HSPC-transplanted retinas and especially in the proximity of YFP+ hybrids (Figure 3C and Supplemental Figure 1, B and C).

We then verified on sections whether Müller-derived hybrids can generate photoreceptors. We performed double staining of retinal sections 1 week after transplantation of BIO-treated or untreated HSPCs in MNU-damaged *Gfap-Cre* eyes (Figure 4, A and B). First, upon HSPC transplantation, we observed only 1 or 2 leftover nuclear layers in the ONL, and these were negative for the expression of recoverin (Figure 4, A and C, and Supplemental Figure 4A). The few YFP+ hybrids detected were positive for the MG marker GS (Figure 4A). In contrast, after transplantation of BIO-treated HSPCs, we found up to 8 nuclear layers in the ONL (Figure 4, B and C, and Supplemental Figure 4A), which included hybrids positive for both YFP and recoverin (i.e., photoreceptors) (Figure 4B).

Overall, these results show that MG can be efficiently reprogrammed into photoreceptors upon fusion with BIO-treated HSPCs, passing through a proliferating and more dedifferentiated intermediate.

Despite many photoreceptors derived from the proliferation and differentiation of reprogrammed MG cells (i.e., BrdU/Rec double-positive cells; Figure 4D), not all of the photoreceptors in

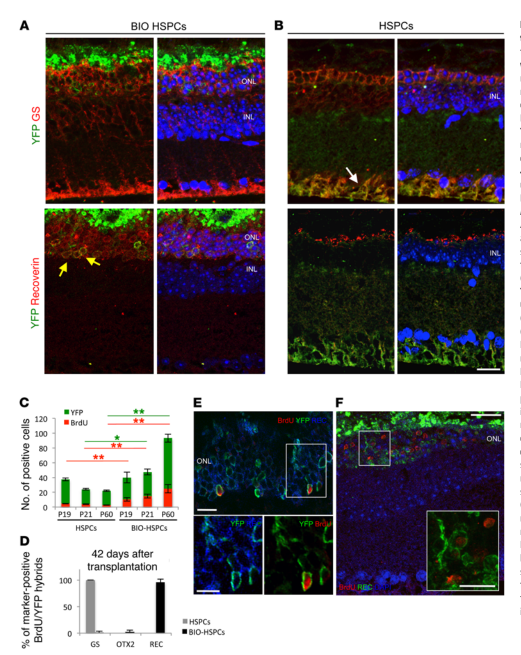


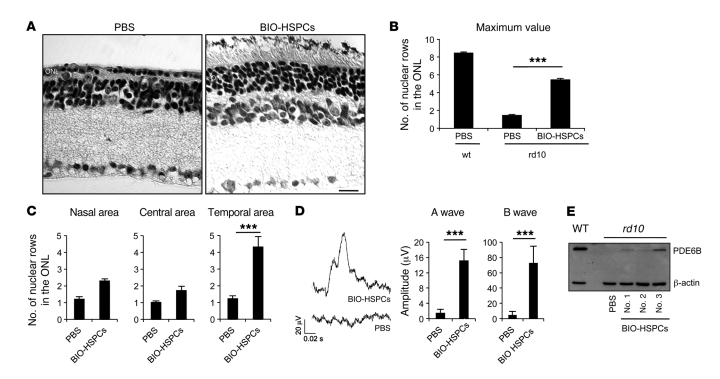
Figure 5. Regeneration of photoreceptors in rd10 mice. (A and B) Representative immunostainings of YFP+ (green) with either GS (red, white arrow) or recoverin (red, yellow arrows) in rd10R26Y retinas 42 days after transplantation of BIO-treated (A) or untreated (B) $\mathsf{HSPCs}^{\mathit{Vav-Cre}}.\ n=3.$ (C) Total number of YFP+ hybrids (green bars) that incorporated BrdU (red bars) detected in $rd10^{\rm R26Y}$ eyes 24 hours (P19), 72 hours (P21), and 42 days (P60) after transplantation of untreated or BIO-treated HSPCs Vav-Cre. Data are represented as mean ± SD. n = 9. **P < 0.001; *P < 0.01, 2-wayANOVA and Bonferroni's post-test. Red and green lines show the statistical significance of the difference between the respective green (YFP) and red (BrdU) bars. (D) Percentages of double YFP*BrdU* hybrids also positive for GS, OTX2, or recoverin stainings 42 days (P60) after transplantation of untreated (gray bars) or BIO-treated (black bars) HSPCsVav-Cre in rd10R26Y retinas. Data are represented as mean \pm SD. n = 9. (E) Representative immunodetection of BrdU+ cells (red) also positive for the photoreceptor marker recoverin (green) in the ONL of MNU-damaged rd10R26Y retinas 42 days after transplantation of BIO-treated HSPCsVav-Cre. Nuclei were counterstained with DAPI (blue). Insert shows higher magnification of the region in the white square. (F) Representative confocal image of YFP+ hybrids (green) also positive for BrdU (red) and recoverin (blue) in the ONL of rd10R26Y retinas 42 days after transplantation of BIO-treated HSPCs Vav-Cre. Images at the bottom represent enlargements of the same YFP+ hybrid (green) also positive for recoverin (blue in the left panel) and for BrdU (red in the right panel) stainings. n = 3. Scale bar: 20 μ m.

