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Interphotoreceptor matrix-poly(ε-caprolactone) composite scaffolds for human photoreceptor differentiation

Petr Baranov¹, Andrew Michaelson², Joydip Kundu², Rebecca L Carrier² and Michael Young¹

Abstract
Tissue engineering has been widely applied in different areas of regenerative medicine, including retinal regeneration. Typically, artificial biopolymers require additional surface modification (e.g. with arginine–glycine–aspartate-containing peptides or adsorption of protein, such as fibronectin), before cell seeding. Here, we describe an alternative approach for scaffold design: the manufacture of hybrid interphotoreceptor matrix-poly (ε-caprolactone) scaffolds, in which the insoluble extracellular matrix of the retina is incorporated into a biodegradable polymer well suited for transplantation. The incorporation of interphotoreceptor matrix did not change the topography of polycaprolactone film, although it led to a slight increase in hydrophilic properties (water contact angle measurements). This hybrid scaffold provided sufficient stimuli for human retinal progenitor cell adhesion and inhibited proliferation, leading to differentiation toward photoreceptor cells (expression of Crx, Nrl, rhodopsin, ROM1). This scaffold may be used for transplantation of retinal progenitor cells and their progeny to treat retinal degenerative disorders.

Keywords
Retina, photoreceptors, interphotoreceptor matrix, polycaprolactone

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Introduction
Transplantation of stem cells or their progeny can be used to treat a large variety of degenerative disorders affecting organs with limited regenerative capacity, such as the nervous system and sensory organs including the eye. Although experimental studies have shown the potential of progenitor cells delivered as single-cell suspensions to integrate into degenerating retina, a tissue engineering approach incorporating biomaterials together with cells is particularly promising due to increased cell survival and directed cell differentiation. This is based on a variety of studies involving both natural biopolymers, such as gelatin, chitosan, alginate, and artificial polymers, including poly(lactic-co-glycolic acid) (PLGA), poly(methyl methacrylate) (PMMA), poly(glycerolsebacate) (PGS), poly(hydroxybutyrate) (PHB), and poly(ε-caprolactone) (PCL). These polymers provide a structural scaffold for cell growth and are intended to mimic the natural tissue niche.

The extracellular matrix structure and topography (e.g. basement membranes, fibrillar matrix) are often considered during the design of tissue engineering scaffolds. Porous scaffolds, which allow cell ingrowth, are representative of matrices encountered in connective/muscular/nerve tissue regeneration, while epithelial cells (including neuroepithelial) grow on a relatively flat basement membrane. PLGA and PCL films are suitable scaffolds for mouse and human...
retinal progenitor cells (hRPCs), resulting in robust cell integration into degenerating retina upon implantation and controlled differentiation. The precise mechanisms of material-driven fate specification are yet to be elucidated, although it is suggested that this phenomenon is mediated by surface physical (such as stiffness) or chemical properties, since PCL induces photoreceptor differentiation from hRPCs independently of microtopography. This is confirmed by work with neuron differentiation on electrospun PCL.15,16 Despite impressive ability of PCL to induce differentiation in these studies, its hydrophobic nature requires additional modification or coating of the surface with charged molecules, such as poly-d-lysine and/or extracellular matrix proteins, including fibronectin, laminin, or collagen to enable efficient cell attachment.17 An alternative is the incorporation of the extracellular matrix proteins18 or total extracellular matrix (ECM) fraction into the polymer. Optimally, the incorporated molecules should provide both adhesive properties and differentiation stimuli to the attached cells. This can be achieved with single proteins (fibronectin, laminin) or with tissue-specific decellularized ECM preparations. In the retina, the interphotoreceptor matrix (IPM) is the specialized matrix surrounding the outer segments of photoreceptors.20 IPM mediates key interactions between the photoreceptors and retinal pigment epithelium (RPE) including adhesion, outer segment stability, nutrient exchange, and trafficking of retinol isoforms in the visual cycle.21,22 Thus, IPM is a logical matrix to explore for incorporation into hRPC culture or cell delivery biomaterials.

The aim of this study is to elucidate the impact of incorporation of IPM on the differentiation process of hRPCs on PCL films. We have incorporated IPM into PCL and have shown that this composite scaffold, without additional modification, has improved adhesive properties and can drive photoreceptor differentiation of hRPCs.

