PEDF and GDNF are key regulators of photoreceptor development and retinal neurogenesis in reaggregates from chick embryonic retina

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Abstract Here, role(s) of pigment epithelial-derived factor (PEDF) and glial-derived neurotrophic factor (GDNF) on photoreceptor development in three-dimensional reaggregates from the retinae of the E6 chick embryo (rosetted spheroids) was investigated. Fully dispersed cells were reaggregated under serum-reduced conditions and supplemented with 50 ng/ml PEDF alone or in combination with 50 ng/ml GDNF. The spheroids were analyzed for cell growth, differentiation, and death using proliferating cell nuclear antigen, terminal deoxynucleotidyl transferase deoxyuridine triphosphate nick end labeling, and other immunocytochemical stainings and semi-quantitative reverse transcription polymerase chain reaction (RT-PCR) methods. PEDF strongly promoted synthesis of the messenger RNAs for blue and violet cone opsins and to a lesser extent on the red and green cone opsins. This correlated with an increase in the number of cone photoreceptors, as determined by the cone cell marker CERN906. Likewise, PEDF nearly completely inhibited rod differentiation, as detected by immunostaining with anti-rho4D2 and RT-PCR. Furthermore, PEDF accelerated proliferation of cells in the spheroids and inhibited apoptosis. As negative effects, PEDF inhibited the normal histotypic tissue formation of retinal aggregates and reduced the frequency of photoreceptor rosettes and IPL-like areas. Noticeably, supplementation of PEDF-treated cultures with GDNF reversed the effects of PEDF on spheroid morphology and on rod differentiation. This study establishes that PEDF strongly affects three-dimensional retinogenesis in vitro, most notably by inhibiting rod development and supporting proliferation and differentiation of cones, effects which are partially counteracted by GDNF.

Keywords Cytokines · Muller glial cells · Photoreceptor degeneration · Retinal spheroids · Tissue formation

Introduction

During retinogenesis, the neural retina develops from the inner layer of the optic vesicle and the retinal pigment epithelium (RPE) from the outer layer. The close proximity of both tissues and their intricate interactions are essential for function, structure, and differentiation of photoreceptors [1–4]. In particular, the development of the photoreceptor outer segments is highly dependent on the RPE, e.g., after RPE removal, the structural organization of the outer segment is disrupted [5]. The interdependence between the RPE and photoreceptors is mediated by the exchange of nutrients, metabolites, and various signal molecules secreted from both cell populations. Numerous growth factors have been identified as RPE-secreted products [6–10],
including pigment epithelium-derived factor (PEDF) [11]. Analysis of the specific roles of these molecules in vivo is complicated because of their multiple and combined effects (“cocktail effects”) on the development, survival, and differentiation of the retina. A previously developed three-dimensional in vitro culture of reaggregated retinal organotypic spheres has been found to represent a most suitable model to dissect the effects of individual growth factors in early retinal development. The reaggregates are derived from dissociated cells isolated from 6-day-old chick retina. Under rotation conditions, the cultures will form three-dimensional rosetted spheroids that attempt to imitate normal retinal development (see Fig. 3). The innermost part of the spheroids contains rosettes that are composed of photoreceptors and thus correspond to the outer nuclear layer of a normal retina. Each rosette is surrounded by a circular outer plexiform layer, followed by sections of an inner nuclear layer (INL). Next to the INL are inner plexiform layer (IPL)-like areas which are cell-free island-shaped areas, consisting of neurites of bipolar, ganglion, and amacrine cells (for review [12]). Within these IPLs, only a few displaced ganglion and amacrine cells are detectable. In addition to the above structures, there are non-organized areas of the spheroids consisting of different retinal cell types and their corresponding fibers. The rosetted spheroids arise first through aggregation of the dissociated retinal cells, followed by cell proliferation, cell migration, differentiation, and programmed cell death. The spheroids are easy to manipulate and their growth can be altered by supplementation with individual growth factor(s) and a cocktail of molecules or by blocking the expression of key proteins using gene knockdown strategies.