BIO-HSPC transplanted retinas were derived from the hybrids, as not all of them were positive for YFP (Figure 4B) or for BrdU (Figure 4D), which suggests that newly generated photoreceptors can contribute overall to blocking the death of the neighboring ones.

In vivo reprogramming of MG upon fusion with Wnt-activated HSPCs can rescue retinal degeneration phenotypes. We next aimed to investigate the possible functional rescue of the retinal degeneration phenotype due to the photoreceptors derived from the hybrids formed by fusion of MG and BIO-treated HSPCs. Although MNU-induced photoreceptor cell death is a useful model for studying the effects of therapeutic approaches to treating retinal degeneration in the short term, this alkylating drug leads to extensive oxidative stress and induction of tumors within 1 month (38, 39). Thus, we determined whether MG-derived photoreceptors can functionally rescue the photoreceptor degeneration phenotype in

a type 10 retinal degeneration mouse model (rd10) over the long term. This is a model of inherited autosomal recessive retinitis pigmentosa, which carries a spontaneous mutation in the rod-specific β subunit of cyclic guanosine monophosphate–phosphodiesterase (cGMP-phosphodiesterase) (PDE6B). The rd10 mice show photoreceptor dysfunction starting from P18 (i.e., 18 days after birth), and full rod photoreceptor degeneration is followed by cone degeneration and remodeling of the inner retina by P40 (40).

First, we confirmed the presence of YFP*DiD* hybrids after transplantation of BIO-treated or untreated DiD-labeled HSPCs^{Vav-Cre} in the subretinal space of P18 *rd10* mice carrying the R26Y transgene (referred to as *rd10*^{R26Y}) when the degeneration started (Supplemental Figure 5, A and B). YFP*DiD* hybrids were not detected in transplanted WT control retinas, which indicated that *rd10* photoreceptor degeneration induced fusion of MG with transplanted



HSPCs (Supplemental Figure 5, A and B). To confirm fusion with MG, we also transplanted HSPCs^{Vav-Cre} in P18 *rd*10^{R26Y}; as expected, 24 hours after transplantation, the hybrids were immunoreactive to the MG marker GS, but not to the photoreceptor marker RHO (Supplemental Figure 5, C and D). The hybrids obtained upon fusion with BIO-treated HSPCs^{Vav-Cre} proliferated (Supplemental Figure 5E) and expressed pluripotent/neural progenitor markers (Supplemental Figure 5G, see below). In contrast, the YFP⁺ hybrids formed upon transplantation of untreated HSPCs^{Vav-Cre} underwent apoptosis (Supplemental Figure 5F) and were not reprogrammed (Supplemental Figure 5G).