Materials and methods

IPM isolation and characterization

The IPM was isolated from adult bovine eyes collected fresh from the abattoir (Research 87 Inc., Boston, MA, USA) within 2 h after slaughter. Briefly, the muscle was removed from the outside of the eye. An incision was made 0.5 cm behind limbus, and a complete circumferential cut was made, being careful not to cut through the vitreous humor. Next, the cornea and vitreous humor were removed from the eye. Phosphate-buffered saline (PBS) was then immediately poured into the eyecup until full. Using a microspatula (Hayman Style), the retina was gently pulled away from the pigmented epithelium. A cut was made through the optic nerve and the retina. The retina was collected using a cut transfer pipette and placed in PBS (Sigma–Aldrich, St. Louis, MO, USA) and then transferred to a Petri dish containing deionized water. After orbital shaking (3 min, 75 r/min, TECHNE Mini Orbital Shaker, TSSM1, Technne Inc., Burlington, NJ, USA), the retinal debris was removed and the native IPM was collected. The collected IPM was centrifuged (5000 r/min; Sorvall Legen X1R Centrifuge, Thermo Scientific, Waltham, MA, USA) at 25°C for 20 min to pellet the IPM. The PBS was then decanted and the pellet was re-suspended within PBS.

PCL and PCL–IPM production

A total of 10% (w/v) solution of PCL was prepared by dissolving 10 g of PCL (molecular weight (mw) 70,000–90,000; Sigma–Aldrich) in dichloromethane (Sigma–Aldrich) with mixing at room temperature for 24 h. IPM, isolated from 12 bovine eyes, was pelleted (pellet volume approximately 500 µL) by centrifuging at 3000g, dehydrated with isopropanol (Sigma–Aldrich), centrifuged, re-suspended in 5 mL of dichloromethane, and then mixed in 100 mL of 10% PCL solution. This preparation was used to prepare two types of scaffolds: films and PCL–IPM-coated plates. The films were prepared as previously described: sufficient volume (5 mL) was poured in the center of a silicon wafer with spinning (Brewer Bioscience, Brewer Science Inc, Rolla, MO, USA) at 2000 r/min for 2 min. The films were dried at 45°C for 30 min, then sterilized by soaking in 95% ethanol for 30 min, and washed in distilled water three times. PCL-coated Petri dishes or coverslips were prepared in the same manner: 60-mm-diameter polystyrene dishes (BD Biosciences, Franklin Lakes, NJ, USA) were fixed on the spinner rotor; 5 mL of PCL (or PCL–IPM) solution was poured in the center, and dishes were spun at 2000 r/min for 5 min. Drying and washing procedures were the same.

Characterization of scaffolds: scanning electron microscopy, contact angle, lectin staining

Scanning electron microscopy (SEM). The surface morphology of PCL and PCL–IPM surfaces was observed using an S-4800 Hitachi scanning electron microscope (Los Angeles, CA, USA) at an operating voltage of 3.0 kV. The samples were then attached to a specimen stub with a carbon adhesive tab, sputter coated with 15-nm gold–palladium, and examined at room temperature.

Contact angle. Contact angles were measured with static drops of water on PCL and PCL–IPM-coated glass coverslips using a contact angle measurement system (Phoenix 300 plus, SEO, Suwon City, Korea) to provide information about hydrophobicity and hydrophilicity of these surfaces. Each data point represents 10 independent measurements. The experiment was repeated twice.

Lectin staining. The lectin staining was performed on PCL and PCL–IPM-coated glass coverslips. The coverslips were...
treated with 0.1% bovine serum albumin (Sigma, St. Louis, MO, USA) to prevent non-specific staining. A total of 50µL of FPNA Fluorescein-Labeled Peanut Agglutinin (FPNA; Vector Laboratories) and 50 µL (Rhodamine-labeled Wheat Germ Agglutinin (RWGA; Vector Laboratories, Burlingame, CA, USA) were then added to each surface for 30 min in the dark. The staining solutions were then removed, and the surfaces were each flooded with 1 mL of PBS. The surfaces were then examined under a fluorescence microscope.

**hRPC isolation, expansion, and characterization**

All work with human material was performed with institutional review board (IRB) approval. hRPCs were isolated from human fetal neural retina at 16 weeks of gestational age, as previously described.23 Briefly, whole neuroretina was peeled from the RPE layer, minced, and digested with collagenase I (Sigma–Aldrich). Cells and clusters were plated onto human fibronectin (Akron, Boca Raton, FL, USA)-coated flasks (Nunclon Delta, Waltham, MA, USA) in Ultraculture Media (Lonza, Walkersville, MD, USA), supplemented with 2 mM L-glutamine (Invitrogen, Carlsbad, CA, USA), 10 ng/mL recombinant human basic fibroblast growth factor (rh-bFGF) (Peprotech, Rocky Hill, NJ, USA), and 20 ng/mL recombinant human epidermal growth factor (rh-EGF) (Peprotech) in low-oxygen incubator (37°C, 3% O2, 5% CO2, 100% humidity). Fibronectin coating was accomplished by incubating the flasks with 10µg/mL fibronectin solution for 1 h at room temperature (1 mL of the solution per 10 cm2 of the surface), followed by single wash with deionized water. hRPCs were passaged at 80% confluency using TrypZean (Sigma–Aldrich), benzonase (EMD Chemicals, Darmstadt, Germany), and Defined Trypsin Inhibitor (Invitrogen). At each passage, cell number and viability were estimated with a hemocytometer, and cells were plated on a fibronectin-coated surface at a density of 20,000 cells/cm2 in medium. All further described work was performed with a single hRPC cell line at passage 9.

