Endothelial Protein C–Targeting Liposomes Show Enhanced Uptake and Improved Therapeutic Efficacy in Human Retinal Endothelial Cells

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Diabetic retinopathy is the most common microvascular complication of diabetes mellitus and corresponds to the leading cause of visual impairment in adults between 20 and 65 years old in the developed world.1,2 Its hallmark features include weakening of tight junctions, formation of microaneurysms on the retinal capillaries, pericyte loss, endothelial cell apoptosis, low cytotoxicity, and a high loading capacity for phospholipids and, frequently, cholesterol, have a range of attractive properties as drug carriers, including ease of assembly, low cytotoxicity, and a high loading capacity for both hydrophobic and hydrophilic drugs.6,12 EPCR was initially thought to be expressed only on the endothelium of large vessels but was then found on smooth muscle cells, neutrophils, keratinocytes, and microvascular endothelia such as that of the brain capillaries.13–15 To the best of our knowledge, it has not previously been identified on human retinal endothelial cells (HRECs).

In this study, we investigated the expression of EPCR on HRECs and the potential of this receptor as a target for drug nanocarriers by using unilamellar liposomes as a model system. Liposomes, uni- or multi-lamellar vesicles comprised of phospholipids and, frequently, cholesterol, have a range of attractive properties as drug carriers, including ease of assembly, low cytotoxicity, and a high loading capacity for both hydrophobic and hydrophilic drugs.16,17 Liposomes have a history of clinical approval18 and have demonstrated promising intraocular results (e.g., sustained drug release, high therapeu-
tic efficacy, and reduced cytotoxicity of the drugs.\textsuperscript{19,20} Here, we report EPCR expression on HRECs in comparison to human aortic endothelial cells (HAECs) and human umbilical vein endothelial cells (HUVECs). We determined the targeting efficiency of functionalized liposomes with EPCR-specific monoclonal antibodies on HRECs and HAECs. Furthermore, we assessed the anti-inflammatory efficacy of EPCR-targeting liposomes when loaded with the corticosteroid prednisolone (either prednisolone 21-hemisuccinate sodium salt [PH] and/or prednisolone 21-palmitate [PP]) through the inhibition of IL-8 and IL-1β and IL-6 expression in stimulated HRECs, as well as the ability to inhibit angiogenesis in vitro using an endothelial cell tube formation assay.

**Materials and Methods**

**Materials**

Cholesterol (Chol), 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC), 1,2-distearyloxy-sn-glycero-3-phosphoethanolamine-N-(methoxy)[polyethylene glycol]-2000 (PEG) and 1,2-distearyloxy-sn-glycero-3-phosphoethanolamine-N-(maleimide)[polyethylene glycol]-2000 (maleimide) were all purchased from Avanti Polar Lipids (Alabaster, AL, USA). 1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine-N-lissamine (DPPE-Atto) was purchased from Atto-TEC (Siegien, Germany). Prednisolone, PH, mannitol, HEPES, NaCl, N,N′-dicyclohexylcarbodiimide, 4-(dimethylamino)pyridine, palmitic acid, and all organic solvents were purchased from Sigma-Aldrich Corp. (St. Louis, MO, USA). ELISA kits for IL-1β, IL-6, and IL-8, along with the Human Cytokine Array kit were obtained from R&D Systems Inc. (Minneapolis, MN, USA), and the Matrigel matrix was purchased from Corning (Flintshire, UK). The monoclonal antibodies Alexa Fluor 488 rat anti-human CD201 clone RCR-252, Alexa Fluor 488 rat IgG1 κ isotype control clone R3-34, and rat anti-human CD201 clone RCR-252 (anti/EPCR) were purchased from BD Biosciences (San Jose, CA, USA).