Next, 42 days after transplantation, at P60, we analyzed the presence of hybrid-derived photoreceptors. Interestingly, in 2-month-old *rd10* mice transplanted with BIO-treated HPSCs isolated from *Vav-Cre/CAG-RFP* double-transgenic animals (where RFP indicates red fluorescent protein; HSPCs^{Vav-Cre/RFP}), many YFP+RFP+ hybrids were detected in the regions of the ONL, including a high number of nuclear rows (Supplemental Figure 5H). Indeed, 42 days (at P60) after transplantation of BIO-treated HSPCs^{Vav-Cre}, we found YFP+ hybrids that were positive for recoverin staining and negative to GS (Figure 5A). In contrast, as expected, few YFP+ hybrids were found in HSPC-transplanted *rd10* retinas of 2-month-old mice. When detectable in sections,

these hybrids were positive for the MG marker GS and negative for recoverin (Figure 5B). Moreover, while only 1 row of nuclei in the ONL was detected in *rd10* eyes transplanted with untreated HSPCs, a much higher number of photoreceptor nuclei was observed in mice transplanted with BIO-treated HSPCs (Figure 5, A and B).

We then performed cell-tracing experiments to investigate whether the hybrid-derived photoreceptors were generated from proliferative events. We followed BrdU+ hybrids after transplantation of BIO-treated and untreated HSPCs $^{Vav-Cre}$ in $rd10^{R26}$. Activation of Wnt signaling in transplanted HSPCs resulted in an increase in the proliferating hybrids at 24 hours (P19), 72 hours (P21), and 42 days (P60) after transplantation with respect to the corresponding times after HSPC transplantation (Figure 5C). In contrast, the YFP+ hybrids decreased after HSPC transplantation (Figure 5C), which suggests that they underwent cell death in the latter case. Brdu+YFP+ hybrids expressed GS after HSPC transplantation at 24 hours, 72 hours, and 42 days and therefore remained as MG (Figure 5D and Supplemental Figure 5, I and J). Instead, after BIO-HSPC transplantation, Brdu⁺YFP⁺ hybrids progressively silenced GS and expressed OTX2 and recoverin, which suggests their differentiation toward a photoreceptor cell fate (Figure 5D and Supplemental Figure 5, I and J). Remarkably, by tracing them, we found a large increase of hybrids positive for BrdU in P60 mice, which indicates that they efficiently proliferated and differentiated toward a mature photoreceptor cell fate (Figure 5, C, E, and F). Of note, not all of the photoreceptors were BrdU⁺ or YFP⁺, which suggests that in the *rd10* model, as in the MNU model, a regeneration process goes along with some preservation effect.

We then counted the number of nuclear photoreceptor rows across sections. Histological analysis of retinal sections from the control left eyes of 2-month-old *rd10* mice treated with PBS confirmed complete photoreceptor degeneration, with only 1 row of photoreceptor nuclei detectable in the ONL (Figure 6, A-C). Instead, up to 5 rows of photoreceptor nuclei were detected in the ONL of the right eyes of a group of mice transplanted with BIO-treated HSPCs, specifically in the temporal area where the cells were transplanted (Figure 6, A-C). Of note, no changes were detected in either the nuclear row numbers in the inner nuclear layer (INL) or in the number of GC nuclei, which suggests that treatment with BIO-HSPCs results specifically in the rescue of photoreceptor cell loss (Supplemental Figure 6, A-C).

To investigate whether functional rescue of the rd10 phenotype was achieved, we recorded scotopic electroretinographic (ERG) responses of rd10 dark-adapted right eyes transplanted with BIO-treated HSPCs. Left eyes treated with PBS were also recorded as controls. Remarkably, we observed that some of the retinas transplanted with BIO-treated HSPCs showed recordable A-waves and B-waves (Figure 6D), albeit these were lower than the WT levels (WT A-wave mean $[\mu V] = 416 \pm 77$ SD; WT B-wave mean $[\mu V] = 826 \pm 107$ SD; n = 4). The A-wave is generated from functional rod photoreceptors, while the B-wave is mainly derived from retinal cells making synapses with photoreceptors (41). Interestingly, the mice that showed an increase in the number of photoreceptor nuclear rows (Supplemental Figure 6D) also showed recordable A-waves and B-waves (Supplemental Figure 6E). Instead, the contralateral untransplanted eyes showed completely flat ERG responses (Figure 6D and Supplemental Figure 6E).

Similar results were obtained with the $rd10^{R26Y}$ mouse strain, where the retinas of mice displaying YFP⁺ hybrids also showed the highest number of rows and were positive for the ERG (Supplemental Figure 6, F and G).