**hRPC adhesion and proliferation on PCL–IPM scaffolds**

hRPCs were plated at a density of 20,000 cells/cm2 in 60-mm Petri dishes containing the following materials: IPM–PCL scaffold, uncoated or fibronectin-coated PCL, tissue culture–treated plastic, IPM-coated plastic, or fibronectin-coated plastic, in 3 mL of medium. Fibronectin coating was accomplished as described above; for IPM coating, we incubated the flasks with the IPM suspension for 1 h. The dishes were then incubated for 30 min (37°C, 3%O2, 5% CO2, 100% humidity) to allow sufficient time for adhesion.24 After 30 min, all non-attached hRPC were collected, and each well was washed with 3 mL of Hank’s Balanced Salt Solution (HBSS) (with Ca++ and Mg++). Collected medium and washes from three wells per group were combined in the same tube and centrifuged at 400g for 5 min. The numbers of hRPC were counted with a hemocytometer, and the ratio of adhered cells was calculated. This experiment was performed three times (for independent IPM, PCL, and PCL–IPM preparations), and every time three independent Petri dishes were used for each condition. Mean and standard deviation (SD) were calculated for each group for the pooled data, and t-test analysis was performed to compare means.

Since we have previously observed a significant decrease in hRPC proliferation on PCL substrates, we were interested to see whether IPM incorporation leads to any change in this effect. To determine this, hRPCs were plated at a density of 20,000 cells/cm2 on PCL–IPM, PCL coated with fibronectin, or tissue culture plastic coated with fibronectin in the medium described above, with the medium changed every other day. At days 3 and 7, hRPC cell number was counted by harvesting all the cells as described above for passing procedure (TrypZean–benzonase). The population doubling (proliferation rate) was calculated based on three separate cell plating experiments. Mean and SD were calculated for each group and the results were compared by t-test.

**SEM**

To study the effect of incorporation of IPM on the hRPC morphology, we plated cells on fibronectin-coated PCL and PCL–IPM coated-cover slips in the medium described above. After 1 and 7 days, coverslips were washed, fixed, and dehydrated. The coverslips were fixed using 4% paraformaldehyde solution and dehydrated using sequential treatment with 35%, 50%, 70%, 95%, and 100% ethanol. After dehydrating with 100% ethanol, the samples were treated with hexamethyldisilazane (HMDS) and allowed to air dry after being attached to a specimen stub. Dried samples were sputter coated with 15-nm gold–palladium, and the hRPC morphology on PCL and PCL–IPM surfaces was observed using a scanning electron microscope (S-4800 Hitachi) at an operating voltage of 3.0 kV.

**Photoreceptor differentiation from hRPC on PCL–IPM scaffolds**

To investigate cell differentiation on PCL–IPM scaffolds, we plated hRPC in the medium described above at a density of 20,000 cells/cm2 on PCL–IPM scaffolds or PCL coated with fibronectin. The medium was replaced every other day. Seven days after plating, live cells were harvested with TrypZean for flow cytometry. hRPCs were fixed in Perm/Fix buffer (BD Biosciences) for 20 min at 4°C. Cells were then washed in Wash buffer (BD Biosciences) and incubated in block buffer (Pharmingen staining buffer with 2% goat serum) for 30 min at room temperature. The cells were stained with primary antibodies for 1 h at room temperature, washed, and stained with secondary conjugated antibodies for 30 min at room temperature. After the final wash, light
scatter and fluorescence signal from each cell were measured by Beckman Epics XL flow analyzer, Beckman Coulter, Brea, CA, USA (10,000 events were recorded). The results were analyzed with FlowJo software (Tree Star, Ashland, OR, USA). The ratio of positive cells within the gated population was estimated based on comparison with species-specific isotype control. For immunocytochemistry, hRPCs were collected after differentiation and replated onto 16-well chamber glass slides, coated with fibronectin. Antibodies used were as follows: anti-Pax6 (1:50; Hybridoma Bank, Iowa City, IA, USA), anti-Lhx2 (1:200; Chemicon, Billerica, MA, USA), anti-Nrl (1:50; Santa Cruz), anti-Crx (1:50; Santa Cruz, Santa Cruz, CA, USA), anti-Recoverin (1:1000; Chemicon), anti-Rhodopsin (1:200; Chemicon), and Anti-Rod Outer Membrane 1 (1:100; Abnova, Walnut, CA, USA).