Using this culture model, the function of glial-derived neurotrophic factor (GDNF) on retinal development was investigated [13, 14]. GDNF, a distant member of the transforming growth factor beta superfamily, is a potent neurotrophic factor for a variety of central and peripheral neurons [15, 16]. Using retinal spheroids, GDNF was shown to regulate proliferation, differentiation, and survival of chicken rod photoreceptors in vitro [13], while it had minor effects on cone photoreceptors. Interestingly, knockdown of a particular receptor for GDNF predominantly affected the development of violet photoreceptors [17]. In particular, GDNF increased proliferation of rod precursors, promoted the onset of their differentiation, increased their number, and prevented programmed cell death.

Here, we focused on another potent factor, PEDF, which is synthesized and secreted by the RPE. Originally, PEDF was isolated from conditioned medium of human RPE cells, inducing neuronal differentiation of human retinoblastoma cells [11, 18]. The 50-kDa PEDF protein belongs to the family of serine protease inhibitors, however, not exerting proteolytic activity [19]. Being expressed in several parts of the central nervous system and eye tissues of humans [20, 21], PEDF has been assigned several physiological roles in various tissues, e.g., an anti-angiogenic effect during neovascularization [22], including mammalian eyes [23, 24]. In central nervous system and peripheral nervous system neurons from rodent and human origins, PEDF acted as survival factor in vitro [25–27]. The significance of PEDF in the developing neural retina is not well understood. PEDF is expressed by more than one retinal cell type in several mammalian retinas, e.g., Muller and ganglion cells in rodents can express and secrete PEDF [24, 28] and possibly protect themselves by this factor [29]. An expression of PEDF in photoreceptors has been detected only in human retinae [24]. Because of its abundant secretion by the RPE, several studies have focused on roles of PEDF for photoreceptor survival. For example, PEDF was capable of protecting rat photoreceptors from death in vitro induced by light, oxidative stress, or glutamate-induced toxicity [5, 30].

In the first part of this study, we applied PEDF to rosetted spheroids of retinal reaggregates to investigate its role in proliferation, differentiation, and survival of specific retinal cell types, in particular the photoreceptors. We showed that, while the overall morphologic tissue development remained inferior to the control after treatment, PEDF increased and sustained proliferation and strongly decreased the number of apoptotic cells. Furthermore, PEDF showed opposing effects to GDNF on the differentiation of rod and cone photoreceptors. A combination treatment with both PEDF and GDNF resulted in synergistic effects on the survival and proliferation of cone photoreceptors and the morphology of the spheres. From these studies, we propose that PEDF and GDNF are key players in the development of rod and cone photoreceptors during retinogenesis.

Materials and methods

Tissue culture

To produce rosetted retinal spheroids, the retinas from 6-day-old chicken embryos (E6, white leghorn) were used. The central parts of the retina were isolated and collected in F12 medium on ice. The retinal tissue was dissociated by tryptic digestion in F12 medium containing 0.05 mg/ml trypsin (Worthington Biochemicals/Cell Systems, Remagen, Germany) for 8 min at 37°C. The remaining cell clusters were mechanically dissociated in Hanks’ balanced salt solution containing 0.5 mg/ml DNase I (Worthington Biochemicals/Cell systems, Remagen, Germany) by 30 to 35 gentle strokes with a round-bored Pasteur pipette. For generation of retinal spheroids, 2 × 10⁶ cells per milliliter were cultured in 35-mm dishes containing 2-ml aggregation
medium (Dulbecco’s modified Eagle’s medium, 2% fetal calf serum, 1% l-glutamine, and 0.15% penicillin/streptomycin, all from Gibco, Berlin, Germany) on a gyratory shaker in an incubator at 37°C and 5% CO₂ (Heraeus Holding GmbH, Hanau, Germany), either in absence of growth factors or presence of 50 ng/ml PEDF, presence of 50 ng/ml GDNF (Sigma, Deisenhofen, Germany), or in a combination of both.

Cryosections and immunostaining procedures

For obtaining cryosections, rosetted spheroids were harvested at the indicated time points and fixed in 4% formaldehyde (Merck, Darmstadt, Germany) for 30 min at room temperature. After the fixative was removed by two washes in phosphate-buffered saline (PBS), rosetted spheroids were soaked in 25% sucrose (Merck, Darmstadt, Germany) and stored at 4°C. Cryosections of 10-μm thicknesses were cut on a cryostat (Microm, Waldorf, Germany), mounted on gelatine-coated slides, and stored at −20°C.