Subretinal injection of P18 pups is a challenging chirurgical approach, and we therefore observed retinal regeneration in a subset of BIO-HSPC transplanted mice. Therefore, we monitored the outcome of the injection by optical coherence tomography (OCT) in a different group of *rd10*-transplanted mice. We confirmed that only the group of eyes successfully transplanted with BIO-treated HSPCs showed photoreceptor regeneration at the histological level (Supplemental Figure 6, A–C). In all these mice, the OCT generated a cataract, and it was not possible to record the ERG.

The ERG A-wave is normally flat in 2-month-old *rd10* mice because of the absence of PDE6B, which is necessary for the closure of cGMP-gated cationic rod channels (42). Therefore, we analyzed the levels of the PDE6B protein in retinas harvested from 2-month-old *rd10* mice transplanted with untreated or BIO-treated HSPCs at P18. WT retinas treated with PBS were also analyzed as controls. The PDE6B protein was completely absent in untreat-

ed *rd10* eyes, while it was present at variable levels in 3 different *rd10* eyes transplanted with BIO-treated HSPCs (Figure 6E). This suggests that WT PDE6B was expressed by the genome of the transplanted HSPCs within the hybrids, thereby rescuing the *rd10* phenotype.

In conclusion, our results demonstrate that, upon fusion with Wnt-activated HSPCs, MG can be efficiently reprogrammed and differentiated into functional photoreceptors in vivo and that this might represent a therapeutic strategy for rescuing retinal degeneration phenotypes.

Discussion

Retinitis pigmentosa is a family of inherited diseases that cause retinal degeneration due to photoreceptor cell death. The *rd10* mouse model reproduces one of the earliest onset and most aggressive forms of recessive retinitis pigmentosa, which is due to mutations in the *Pde6b* gene (43, 44). Very limited therapeutic approaches are currently available for the treatment of this blindness. Indeed, many efforts have been made to block or delay degeneration of photoreceptors (45, 46); however, to date, very limited success has been obtained in terms of their regeneration in mouse models of retinitis pigmentosa.

Here, we have shown that MG can be reprogrammed in vivo and, in turn, efficiently differentiated into photoreceptors via cell fusion with HSPCs in a drug-induced model of retinal degeneration and in rd10 mice, provided that Wnt signaling was previously activated in the transplanted cells. BrdU⁺ photoreceptors originated from the MG-derived hybrids were tracked over the long term, i.e., 42 days after transplantation of BIO-HSPCs in rd10 mice. On the other hand, we also clearly observed a protective effect of the resident photoreceptors upon transplantation of BIO-HSPCs, which can be due to hybrid-mediated indirect protection or to a direct effect mediated by BIO-HSPC-secreted factors. Therefore, the hybrid-derived photoreceptors likely blocked or delayed the overall retina degeneration; indeed, not all the surviving photoreceptors in the long term after transplantation were YFP positive. Some of them were resident photoreceptors that were probably protected from cell death. Our data also suggest that, while in the degenerated retinas MG appears to extend into the remodeled ONL, upon transplantation of BIO-HSPCs, the ONL is regenerated and MG staining appears in sections as in a WT retina.

We should also consider the possibility that MG phagocytoses BIO-HSPCs and that this could result in either regenerative or protective effect toward the ONL neurons. However, MG cells were shown to phagocytose apoptotic retinal cells during development (47) as well as apoptotic photoreceptors in degenerated mouse retinas (48, 49). Here, we observed expression of RFP and *Pde6b* long term after transplantation, suggesting that the genome derived from donor transplanted BIO-HSPCs is present and is expressed in the YFP⁺ neurons derived from MG; therefore, phagocytosis is unlikely to occur in our experimental setting. On the other hand, if some BIO-HSPC-transplanted cells are phagocytosed by MG, these will just die and will likely not be the cells that regenerate the photoreceptors.