Results

PCL and PCL–IPM production and characterization of scaffolds

IPM was successfully isolated from bovine eyes. IPM collection was straightforward as it dissociates from the retina when placed in deionised (DI) water,20 and numerous macroscopic sheets can be observed and collected. The IPM was successfully incorporated into the polycaprolactone (no chunks of the matrix were observed after mixing with PCL) to produce two types of scaffolds (Figure 1(a)): coated Petri dishes, which eliminate the need to fix the PCL film inside the culture vessel for culture and analysis, and thin films, which can be further used as cell carriers for transplantation purposes.

As was demonstrated by staining with PNA, IPM honeycomb structure observed in collected material (Figure 1(e) and (h)) is lost during PCL–IPM preparation (Figure 1(f)). SEM (Figure 1(b) and (d)) revealed pits in both PCL and PCL–IPM, varying in size from 1 to 7 µm. These pits likely appear during evaporation of the solvent (dichloromethane) from the scaffold. There were no identifiable differences between the topography of PCL and PCL–IPM scaffolds.

hRPC adhesion and proliferation on PCL–IPM scaffolds

IPM incorporation slightly changed the hydrophobic properties of the scaffold, as reflected by mean contact angles of 64.5 in PCL versus 60.2 in PCL–IPM (Figure 2(b)). Although this moderate change was significant, it is unlikely to be the only mechanism causing a profound increase in hRPC adhesion (Figure 2(a)) to the hybrid scaffolds. Coating of plastic with IPM by centrifugation led to a slight increase in adhesion, but we were not able to maintain IPM attachment to the PCL during culture. The inhibitory effect of polycaprolactone on hRPC proliferation did not change (Figure 2(c)) after IPM incorporation: population number did not increase on polycaprolactone scaffold with IPM to a greater extent than on PCL alone. Due to high proliferation rate, it is not possible to get non-confluent monolayer of hRPC on fibronectin-coated plastic—cells reach overconfluent state, start to form clumps, and detach, resulting in inconsistent cell count.

SEM

SEM analysis (Figure 3) showed that hRPCs attach and grow processes on the surface of PCL (Figure 3(a) and (b)) and hybrid PCL–IPM (Figure 3(c) and (d)) films. We have observed long cellular processes on all preparations. One-week culture led to stratification (increased area) of cells (Figure 3(b) and (d)), which may be related to additional ECM production and deposition.

Photoreceptor differentiation from hRPC on PCL–IPM scaffolds

One-week culture of hRPC on the PCL scaffolds (both fibronectin-coated and enriched with IPM) led to a significant decrease in the expression of early progenitor markers, such as Pax6 and Lhx2 (Figure 4(a)), while photoreceptor markers (Crx, Nrl, Rhodopsin, Rom1, Recoverin) were upregulated. The immunocytochemical analysis (Figure 4(b)) confirmed the differentiation of hRPC into rod photoreceptors. We did not identify any differences in differentiation pattern between PCL–IPM and PCL coated with fibronectin groups, which suggests that IPM provides stimuli sufficient for photoreceptor differentiation.

Discussion

Recent advances in retinal regeneration have focused on retinal progenitor and photoreceptor precursor cell transplantation. Several approaches have been investigated,25 including suspension26 and composite graft13 transplantation. Despite a complicated delivery procedure, composite graft transplantation is a promising strategy due to the structural organization and differentiation of cells. The scaffold of choice for such grafts should have properties similar to the ECM of the target tissue. This can be achieved with synthetic or biological polymers with biomimetic structures or chemical composition. One method of scaffold development which has demonstrated great potential is use of decellularized ECM27 for reconstruction of organs, including vessels,28 trachea,29 heart,30 lungs,31 and liver.32,33 These scaffolds not only provide support for the cellular component but also drive differentiation of progenitors and serve as a meshwork for new ECM deposition.