**PP Synthesis, Purification, and Characterization**

The synthesis of PP was achieved using Steglich esterification. Prednisolone, palmitic acid, N,N′-dicyclohexylcarbodiimide, and 4-(dimethylamino)pyridine (1:2:2 molar ratio) were all dissolved in excess dichloromethane under an N\textsubscript{2} atmosphere and left stirring at room temperature for 18 hours. The reaction was followed by thin-layer chromatography using a chloroform:methanol (95:5) mobile phase. PP appeared as a streaked spot close to the baseline. Dichloromethane was evaporated under rotation and reduced pressure to leave a crude product. Crude product was redisolved in a chloroform:methanol (95:5) mixture and purified on a silica gel column by using a chloroform:methanol (95:5) mobile phase. The resulting purified PP had a high yield (>85%) and high purity (>97%). Purification was determined by both analytic high pressure liquid chromatography (HPLC) and \textsuperscript{1}H nuclear magnetic resonance (NMR) spectroscopy. \textsuperscript{1}H NMR (CHCl\textsubscript{3}) key characteristic peaks include the following: δ 0.88 (t, 3H, 37), 0.98 (s, 3H, 18), 1.25 (s, 3H, 25–36), 1.45 (s, 3H, 19), 4.48 (m, 1H, 11), 4.87 (d, 1H, J = 17.4 Hz), 21.497 (d, 1H, J = 17.4 Hz), 21.600 (t, 1H, 4), 6.27 (dd, J = 10.09, 1.89 Hz), and 7.27 (dd, J = 10.1 Hz, 1). The remaining corticosteroid ring \textsuperscript{1}H chemical shifts followed similar values (e.g., δ 1.6–2.6 covers protons at 6, 7, 8, 12, 14, 15, 16, and 23) as previously reported for prednisolone, similar corticosteroids, and functionalized corticosteroids where functionalization occurred at position 21 (see Supplementary Fig. S1 for \textsuperscript{1}H NMR spectrum and atom numbering).\textsuperscript{21–23}

**Liposome Preparation and Loading**

All liposomes were prepared by mixing the lipids in a tertiary butanol:water (9:1) mixture and lyophilized overnight in a Christ Epsilon 2-4 LSCplus freeze dryer (Buch & Holm, Herlev, Denmark). Subsequently the lipid mixtures were rehydrated with 10 mM HEPES saline buffer or with a 200 mM calcium acetate buffer (Sigma-Aldrich Corp.) for the remote loading of PH. Lipid suspensions were prepared at 15 mM and hydrated at 60°C with gentle vortexing for 60 minutes, followed by extrusion through a 100-nm filter at 60°C.\textsuperscript{24} For the liposome formulations with membrane-loaded PP, the extrusion was performed in 3 steps through 400-nm, 200-nm, and finally through 100-nm filters. Remote loading was performed using a protocol previously described.\textsuperscript{25,26} Briefly, the liposomes were subjected to 3 repeated 1-hour dialysis cycles by using a ratio of liposome dispersion to dialyzing medium (5% glucose) 1:200, followed by a forth dialysis step overnight of 1:400. PH was dissolved in 5% glucose and was mixed with liposomes in a drug to lipid ratio of 1:2. Remote loading was achieved by incubation of the liposomes with the drug for 20 minutes at 60°C to 65°C, with continuous stirring. Nonencapsulated drug was removed by 2-step dialysis (dialysis tubing, MWCO 10 kDa; Sigma-Aldrich Corp.) using a ratio of 1:500 sample to diluent (10 mM HEPES, pH 7.4, 150 mM NaCl), with the first step lasting for 4 hours at 4°C and the second step overnight at 4°C. For the uptake studies, the different systems that we prepared were of the following molar ratios: (1) DPPC/Chol/PEG/maleimide/DPPE-Atto, 55:92:40:3:1:0.08, (2) DPPC/Chol/DPPE-Atto, 59:92:40:0:0.08, and (3) DPPC/Chol/PEG/DPPE-Atto, 55:92:40:4:0.08. For the flow cytometry experiments, DPPE-Atto 655 was used and for the microscopy images, DPPE-Atto 488 was used. We did not observe any size or charge differences of the liposomal carriers when replacing fluorophores. For the efficacy studies, we removed fluorophore-conjugated phospholipids so the formulations for these experiments were DPPC/Chol/PEG/maleimide (56:40:3:1) and DPPC/Chol/PEG (56:40:4), and we included an extra formulation of DPPC/Chol/PEG/maleimide/PP (51:40:3:1:5) with a 5% molar concentration of PP in the bilayer. See Tables 1 and 2 for all liposome formulations.