What signaling controls regeneration in zebrafish and tadpoles (19, 50, 51). We have previously shown that ganglion and amacrine neurons in the mouse retina can be regenerated by the hybrids formed after Wnt-activated HSPCs (28). How is it possible that the hybrids can generate either photoreceptors, GCs, or amacrine neurons? It appears likely that this is dependent on the signaling pathway induced, on the type of damaged cells in need of being replenished, and also on the site of injection. Here, we transplanted HSPCs very close to the photoreceptor layer; we also attempted transplantation in the vitreous, but we did not observe regeneration. Furthermore, transplantation at late times, when the ONL was completely lost, was not effective in rescuing the increase in nuclear rows in the ONL, which indicates that it is essential to transplant mice when some photoreceptors are still present. A fully degenerated photoreceptor layer cannot be reverted, which indicates that the time of intervention is critical. However, in human pathology, where degeneration is much slower than in rd10 mice, it is possible that the time window to transplant the cells will be considerably larger. It is also worth considering that different signals can be activated by damaged neurons and they can induce alternative differentiation responses; for example, MG regenerates photoreceptors when these specific cells are destroyed in goldfish (52). These signals might in principle be targeted to enhancing the efficacy of the therapy.

In vitro-generated neurons after transdifferentiation of somatic cells are an exciting tool for regenerative medicine; however, engraft problems are a bottleneck for this approach. Strategies based on in vivo conversion of resident cells through the overexpression of key transcription factors might overcome this limitation; nevertheless, to reach clinical applications, it is imperative to obtain a safe in vivo delivery system to specifically target cells where the expression of reprogramming factors can be turned off. In this context, cell fusion with endogenous glial cells represents a promising therapeutic approach that might overcome all of these hurdles. Fusion of transplanted HSPCs might be used to deliver a specific combination of key factors to the targeted cells, once shown to be efficient for inducing conversion of MG into functional photoreceptors. Alternatively, HSPCs can deliver WT alleles after fusion with MG carrying recessive mutations, therefore providing a transversal strategy to correct genetic defects in a variety of patients. HSPCs have also been shown to stabilize retinal blood vessels in rd10 mice, thereby providing a neurotrophic beneficial effect for the preservation of the retinal nuclear layers (53). Importantly, clinical trials have already shown the safety of HSPCs (54).

Still, some questions remain open in our studies. We observed some single-color hybrids (only YFP+), which indicates silencing of the RFP transgene or ploidy reduction. We previously showed that in vivo-formed hybrids appear as synkaryons or heterokaryons, thereby containing 1 or 2 nuclei (28). Multipolar mitosis with ploidy reduction of hybrids was seen in the liver after fusion of hepatocytes with BMDCs (55). On the other hand, heterokaryons have been found in the CNS after BM transplantation (56), which indicates that neurons can cope with high ploidy content.

Here, we also provided evidence that Wnt activation increases the proliferation potential and survival of the hybrids, which in turn downregulate the 2 cell-cycle regulators p27^{Kip1} and p57^{Kip2}. These 2 factors control retinal progenitor cell-cycle exit and differentiation during retinal development (14). Interestingly, p27^{Kip1} is downregulated very early in MG in regenerating fish retinas (57).

We previously showed that upon retinal injury, MG can fuse with endogenous mobilized BMDCs (28), which suggests that this might also be a physiological response to the damage in mammals and lower vertebrates.

In conclusion, we have identified a route to reprogramming MG in vivo in order to functionally regenerate photoreceptors in a mouse model of retinitis pigmentosa. This approach can be potentially exploited in the future for the treatment of patients.

Methods

Animal care and treatment. In this study, we used the following male and female transgenic mice: Vav-Cre (58), CAG-RFP (59), Gfap-Cre (60), R26Y (61), RHO-Cre (32), Calr-Cre (33), and nestin-Cre (62). Mice were purchased from Jackson Laboratory. R26Y mice were a gift of M. Studer (Institute of Biology Valrose, Nice, France). For MNU-induced photoreceptor degeneration, adult recipient mice (8 to 12 weeks old) were treated with an i.p. dose of sterile MNU (60 mg/ kg body weight; Sigma-Aldrich) 24 hours before cell transplantation. For BrdU tracking of proliferating hybrids, mice were treated with a 2 i.p. dose of BrdU (15 mg/kg) every 12 hours, with the first pulse at the moment of the cell transplantation. Mice were maintained under a 12-hour light/12-hour dark cycle, with access to food and water ad libitum. For the whole study, animals were randomly assigned to treatment groups. We chose groups of a minimum of 6 mice because that was the minimal number of animals required for reaching statistical significance. General anesthesia was induced when needed with i.p. injections of ketamine (70 mg/kg, Imagene; Merial) and medetomidine (10 mg/kg, Domitor; Pfizer Animal Health).