For immunostaining, sections were dried at 37°C and pre-incubated in blocking solution, containing 3% bovine serum albumin, 0.1% Triton-X-100 (all from Sigma, Deisenhofen, Germany) in PBS for 30 min at room temperature. Then, tissues were incubated with the primary antibody for 75 min, followed by three washes in PBS. The rod-specific monoclonal antibody rho4D2 (a generous gift from D. Hicks, Strasbourg, France) was used at a dilution of 1:1,000 in blocking solution, and the red- and green-specific antibody CERN906 (a generous gift of W. DeGrip, University of Nijmegen, The Netherlands) was used at a dilution of 1:500 in blocking solution. For detection of the primary antibody, sections were incubated with donkey anti-mouse-conjugated Cy3 and goat anti-rabbit-conjugated-Cy2 (both 10 μg/ml; Dianova, Hamburg, Germany) for 1 h. For detection of proliferation, the staining reagent proliferating cell nuclear antigen (PCNA) was used at a dilution of 1:250 in blocking solution; cryosections were incubated for 3 h with PCNA. Before pre-incubating in blocking solution, sections were incubated with 4 N HCl for 10 min. For detection of apoptosis, sections were terminal deoxynucleotidyl transferase deoxyuridine triphosphate nick end labeling (TUNEL)-stained according to the manufacturer’s instructions (Roche Molecular Biochemicals, Mannheim, Germany). Between the last washes, cell nuclei were stained with 4′,6-diamidino-2-phenylindole (DAPI; 0.1 mg/ml 4′,6-diamidine-2-phenylindol-dihydrochloride in PBS; Merck, Darmstadt, Germany). Finally, sections were dried and embedded in Kaiser’s glycerine gelatine (Merck, Darmstadt, Germany). It should be noted that, due to limited availability of spheroids, cryosections have been taken not only from their equatorial level. Therefore, different aerial sizes of cryosections as shown in figures do not necessarily indicate a smaller spheroid volume.

RT-PCR

For semi-quantitative reverse transcription polymerase chain reaction (RT-PCR), total RNA was isolated from retinal cultures at appropriate stages and transcribed into complementary DNA (cDNA) by using the reverse transcription system from Promega (Mannheim, Germany). Target cDNA was amplified by PCR with specific primers. PCR was performed using parameters of 94°C for 1 min, 57°C for 1 min, and 72°C for 1 min for 29 cycles. Primers for PCR were 5′-GGACAGTTCAAGGCACTGT-3′ and 5′-CTTCTGTGTGGCTGTGATG-3′ for GAPDH, 5′-TCACC ATCCAGCAACAGAAA-3′ and 5′-GACCAAGTTCCCAT AGCAGA-3′ for rhodopsin, 5′-ATCAACCAGATCTCG GGGTAG-3′ and 5′-CAGAAGCAGTAGGCCACG AT-3′ for red opsin, 5′-CATGATGGGGATAGCTTTCA-3′ and 5′-TGG TGATCATGGCAATTACGG-3′ for green opsin, 5′-CGCTCT GAGTGATGAGGAGA-3′ and 5′-AGCTCCGTTCCTCC ACCTTAAT-3′ for blue opsin, and 5′-CGTGCGCTCTCA CCTACAG-3′ and 5′-TACTCGCTGCCGTATTTCTGT-3′ for violet opsin. The relative expression of messenger RNA (mRNA) was quantified as a percentage of GAPDH band intensity at the indicated stages.

Cell counting and statistical analysis

To determine the number of immunostained cells, frozen sections were stained with DAPI and rho4D2 antibody. Then, the percentage of immuno-labeled cells per section of a single spheroid was calculated in relation to DAPI-positive cells of the same spheroid. At least six cryosections of different spheroids derived from two individual experiments were analyzed. Data are presented as the mean±SD and compared by a two-tailed Student’s t test. Note that the standard deviation represents the difference between calculations of individual spheroids.

To determine the relative expression of mRNA, the percentage was calculated in relation to GAPDH band intensity for each culture stage. Each experiment was carried out at least three times.