In previous studies with IPM,31,34 some benefits and limitations have been noted. IPM is the matrix surrounding the outer segments of photoreceptors and is naturally acellular upon collection as described above (as noted by...
nuclear staining, data not shown). However, in contrast to decellularized bone,\textsuperscript{35} blood vessels,\textsuperscript{28} trachea,\textsuperscript{29} heart,\textsuperscript{30} tooth bud,\textsuperscript{36} or lacrimal gland, IPM does not possess sufficient mechanical integrity to be easily handled, although we were able to collect it as a suspension. It is possible to use these suspended IPM particles to coat different surfaces by centrifugation, although they tend to detach from the surface, limiting experimental studies. These water-insoluble IPM particles can be suspended in organic solvents, such as dichloromethane or chloroform (trichloromethane), also used for PCL preparation.

This has allowed us to functionalize scaffolds previously described for retinal tissue engineering. Based on our experience with retinal progenitor cells and different types of synthetic polymers\textsuperscript{14} (PLGA/poly-l-lactic acid (PLLA),\textsuperscript{9} PMMA,\textsuperscript{10} PGS,\textsuperscript{37} and PCL\textsuperscript{38}), we have focused in these studies on PCL due to its biocompatible, biodegradable, and physical properties. We have previously shown that microtopography properties of PCL have only a slight effect on hRPC differentiation.\textsuperscript{39}

Although we were able to incorporate IPM, as shown by positive staining for lectin,\textsuperscript{40} the honeycomb structure of IPM was lost during this process. The preservation of

Figure 1. PCL and PCL–IPM plates and films. The manufacture of polycaprolactone in 10-cm Petri dish format (a, left) allows the scale up of the cell differentiation, eliminating the need of fixing the thin film (a, right) on the plate. SEM analysis (b–d) on different magnifications has not revealed any differences in the structure of the PCL and PCL–IPM scaffolds. Honeycomb structure of lectins revealed by PNA staining (green) in freshly isolated IPM (e, h) was not preserved during PCL–IPM scaffold preparation, resulting in diffuse staining (f). PCL without any additives (g) was used for control. Bars are 100 µm.

PCL: poly(ε-caprolactone); IPM: interphotoreceptor matrix; SEM: scanning electron microscopy, PNA: peanut agglutinin.
this structure is one of the aims for future studies, since it is suggested to be critical for outer segment homeostasis and photoreceptor–RPE interaction. IPM incorporation into PCL leads to a decreased contact angle and increased hRPC adhesion. Although the adhesion is lower compared to fibronectin-coated surfaces, it is sufficient to achieve retinal progenitor cell survival and differentiation. PCL–IPM inhibits proliferation of hRPC in the same manner as PCL. This inhibition of proliferation is required for differentiation and maturation of photoreceptors. After 1 week on PCL–IPM, retinal progenitor cells obtained photoreceptor phenotype and functional properties. We observed expression of the photoreceptor-specific transcription factors (Crx—both rods and cones; Nrl—rods) and visual pigment rhodopsin together with structural protein ROM1. The differentiation pattern of hRPC was similar on PCL–IPM and fibronectin-coated scaffolds.

We suggest that the designed scaffold can be readily used as a bioreactor for the differentiation of photoreceptors and photoreceptor precursors, although its utility for transplantation has to be addressed in future studies, primarily due to the immunogenicity of IPM components, including interphotoreceptor retinoid-binding protein.

**Conclusion**

Here we have described an approach to functionalize PCL biopolymer with a water-insoluble IPM. This approach overcomes both the lack of cell adhesion, specific for hydrophobic PCL, and water insolubility of IPM, which complicates its use in vitro. We have shown that the resulting composite scaffolds drive the hRPC fate toward photoreceptors, allowing one to create a graft for subretinal transplantation and retinal regeneration, although the immune component of IPM transplantation should be addressed. We suggest that incorporation of ECM into PCL film may be a useful approach if composite sheet grafts are required, as it is in case of epithelium or neuroepithelium regeneration.
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Declaration of conflicting interests
The authors declare that there is no conflict of interest.

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Figure 4. Differentiation of hRPC on PCL–IPM. (a) Flow cytometry analysis showed the downregulation of early eye development transcription factors Pax6 and Lhx2 in hRPC after 7 days of differentiation on both PCL coated with fibronectin (blue) and PCL–IPM (green) versus undifferentiated hRPCs (red). These results correlated with increase in photoreceptor-specific transcription factors (Crx and Nrl) and rod-specific rhodopsin and ROM1. Isotype controls are shown in black. (b) Immunocytochemical analysis of hRPC after differentiation confirmed the flow cytometry findings with individual cells expressing rhodopsin and ROM1 in polarized manner.

hRPC: human retinal progenitor cell; PCL: poly(ε-caprolactone); IPM: interphotoreceptor matrix.
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