HSPC isolation and treatment. Donor adult mice (8 to 12 weeks old) were sacrificed by cervical dislocation, and total BM was flushed out from femurs and tibia. HSPCs (Lin⁻) were purified from total BM cells by untouched isolation on a magnetic sorter using the Lineage Depletion Kit (Miltenyi Biotec) according to the manufacturer's protocol. Cells were cultured in StemSpan SFEM medium (STEMCELL Technologies) at a concentration of 1×10^6 cells/ml in the presence of 100 ng/ml recombinant mouse SCF (R&D Systems), 20 ng/ml mouse IL-3 (Peprotech), 20 ng/ml human IL-6 (Peprotech), and 10 ng/ml FLT3 ligand (Peprotech). Cells were treated with $1\,\mu\text{M}$ BIO (Calbiochem) for 24 hours or untreated as control before transplantation. Thirty minutes before transplantation, cells were incubated with Vybrant DiD Cell-Labeling Solution (Life Technologies) for 30 minutes when necessary, washed twice in PBS, and resuspended to a final concentration of 1×10^5 cell/ μ l for transplantation.

Subretinal transplantation of cells. Anesthetized recipient animals received a single 1 μ l administration of HSPC suspension (1 × 10 5 cells/ μ l) by subretinal route. Injections were performed under an operating microscope (Zeiss OPMI Pico; Carl Zeiss Meditec GmbH) with the aid of a 5- μ l Hamilton microsyringe (Hamilton Co.). Each mouse was kept on a 37 $^{\circ}$ C pad after it was anesthetized, during and after surgery. Briefly, a scalpel was used to make a small cut through the sclera in a temporal position. Then, a 33-gauge bevelled (45 $^{\circ}$) needle located on a Hamilton syringe system was inserted in the hole between the epithelium and the retina with a tangential angle with the flat part of the bevel toward the retinal side. Cell suspension was injected with a slow and constant pressure. The needle was left in position for another 5 to 10 seconds to allow pressure in the injection system to equilibrate with the eye pressure and then removed by slowly pulling backwards

to allow the hole in the retina to reseal and to avoid damaging the retina and corneal endothelium during removal.

OCT. To further confirm the correct delivery of the cells in the subretinal space, noninvasive in vivo assessment of retinal detachment formed upon subretinal injections was performed by OCT of 2 groups of mice (group 1, n = 12; group 2, n = 16; 4 different litters in total) subretinally injected by 2 different operators. Two independent experiments were sufficient to confirm the same percentage of success of the subretinal injection procedure (32.2%). For this purpose, groups with different variances were compared. The assessment was performed when mice were under general anesthesia as described in the Animal care and treatment section. Combined confocal scanning laser ophthalmoscopy (cSLO) and SD-OCT imaging were performed using a commercially available device (Spectralis HRA OCT system, Spectralis Eye Explorer 5.6b, Heidelberg Engineering). To adapt the system to the mouse eye, a commercially available +25D lens was fit to the system (Heidelberg lens, HE 50744) and the length of the reference pathway was manually adapted according to the manufacturer's instructions. The cSLO system used was an HRA2, using solid source for 488 and 514 nm and an infrared diode laser for 785 and 815 wavelengths. In order to improve the image acquisition quality, mice wore a PMMA afocal contact lens, with a radius and diameter of 2.7 mm, respectively. Contact lenses were purchased from Cantoor Nissel (Market Place Brackley). Tear substitutive drops were continuously added to improve transparence (Systane, Alcon Labs). Mice were analyzed for cSLO/ SD-OCT with a single scan looking for the injection site.

The correlation analysis between the positive delivery of cells in the subretinal space and the increase in the number of nuclear rows in the ONL, INL, and GC layer (GCL) was carried out blindly before inspection of the histological results.

Electroretinogram tests. For the ERG tests, treated 2-month-old mice were dark adapted for 3 hours, anesthetized, and placed in a stereotaxic apparatus under dim red light. ERGs were evoked by flashes of different light intensities ranging from 10⁻⁴ to 20 cd m⁻² s⁻¹ generated through a Ganzfeld stimulator (Lace Elettronica). Three different responses evoked by light were averaged for each luminance step to minimize the noise. The electrophysiological signals were recorded through gold plate electrodes inserted under the lower eyelids referred to a needle electrode inserted s.c. at the level of the corresponding frontal region. The different electrodes were connected to a 2-channel amplifier. The amplitudes of the A- and B-waves measured for each eye upon stimulus with the higher intensity flash under scotopic condition were plotted.