Microscopy and photography

Photomicrographs of sections were taken with an Axioskop microscope (Carl Zeiss, Jena, Germany) combined with a charge-coupled device three-color (CCD-3) digital camera (Intas, Göttingen, Germany). Photomicrographs were processed on a computer with documentation and analysis software (Diskus Histologie MAN F70B, Hilgers, Königs- winter, Germany; Multianalysts 2.0, Biorad, München,
Germany; Adobe Photoshop CS, Adobe, San Jose, CA, USA; and Excel, Microsoft, Redmond, WA, USA).

Results

PEDF enhances proliferation

When PEDF, at a concentration of 50 ng/ml, was added to rosetted spheroids developing from E6 chick retinae under serum-reduced culture conditions, the number of proliferating cells (PCNA+ cells) was significantly increased (Fig. 1). This proliferative stimulation was most pronounced at early culture stages. At day in vitro (div) 2, 28% of cells in controls were proliferating, while in PEDF-treated cultures 38% proliferating cells could be identified (Fig. 1e). The stimulatory effect on proliferation became much more pronounced at days 4 and 6. Compared to controls, the number of proliferating cells was now more than doubled after PEDF treatment. Thus, in the presence of PEDF, 17% of all cells were proliferating at day 4 and 8% at day 6, compared with 6% and 3% in the respective controls. At day 8, proliferation had become very low in both cultures and did not differ significantly between treated and non-treated cultures (Fig. 1e).

Apoptosis is strongly decreased by PEDF

Using the TUNEL assay, a very dramatic inhibitory effect of 50 ng/ml PEDF on programmed cell death was revealed in our retinal cell cultures (Fig. 2). Histologically, cells of the entire spheroid were evenly protected from apoptosis (cf. Fig. 2b, d). The percentage of cells undergoing apoptosis in PEDF-treated and non-treated cultures was quantified in relation to the total number of DAPI-positive cells at day 10 (Fig. 2e). The percentage of apoptotic cells was only 8% in presence of PEDF, while in controls 32% of cells were identified as apoptotic.

PEDF inhibits spatial organization of retinal spheroids

The morphology of spheroids as characterized by typical structures like rosettes and IPL-like areas was also strongly affected (Fig. 3; see also Figs. 1 and 2). In Fig. 3a–c, we show a triple-stained retinal spheroid, stained with DAPI (Fig. 3a), anti-rho4D2 (specific for rod photoreceptors, in Fig. 3b), and anti-CERN906 (specific for red and green cones, in Fig. 3c), with one small photoreceptor rosette (broken circle), composed of rods (Fig. 3b) and cones (Fig. 3c). Near the surface of the sphere, there are four large circular IPL-like areas, each containing a few isolated cells (Fig. 3a, IPL, arrow). In the presence of PEDF, these histotypical structures found in the untreated controls were very rare (Fig. 3d–f; Figs. 1 and 2). Instead, the sphere mostly appeared as a non-organized cell aggregate, suggesting that PEDF strongly interfered with development of the normal tissue organization of the spheroids.

PEDF decreases rho4D2+ cells and rhodopsin mRNA expression

Since several neurotrophic factors affect differentiation of various retinal cell types, including photoreceptors, we were interested in the effect of PEDF on rod and cone photoreceptor development in the spheroids (Fig. 3). PEDF strongly decreased the staining for rod photoreceptors as compared to the controls (Fig. 3b, e). In the non-treated controls, groups of rho4D2-positive cells are shown in the center of the rosette as well as at the periphery and in intermediate locations (Fig. 3b). In contrast, spheroids treated with PEDF...
are very weakly stained for rho4D2, with only a few immuno-positive cells present (Fig. 3e). Based on DAPI staining of the identical control and experimental spheroids (Fig. 3a, d), the percentages of rho4D2-positive cells were quantified in controls and in PEDF-treated cultures (e). Each data point represents the mean±SD of multiple sections (n=6). **P<0.001. Scale bar, 100 μm.