**Liposome Characterization**

The diameter and surface charge of the liposomes were measured on a Brookhaven ZetaPALS zeta potential analyzer (New York, NY, USA) as previously described.\textsuperscript{27} Briefly, diameters (measured by dynamic light scattering) and zeta potentials were measured in 10 mM HEPES in a 5% glucose solution. Phospholipid concentrations were determined by quantifying the phosphorous content in the liposome samples by using inductively coupled plasma mass spectrometry. For further confirmation of the liposome size distribution measured by dynamic light scattering, we used Cryo-transmission electron microscopy (TEM). For each experiment, 3 μL of lipid dispersions at a concentration of 3 mM was placed on a lacy carbon 300 mesh copper TEM grid, blotted and plunged frozen in liquid ethane by using a FEI Vitrobot Mark IV (Thermo Fisher Scientific, OR, USA). Samples were transferred to the TEM at –175°C by using a Gatan 626 single tilt cryo-transfer holder (Gatan, Inc., Pleasanton, CA, United States) and were imaged using a FEI Tecnai G2 20 TWIN TEM operated at 200 keV in low dose mode with a FEI high-sensitive 4k × 4k Eagle camera. The loading of both PH and PP was quantified by reverse phase HPLC, and the loading efficiency was calculated by the drug-to-lipid ratios before and after purification.
**Liposome Functionalization**

Rat anti-human CD201 clone RCR-252 and isotype IgG control (rat IgG1; BD Biosciences) antibodies were diluted to a final concentration of 0.45 mg/mL in 0.2 M sodium borate buffer (pH 8.5) mixed with 2-iminothiolane (Traut's reagent; Sigma-Aldrich Corp.) at a molar ratio of 1:20. The thiolation reaction proceeded for 60 minutes at room temperature, and the thiolated antibodies were then transferred to Amicon Ultra 4-mL spin filters (MWCO 30 kDa; Merck Life Science, Søborg, Denmark). The spin filters were filled with cold 10 mM HEPES buffer and centrifuged at 2500g for 30 minutes at 4°C. This process was repeated 2 times to purify the thiolated antibodies from unreacted 2-iminothiolane. The thiolated antibodies were immediately mixed with DSPE-PEG-maleimide liposomes in a molar ratio of 10:1 (of the available maleimide groups) and the liposome-antibody mixture was incubated overnight on a rocking table (180 rpm) in the dark to allow for conjugation. After the incubation, the antibody-functionalized liposomes were separated from the nonbound antibodies by size-exclusion chromatography using a Sepharose CL-4B column (Sigma-Aldrich, Corp.) with a 10 mM HEPES, 150 mM NaCl (pH 7.4) buffer as the fluid phase. The size-exclusion chromatography fractions were analyzed by the BCA assay and at 4°C.

**Release Profile In Vitro**

The dialysis method was used to characterize the in vitro release behavior of the PH-loaded liposomes. We loaded 1 mL of 6 mM liposomes into dialysis bag (MWCO 10 kDa) and incubated them in 9 mL of 10 mM HEPES saline buffer at 37°C with continuous stirring for 120 hours. At predetermined timepoints, 200 μL of release media was sampled and equal volumes of fresh media was added. The cumulative release of PH was calculated after analyzing the samples by HPLC (n = 4).

**Cell Culture**

HRECs were purchased from Innoprot (Bizkaia, Spain), and the HAECs and HUVECs were purchased from LGC Standards GmbH (Wesel, Germany). The cells were grown to 85 to 90% confluence in the endothelial cell medium (Innoprot) supplemented with 5% (vol/vol) fetal bovine serum and 1% (vol/vol) penicillin/streptomycin in an environment of 95% O₂ and 5% CO₂. For the endothelial cell tube formation assay, the cells were cultured on Matrigel matrix (Corning). 96-well plates were coated with Matrigel (50 μL/well) at 4°C and were incubated at 37°C for 30 minutes. Subsequently, 15 × 10³ HRECs were seeded in each well with a final volume of 200 μL endothelial cell medium.

**Cell Viability (MTS Assay)**

Cell proliferation and viability were evaluated by an MTS assay. HRECs, HAECs, and HUVECs were seeded into 96-well plates at a density of 3 × 10⁵ cells per well for 100 μL. Twenty-four hours later, the cells were exposed to PH at concentrations from 1 nM to 1 mM for 4 hours. The cells were incubated for 48 hours and then 20 μL MTS reagent was added to each well. The absorbance at 490 nm was measured by a microplate reader 4 hours later. The same process was followed in order to determine the cytotoxicity of our liposomal system. The cells were incubated with liposomes at lipid concentrations of 100 nM to 500 μM.