Hybrid isolation for gene expression and flow cytometry analysis. For FACS analysis, retinal samples were isolated, disaggregated in trypsin by mechanical trituration, and filtered. Samples were analyzed by flow cytometry in an LSR I instrument or LSR Fortessa (BD) with FlowJo (Tree Star Inc.) or FACSDiva (BD) software in order to assess the percentage of DID+YFP+ hybrids. In all samples, DAPI (Sigma-Aldrich) was added to exclude dead cells from the analysis.

For gene expression analysis, a BD FACSAria II sorting machine (BD) was used to isolate either the double DiD+YFP+ or RFP+YFP+ hybrids. DAPI (Sigma-Aldrich) was added to exclude dead cells. Total RNA was extracted using RNA Isolation Mini or Micro Kits (QIAGEN) according to the manufacturer's protocol. The eluted RNA was reverse-transcribed with SuperScript III (Invitrogen), and real-time quantitative PCR (qPCR) reactions using Platinum SYBR

Green qPCix-UDG (Invitrogen) were performed in an LightCycler 480 (Roche) real-time PCR machine according to the manufacturer's recommendations. The oligos used are listed in Supplemental Table 1. *Runx1* and *Gata1* oligos were purchased from QIAGEN. All of the experiments were performed in triplicate, and differences in cDNA input were compensated by normalizing to the expression of *Gapdh*. Relative mRNA levels were evaluated as the fold changes with respect to the untreated control or to retinal samples depleted by sorting of either the double DiD+YFP+ or RFP+YFP+ hybrid population.

Fixing, sectioning, immunohistochemistry, and Western blotting. Eyes were fixed by immersion in 4% paraformaldehyde overnight and then embedded in paraffin. Serial transversal sections of 10-µm thickness were processed for TUNEL staining (In Situ Cell Death Detection Kit, Roche) according to the manufacturer's protocol. For immunostaining, sections were treated with serial washing in Xilene for 2 minutes, EtOH 100% for 1 minute, EtOH 95% for 1 minute, EtOH 90 for 1 minute, EtOH 80% for 1 minute, EtOH 70% for 1 minute, EtOH 50% for 1 minute and H₂O. Slices were placed in a plastic rack with an antigen retrieval buffer (NaCitrate 0.1 M Triton X-100 0.1%) and boiled for 4 minutes in a domestic microwave oven. After a wash with cold water, sections were blocked in NGS 10% for 30 minutes and NGS 1% for 30 minutes. Primary antibodies were incubated in PBS overnight at 4°C. The primary antibodies used were as follows: rabbit anti-glutamine synthetase GS (1:300, Sigma-Aldrich, G-2781) mouse anti-PCNA (1:200, Sigma-Aldrich, P8825), chicken anti-GFP (1:500, Abcam, AB13970), rabbit anti-recoverin (1:500, Millipore, AB5585), mouse anti-BrdU (1:200, Sigma-Aldrich, B8434), rabbit anti-OTX2 (Abcam, AB21990), rabbit anti-calbindin D-28K (1:50, Sigma-Aldrich, C7354), mouse anti-calretinin (1:100, Millipore, MAB1568), mouse anti-SMI-32 (1:100, Millipore, NE1023), rabbit anti-PKC-α (1:200, Abcam, AB32376), and mouse anti-β-tubulin III (clone TU-20, Millipore, MAB1637). The secondary antibodies used were as follows: anti-chicken Alexa Fluor 488 (1:1000; Molecular Probes, Invitrogen), anti-chicken Alexa Fluor 568 (1:1000; Molecular Probes, Invitrogen), anti-mouse Alexa Fluor 568 (1:1000; Molecular Probes, A11031, Invitrogen), anti-rabbit Alexa Fluor 568 (1:1000; Molecular Probes, A11036, Invitrogen), anti-mouse Alexa Fluor 633 (1:1000; Molecular Probes, A21050, Invitrogen), and anti-rabbit Alexa Fluor 633 (1:1000; Molecular Probes, A21070, Invitrogen). Nuclei were counterstained with DAPI (Vectashield, Vector Laboratories). Photographs were taken using either an Axioplan microscope (Zeiss) or a Leica laser SP5 or SPE confocal microscopy systems.