These findings were independently confirmed by semi-quantitative RT-PCR (Fig. 4). The temporal expression of endogenous rhodopsin mRNA was determined between days 2 and 10 in controls (Fig. 4a) and in PEDF-treated cultures (Fig. 4b). The percentage of rhodopsin mRNA was calculated relative to the expression of GAPDH mRNA and the quantitative analysis is presented in Fig. 4c. In the absence of PEDF, there was a gradual increase in rhodopsin mRNA from 7% to 22% in the cultures at days 2 to 6 with peak levels at day 6. This was followed by a decrease to about 3% by day 10. In the presence of PEDF, a quantitatively and temporally different expression pattern was revealed over this time in culture. Rhodopsin expression showed a very high level of 24% already at day 4. But then, rhodopsin expression was quickly and efficiently downregulated, leading to a sharp decline to 0.2% by day 10. These studies indicate that PEDF accelerated
rhodopsin expression for a short period of time, then almost completely inhibited it.

PEDF increases CERN906⁺ cones

While PEDF showed dramatic inhibitory effects on rho4D2-positive cells and on rhodopsin mRNA expression in the spheroids, the number of CERN906-positive cone photoreceptors was significantly increased, as revealed by the red/green cone-specific antibody CERN906 (Fig. 3c, f). Similar to the arrangement of rods, CERN906-positive cells were found inside the rosette, and a relatively high number was also seen at the periphery in control cultures (Fig. 3c). After treatment with PEDF, many CERN906-positive cells were found distributed all over the aggregate (Fig. 3f). At all time points examined, the number of red and green CERN906-positive cone cells was higher in PEDF-treated cultures (Fig. 3h). At day 8, there were 15% CERN906⁺-positive cells in the presence of PEDF and 9% in the absence of this neurotrophic factor. At day 10, the number of cones represented 20% and 8% of the total cell population with and without PEDF treatment, respectively. Note that while, in the control cultures, red and green cone cell number decreased from day 8 to 10, in the presence of PEDF, the highest number of CERN906-positive cells was observed at day 10, suggesting a long-lasting effect of PEDF on cone cell differentiation and survival.

Subpopulations of cones are selectively affected by PEDF

The effects of PEDF on specific types of opsins were analyzed by semi-quantitative RT-PCR (Fig. 5). We analyzed the temporal expression of RNAs for red, green, blue, and violet cone opsins in the controls (Fig. 5a) and in PEDF-treated cultures between days 2 and 10 in culture (Fig. 5b). PEDF increased the expression of all four cone opsin RNAs, although each showed a distinctive pattern. The data are quantified and compared in Fig. 6a–d.

The expression of red cone opsin RNA following PEDF treatment showed a rapid increase and then stayed at an almost steady level of approximately 20%, while in controls it decreased after div 6 (Fig. 6a). The expression of green cone opsin was very similar to that of red cone opsin (Fig. 6b vs. a). PEDF induced a rapid increase in expression of blue cone opsin RNA between div 2 and 6 (approx...
imately sevenfold at div 2), after which time the levels decreased (Fig. 6c). Finally, violet cone opsin expression was influenced by PEDF in a pattern similar to that for blue cone opsin, e.g., at div 6 PEDF led to an approximately sevenfold increase of violet opsin expression (Fig. 6d; for its high expression in the control at div 8, see “Discussion”). In summary, while all four types of cone opsin expression were increased by PEDF, most pronounced effects were documented for blue and violet opsins.

GDNF in combination with PEDF restores the spatial organization of retinal spheroids

Since application of PEDF disrupted the spatial organization of rosetted spheroids (Figs. 1, 2, and 3) and had very different effects to those we have previously reported for GDNF, we were interested in the effects of a combined application of PEDF and GDNF. Application of 50 ng/ml GDNF plus 50 ng/ml PEDF completely restored the morphology of rosetted spheroids (Fig. 7d–f). The number and size of IPL areas appeared normal. Thus, GDNF counteracted the disrupting effects of PEDF on tissue formation of the spheroids.