**Detection of EPCR in HRECs by Flow Cytometry and Western Blot**

In order to determine whether HRECs express EPCR or not, we used flow cytometry and Western blot assays. For the flow cytometry, HRECs, HAECs, and HUVECs were seeded in T75 flasks (Thermo Scientific) and grown to 85 to 90% confluency in the endothelial cell medium (Innoprot) supplemented with 5% (vol/vol) fetal bovine serum and 1% (vol/vol) penicillin/streptomycin in an environment of 95% O₂ and 5% CO₂. When the cells reached confluency, they were washed in PBS, 3 mL of trypsin-EDTA was then added to each flask, and flasks were incubated for 5 minutes. The cell suspensions were centrifuged (2000g, 20°C, 5 minutes), the collected cell pellet was resuspended in cold PBS, and this process was repeated once more. The washes were collected in Eppendorf (Horsholm, Denmark) tubes with a concentration of 5 × 10⁶ cells per tube per 500 μL of PBS. Cells were then incubated with 10 μg/mL of Alexa Fluor 488 rat anti-human CD201 clone RCR-252 or Alexa Fluor 488 rat IgG1 κ isotype control clone R3-34 for 30 minutes at 4°C (BD Biosciences). The cell suspensions were washed twice with PBS, transferred to flow cytometry tubes, and analyzed using a BD Accuri C6 autosampler flow cytometer (BD Biosciences). In each experiment, 20,000 events were analyzed and at least 3 independent experiments were performed.

For the Western blot assay, the HRECs and the HAECs were seeded in T25 flasks (Thermo Scientific) and grown to 85 to 90% confluence. The cells were then washed twice with PBS, transferred to cold PBS, resuspended in cold lysis buffer, and washed 2 times with PBS. The cell suspensions were centrifuged (12,000g, 30°C, 5 minutes) and the collected cell pellets were resuspended in lysis buffer. The lysates were clarified by centrifugation (16,000g, 30°C, 10 minutes) and stored at −80°C. The protein concentrations were determined using the BCA assay and at least 3 independent experiments were performed.