For Western blotting, whole cell lysates were prepared by resuspending retinas in 20 mM Tris–HCl, pH 7.5, 150 mM NaCl, 1 mM MgCl₂, 0.1 mM CaCl₂, 10% glycerol, 0.1% Triton X-100, and protease inhibitors (phenylmethylsulfonyl fluoride and cOmplete protease inhibitor cocktail; Roche Molecular Biochemicals), followed by syringing through a 23-gauge needle. Cell lysates (30 µg/lane) were electrophoresed through a 10% polyacrylamide-SDS gel and proteins transferred to a polyvinylidene difluoride (PVDF) membrane. Membranes were immunostained overnight at 4°C with primary anti-PDE6 β (1:500, Abcam AB5663) and anti- β -actin (1:5,000, Abcam AB8226) antibodies and 1 hour at room temperature with secondary anti-mouse IgG (1:2000, GE HealthCare, NA931V) and anti-rabbit IgG (1:2000, GE HealthCare, NA934V) antibodies. The signal was visualized using Pierce ECL Western Blotting Substrate (Thermo Scientific) chemiluminescence detection systems.

Statistics. Percentages of PCNA/YFP or TUNEL/YFP double-positive cells relative to total YFP+ detected in the retinas as well as the percentage of YFP/BrdU/marker-positive hybrids relative to the total YFP/BrdU cells detected in the retinas were evaluated by staining the tissue sections with anti-GFP and different antibodies or TUNEL and analyzing photographs acquired with ×40 magnification. At least 3 different retinal sections for each eye in 3 different independent experiments were evaluated and results were reported as mean ± SD. Experiments performed in the whole work with a higher number of duplicates are specified in the text. For the counting of the number of nuclear rows in either the ONL or in the INL and of number of nuclei in the GCL, photographs of the temporal, central, and nasal areas of 9 different serial sections for each eye were acquired with a ×40 magnification and analyzed. Differences were examined using a 2-tailed unpaired Student's t tests or, in the case of multiple comparisons, using 2-way ANOVA test and Bonferroni's post-test. P < 0.05 was considered significant. The distribution and variance were normal and similar in all groups statistically compared.

Study approval. All of the procedures on mice were reviewed and approved by the Comité Ético de Experimentación Animal del Parc de Recerca Biomèdica de Barcelona (Spain) and were performed according to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

Author contributions

DS designed and performed the experiments, analyzed data, and wrote the paper. GS performed experiments and analyzed

data. UDV performed ERG. NR performed cell sorting. IP performed the OCT analysis on a group of mice. MN supervised the OCT analysis and performed the quality control for the experiments on the rescue of the *rd10* phenotype, which were carried out blind. MPC analyzed the data, wrote the paper, and supervised the project.

Acknowledgments

We would like to thank Andrés Fernández Garcia for support and critical discussions, Francesco Sottile and Ilda Theka for critical reading of the manuscript, the Histological and Microscopy Facilities of CRG and BLOSS Group Company technicians for assistance in the OCT analysis. This work was supported by an ERC grant (242630-RERE to M.P. Cosma), the Ministerio de Economia y Competitividad y FEDER (SAF2011-28580, and BFU2015-71984, BFU2015-71984-ERC to M.P. Cosma), an AGAUR grant from Secretaria d'Universitats i Investigació del Departament d'Economia i Coneixement de la Generalitat de Catalunya (2014SGR1137 to M.P. Cosma), 976_Velux Stiftung (to M.P. Cosma), TV3'12_120530_ FUNDACIO MARATO TV3, Ferrer International (to M.P. Cosma and D. Sanges), and Subprograma Juan de la Cierva - Ministerio Ciencia e Innovación (to D. Sanges).

Address correspondence to: Maria Pia Cosma, Center for Genomic Regulation (CRG), C/Dr.Aiguader, 88, 08003 Barcelona, Spain. Phone: 34.9.33160370; E-mail: pia.cosma@crg.es.

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