GDNF counterbalances the effects of PEDF on rod photoreceptors

Since GDNF reversed the effects of PEDF on the morphology of spheres, we investigated the effect of combined treatment on photoreceptor regulation (Fig. 7). For both rods and cones, the combined treatment with GDNF and PEDF promoted a significant increase as compared with PEDF treatment alone (Fig. 3). While, by day 8, the percentage of rho4D2-positive cells was still about one third lower than in the control, by day 10 it was nearly doubled (Fig. 7g). The effects of a combined GDNF/PEDF treatment were at least as significant for CERN906-positive cone cells (Fig. 7c, f): at both days 8 and 10, the percentage of red/green cones was approximately doubled (Fig. 7h). Moreover, whereas in controls the percentage of CERN906-positive cells decreased from day 8 to day 10, in GDNF/PEDF-treated cultures it stayed constant over the remaining culture period at 13%.

Discussion

We have used a three-dimensional reaggregate model of the embryonic chick retina to investigate effects of PEDF on retinal tissue formation in vitro. Our findings have documented distinct effects of PEDF on the regulation of cell proliferation and apoptosis, as well as on the differentiation of rod and cone photoreceptors. Together, these effects resulted in an altered three-dimensional tissue structure.

PEDF increases spheroid size but inhibits tissue structure

Rosetted spheroids of retinal cells develop a very characteristic histotypic structure (see “Introduction”). Since we raised our cultures here under serum-reduced conditions, this typical structure was not always completely represented.
in each and every spheroid. Nevertheless, it was most obvious that the application of exogenous PEDF to cultures led to dramatic morphologic changes in rosetted spheroids. The most obvious difference was a lack of detectable rosettes. In addition, only a few IPL-like areas were formed, and these were small in size and disrupted in organization. At present, we investigate whether PEDF disrupts spheroid morphology by acting on Muller glial cells [12, 31] since this histologically essential cell type [32] is affected by several factors [33–36].

PEDF stimulates proliferation and inhibits apoptosis in rosetted spheroids

Trophic factors can influence diverse developmental processes and regulate cell proliferation and/or cell death of various cell populations in rodents [30, 37, 38]. Mainly studied in mammals, PEDF has been found to have potent anti-proliferative and pro-differentiation effects [39, 40]. However, in our chicken spheroid culture, the application of PEDF induced an increase in cell proliferation, probably due to species differences. Moreover, since proliferation in this model lasts only for a few days, this apparent difference in effect could also be due to an acceleration rather than an increase of proliferation, so that in effect the system would be pushed towards earlier differentiation. The application of PEDF led to a significant decrease in the number of apoptotic cells; an effect that lasted over extended culture periods. This finding is remarkable since growth factors that prevent programmed cell death foster hopes for the development of therapies for degenerative diseases. Not only could cell death be downregulated in cell culture systems supplemented with neurotrophic factors but also in animal models, in which growth factors were injected into degenerated tissues [41]. Our findings strongly support earlier studies (see below) that PEDF could become a candidate to prevent cells from dying in a blinding disease.

PEDF acts as a pro-differentiating factor for cone photoreceptors

The interaction between the retina and the RPE is essential for the structural and functional integrity of photoreceptors (see “Introduction”), and proteins secreted from the RPE support the survival of photoreceptors [42–44]. One of the most important of these is PEDF. It was shown earlier [5] that, after the application of PEDF, deleterious effects of RPE removal on rod photoreceptor ultrastructure, the composition of outer segment membranes, and the expression of opsin were reversed. Under in vitro conditions, PEDF protected photoreceptors from cell death induced by oxidative stress or light [30, 45, 46]. Also, injection of PEDF in eyes with an inherited retinal degeneration caused a significant protection and delay in the cell death of photoreceptors [47].

The role of PEDF in the developing retina is much less well characterized. PEDF is expressed early in retinal development [24], but its ability to influence retinal cell development has not previously been studied. Using both immunocytochemistry and RT-PCR, we found that PEDF decreased the number of rhodopsin-positive cells in the
spheroids by as much as two thirds. An early spurt of rhodopsin expression at day 4 i.e., which quickly was downregulated again (Fig. 4c), could indicate that PEDF pushes cells towards differentiation of rods but then reverts them to go into a cone lineage. Since rhodopsin is one of the first cell-type-specific genes expressed by developing rods, the results imply that PEDF is preventing retinal progenitors from adopting a rod phenotype.