**Table 2.** Liposome Formulations and Characteristics Used for the Efficacy Studies

<table>
<thead>
<tr>
<th>Formulations</th>
<th>Abbreviation</th>
<th>Size, nm</th>
<th>PDI</th>
<th>Zeta Potential, mV</th>
</tr>
</thead>
<tbody>
<tr>
<td>DPPC:Chol:PEG:DPPE-Atto (56:40:4:0)</td>
<td>P-NL</td>
<td>149.5 ± 1.1</td>
<td>0.04 ± 0.3</td>
<td>−16.82 ± 1.12</td>
</tr>
<tr>
<td>DPPC:Chol:PEG:maleimide-anti/EPCR (56:40:3:0)</td>
<td>EPCR-NL</td>
<td>162.7 ± 0.9</td>
<td>0.05 ± 0.03</td>
<td>−14.19 ± 1.32</td>
</tr>
<tr>
<td>PP:DPPE-PEG:maleimide-anti/EPCR (55:40:3:1)</td>
<td>EPCR-NL</td>
<td>101.2 ± 1.3</td>
<td>0.03 ± 0.01</td>
<td>−15.49 ± 1.71</td>
</tr>
<tr>
<td>DPPC:Chol:PEG:maleimide-anti/EPCR (56:40:3:1)</td>
<td>EPCR-NL</td>
<td>160.3 ± 1.5</td>
<td>0.03 ± 0.02</td>
<td>−15.93 ± 1.65</td>
</tr>
</tbody>
</table>
90% confluence. For transfection, when cells reached 70 to 80% confluency, they were incubated with EPCR/PROCR silencer predesigned siRNA (Thermo Fisher Scientific) Lipofectamine 2000 (Thermo Fisher Scientific) complexes. Briefly, 500 pmol of siRNA was diluted in 250 μL Opti-MEM I reduced serum medium and 5 μL Lipofectamine 2000 was diluted in 250 μL of Opti-MEM I reduced serum medium, and both solutions were incubated for 15 minutes at room temperature. After the 15 minute incubation, the diluted siRNA and the diluted Lipofectamine 2000 were combined and incubated for 15 minutes at room temperature so as to allow complexes to form. Subsequently, the siRNA-Lipofectamine 2000 complexes were added to the T25 flasks at a total of 5 mL medium and incubated at 37°C in a humidified CO2 incubator for 24 hours. Then, the cells were washed 2 times with cold PBS, lysed with lysis buffer (25 mM Tris-HCl, pH, 7.4; 0.15 M NaCl; 1% Triton X-100) with complete Mini Protease Inhibitor Cocktail (Sigma-Aldrich Corp.) and Phosphatase Inhibitor Cocktail 2 (Sigma-Aldrich Corp.), then placed on a shaker at 4°C for 30 minutes. After centrifugation at 12,000g for 10 minutes, the protein concentration of the supernatant was measured by BCA assay (Thermo Fisher Scientific). Subsequently, equal amounts of 20 μg of protein were loaded and separated by 4 to 12% Bis-Tris gel electrophoresis (Thermo Fisher Scientific). The proteins were transferred to a nitrocellulose membrane (Thermo Fisher Scientific). Following the transfer, the immunoblot was incubated with Odyssey blocking buffer (LI-COR Biosciences, Cambridget, UK) at room temperature for 1 hour with agitation. EPCR was detected by monoclonal rat anti-EPCR/CD201 (Cambridg, UK) at room temperature for 1 hour, 2 hours, or 3 hours. For the experiments where we incubated the Atto 655–labeled carriers at a liposomal concentration of 75 μg of rat anti-human CD201 clone RCR-252 (Abcam, Cambridge, UK) at 1:50 dilution in which the membrane was incubated overnight at 4°C. The membrane was washed 3 times with tris-buffered saline and Tween 20 for 5 minutes, and then the second antibody (goat anti-rat IgG [H+L]; 1:300; Sigma-Aldrich) was added to each well and incubated for 1 hour at room temperature. Eventually, each well was washed twice with PBS, and confocal imaging was performed using a Leica TC SP5 confocal laser scanning microscope (Leica Microsystems GmbH, Wetzlar, Germany).

Quantification of Liposome Uptake by Cells Using Flow Cytometry

We evaluated the uptake of fluorescently labeled liposomes by endothelial cells with flow cytometry. HRECs, HAECs, and HUVECs were seeded at a density of 12 × 10^4 cells per well in 12-well plates (Thermo Scientific) in 1 mL of the culture medium at 37°C under an atmosphere with 5% CO_2. After 24 hours, the cells were incubated with the Atto 655–labeled carriers at a liposomal concentration of 75 μM in medium for 1 hour, 2 hours, or 3 hours. For the experiments where we blocked EPCR, we added 5 μg of rat anti-human CD201 clone RCR-252 in each well and incubated for 20 minutes. Then, we added the liposomes without washing off the antibody. Subsequently, the cells were washed with PBS and then trypsinized. The cell suspension was centrifuged (200g, 20°C, 5 minutes), and then the collected cells were resuspended in 1 mL PBS. The latter process was repeated 3 times, and then the cells were transferred to flow cytometry tubes and analyzed using the Gallios flow cytometer (Beckman-Coulter, Copenhagen, DK). Cells were gated based on the forward/side scatter plot to eliminate cell debris from the subsequent analysis. At least 10,000 events were analyzed in each experiment and a minimum of three independent experiments were performed. The fluorescence intensity of the liposome treated cells was corrected based on the autofluorescence of nontreated cells and the resulting data was analyzed using FlowJo software (FlowJo LLC, Oregon, USA).