On the other hand, PEDF increased the number of differentiated cone photoreceptors. By RT-PCR analyses, we identified stimulatory effects of PEDF on red, green, and much more strongly on blue and violet opsins. Using an antibody that recognized red and green opsin cones to label sections of spheroids, we found that the number of these cells was substantially increased by PEDF. This suggests that the major effect of PEDF is on the formation of differentiated cones and not just on the level of opsin expression by differentiated cells. How can the differential effect of PEDF on rods and cones be explained? Cones are born before rods, both in vivo and in spheroids [48–50]. We suggested earlier that PEDF may be accelerating differentiation of retinal cells. If this is the case, PEDF may be driving cells into earlier formed cell types (cones) leaving a decreased number of progenitors to form later cell types (rods). Applying earlier markers for photoreceptor precursors should further clarify this point.

**GDNF reverses the effects of PEDF**

In most in vivo situations, many neurotrophic factors are present and may act simultaneously and in concert (cocktail effects). Different factors can act sequentially, or simultaneously and additively (synergistically), or in a mutually inhibitory fashion [51–53]. It is also possible that several populations of cells need to cooperate in order to achieve combined effects of neurotrophic factors. For instance, subpopulations of developing sensory and motoneurons are dependent on the simultaneous action of GDNF and BDNF [54]. A combination of GDNF and CNTF achieved a higher protection of photoreceptors in several model systems [55]. GDNF and BDNF supported the survival of ganglion cells much more efficiently, as compared with either factor applied separately [56]. Analyzing cocktail effects can be extremely difficult, in particular under in vivo conditions. Retinal spheroids provide a useful three-dimensional tissue environment easily amenable for such studies.

We have previously shown that GDNF strongly supports rod development in spheroids but has minimal effects on cones, an effect opposite to that found for PEDF. Therefore, we applied GDNF together with PEDF to investigate possible interactions of GDNF and PEDF and to test whether one action was dominant. For rods, GDNF effectively reversed the inhibition by PEDF. Noticeably, the number of rods was still lower than in controls at the earlier time tested (div 8) but then was much increased by div 10, supporting again the above assumption that PEDF at earlier time points shifts the rod to cone ratio towards cones (see above). Addition of GDNF did not inhibit the action of PEDF in promoting red- and green cone opsin expression. Thus, a cocktail of the two factors eventually resulted in increased differentiation of both photoreceptor cell types. Overall, the addition of GDNF to PEDF in the spheroid cultures resulted in a restoration of the tissue formation and morphology to that seen in the control.

One possible explanation of the results is that while PEDF promotes differentiation of early progenitors, it could inhibit differentiation of late progenitors. GDNF promotes differentiation in a way that overcomes the inhibitory effect of PEDF. This could be by direct action on the progenitors or by an indirect action through other retinal cells, e.g., Muller glial cells. Muller cells are important for retinal tissue formation and their destruction leads to retinal disorganization in vivo and in vitro [32]. With their morphology reaching through the entire retinal width, Muller cells could well be cellular mediators for the effects of GDNF and/or PEDF on both photoreceptors and cells of the inner retina. For instance, a rescue of photoreceptors by GDNF was mediated via retinal Muller cells, possibly due to upregulating FGF-2, which in turn could act on photoreceptors [57]. This mechanistic scheme puts Muller cells into the center of GDNF actions in the retina and could well represent the cellular locus where the two factors compete with each other.

In summary, we have found four major effects of PEDF on the formation of retinal spheroids. First, it promoted cell proliferation and inhibited apoptosis. Second, it inhibited the overall spatial organization of rosetted spheroids. Third, PEDF inhibited the differentiation of rhodopsin-positive cells. Fourth, PEDF promoted the differentiation of all classes of cones (or their precursors), albeit with different kinetics and efficiencies. While some of the inhibitory effects of PEDF could be (in part) reversed by GDNF, the combination of the two factors resulted in the best differentiation of both classes of photoreceptors (note that differentiation of photoreceptors in spheroids never is complete). The use of neurotrophic factors or combinations of factors suggests fascinating possibilities to develop therapies for degenerating eye diseases, like retinitis pigmentosa [58–60]. A combination of PEDF and GDNF might be one such therapy and should be investigated further.

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