Confocal Laser Scanning Microscopy

Further estimation of the uptake of the different carriers was performed by confocal laser scanning microscopy. In that case, 25 × 10^3 cells were seeded in a μ-Slide 8 well (ibidi GmbH, Planegg, Germany) in 300 μL of culture medium as a 2D culture or on Matrigel matrix. After 24 hours, the cells were incubated with 75 μM of the different Atto 488–labeled carriers for 3.5 hours. Subsequently, 20 mL of TO-PRO-3 (1 μM; Thermo Fisher Scientific) dye was added to stain the nuclei, and the suspension was reincubated for an additional 30 minutes. The medium was then removed, and the cells were washed twice with 300 μL PBS, before they were fixed with 4% paraformaldehyde for 10 minutes at room temperature and washed again twice with 300 μL of PBS. After fixation, and only for the 2D cultured cells, 300 μL of Phalloidin-TRITC in 10 mM HEPES (1:300; Sigma-Aldrich) was added to each well and incubated for 1 hour at room temperature. Eventually, each well was washed twice with PBS, and confocal imaging was performed using a Leica TC SP5 confocal laser scanning microscope (Leica Microsystems GmbH, Wetzlar, Germany).

Stimulation of Cells by High Glucose and Cytokine Quantification

To study the anti-inflammatory efficacy of our carrier, we stimulated an inflammatory response in the cells by culturing them in high glucose conditions. For these studies, only the primary cell lines were used. HRECs and HAECs were seeded in 12-well plates at a concentration of 30 × 10^3 cells/well under either normal glucose (5.5 mM), mannitol (20 mM; mannitol-osmolality control), or high glucose (25 mM) conditions for up to 3 days. Twenty-four hours after having been seeded in high glucose, the cells were treated with the following: (1) 25 μM prednisolone hemisuccinate in the form of free drug, (2) 25 μM prednisolone hemisuccinate encapsulated in the targeting liposomal system, with or without PreP on the membrane, or (3) 25 μM prednisolone hemisuccinate encapsulated in nontargeting liposomes (P-NL) for 4 hours, then washed and incubated at 37°C with 5% CO_2. Subsequently, we collected the cell supernatant 24 hours following the treatment, centrifuged them at 3000g for 5 minutes, and saved the samples at −80°C. New medium was added to the wells and incubated at 37°C with 5% CO2 for another 24 hours when we again collected the supernatant, 48 hours after treatment. Concentrations of IL-1β, IL-8, and IL-6 protein in culture supernatant were determined using an ELISA.

Endothelial Cell Tube Formation Assay

Angiogenesis was studied by assessment of tube formation on HRECs seeded onto Matrigel matrix (Corning), as has been previously described. After coating the 96-well plates with Matrigel, 15 × 10^3 HRECs were seeded in each well with a final volume of 200 μL. To determine the effect of the drug and the liposomes on the capillary tube formation, HRECs were incubated under 6 different conditions from the onset of the culture: (1) no treatment (control), (2) empty liposomes, (3) 50 μM prednisolone hemisuccinate as free drug, (4) 50 μM prednisolone hemisuccinate encapsulated in the targeting liposomal system, with or without PP, or (5) 50 μM prednisolone hemisuccinate in nontargeting liposomes. Phase-contrast images (5× magnification) of the center of each well were taken 4 hours, 8 hours, and 24 hours after seeding the cells, and tube formation was quantified using the Angiogenesis Analyzer tool for ImageJ (reported as the total length of the tubelike formations and the number of segments formed) (National Institutes of Health, Bethesda, MA, USA).
Cell Experiment Statistics

Data were expressed as mean ± SEM and a one-way ANOVA with a confidence level of 95% (α = 0.05), followed by a Tukey's multiple comparison post hoc test (P < 0.05) were used in order to analyze significant differences in the in vitro assays.

RESULTS

Characterization of Liposomes

Size distribution, polydispersity index (PDI), and zeta-potential were measured for each formulation (see Tables 1 and 2), and the structure of the empty and loaded liposomes was further characterized by cryo-TEM imaging (Fig. 1A). Liposome formulations without PP had diameters ranging from 150 to 165 nm, and from the cryo-TEM images, it was observed that most of the liposomes were unilamellar and spherical in shape. The precipitation of PH in the core of the liposomes was also observed. The formulation with PP had slightly smaller diameters of approximately 100 nm. The loading efficiency for PP was 45% (± 3.2%), whereas for PH was 70% (± 2.8%). Moreover, the cumulative release profile of PH for both antibody functionalized and nonfunctionalized liposomes showed sustained release for 5 days without any initial burst, but with a slightly higher release rate for the nonfunctionalized liposomes compared to the EPCR-targeting ones (Fig. 1B). PH release profiles from targeting and nontargeting liposomes could be fitted best with a first order exponential (P-NL1, R² = 0.969; EPCR-NL1, R² = 0.947), which indicates that the release rate of drug is dependent on the drug concentration.31

EPCR Targeting for Enhanced Retinal Drug Delivery

To determine whether or not HRECs express EPCR, we selected 2 endothelial cell types (HUVECs and HAECs) that have already shown strong expression of the receptor, and we compared all 3 with flow cytometry. From the analysis of these experiments, we observed a shift of 2 orders of magnitude in the median fluorescent intensity (MFI) levels of the HRECs that were incubated with the anti-EPCR monoclonal antibody, compared to the isotype-treated cells or the nontreated ones. This shift was similar to shifts observed for HAECs and HUVECs, which can be approximated as 99.9% of HRECs expressing EPCR at similar levels to the other 2 endothelial cell lines (Fig. 2A). See Supplementary Fig. S2 for fluorescence-activated cell sorting scatter plots.

The comparison between HRECs and HAECs through Western blots indicated similar immunoblotting for both endothelial cell types at 49 kDa and 38 kDa (Fig. 2B). The double band can be explained by the fact that EPCR has 4 N-linked glycosylation sites that can be heterogeneously modified, which results in a glycosylated mass of ≤49 kDa that...
varies among cell lines. As another approach to confirm the cell surface expression of EPCR from HRECs, we transfected HRECs with EPCR knockdown siRNA and immunoblotted the total protein extract (~20 μg) of both transfected and nontransfected HRECs. In this case, we can see strong binding of the anti-EPCR monoclonal antibody at 49 kDa and 38 kDa for HRECs cells, but for the transfected cells, there is a 2-fold reduction at the normalized signal of the 49 kDa band and a noticeable decrease at the 38 kDa band (Fig. 2C).

**EPCR-Targeting Effects on Cellular Uptake**

Once we had established the expression of EPCR on the surface of HRECs, we investigated the potential of targeting this receptor with antibody-functionalized liposomes. For the

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**FIGURE 2.** EPCR expression in endothelial cells. (A) Flow cytometry analysis of EPCR expression in HRECs. Cells were incubated with isotype control IgG or anti–EPCR monoclonal antibody (RCR-252), both conjugated with Alexa Fluor 488, and the autofluorescence of the cells is indicated from the nontreated ones. EPCR expression estimated by MFI determined by flow cytometry analysis from 3 independent experiments is shown on the left, and example flow cytometry histograms from 1 of the experiments on the right. (B) Immunoblotting of cell surface EPCR by RCR-252. Western blot (left) where lanes 1–3 represent HRECs and lanes 4–6 represent HAECs. Densitometry of Western blot data is shown on the right. (C) Western blot (left) where lanes 1–3 represent HRECs and lanes 4–6 represent transfected HRECs with EPCR knockdown siRNA. Densitometry of this is shown on the right.
first set of experiments, liposomes functionalized with either the EPCR-specific antibody or isotype were incubated for 4 hours with cells. As a second negative control, EPCR was blocked by incubating the cells with the monoclonal anti-EPCR antibody before adding the EPCR-targeting liposomes. EPCR-targeting liposomes associated strongly with both HRECs and HAECs, which corresponded to an 8.5-fold increase in the MFI compared to isotype IgG liposomes and a 5.2-fold increase compared to the cells with the blocked receptor (Fig. 3A). Subsequently, we wanted to compare the binding and uptake of the EPCR-targeting liposomes with other nontargeting liposomes, so we made nonfunctionalized/blank (B-NL) and P-NL liposome formulations. We then performed a time study in order to investigate how uptake evolves with different incubation times. Analysis of the flow cytometry data showed that the EPCR-targeting liposomes (EPCR-NL) displayed significantly higher uptake with the HRECs and HUVECs after 1 hour of incubation (Fig. 3B) than P-NL (3.2-fold) and B-NL (5.7-fold). The results for HAEcs were similar, with P-NL being higher than B-NL (2.3-fold) and P-NL (4.6-fold). Furthermore, we found that the longer the incubation time, the higher the uptake by cells with all the different formulations. However, EPCR-NL repeatedly showed more than 3-fold higher MFI compared to P-NL after 2 hours and 3 hours of incubation with HRECs and more than 6-fold higher than B-NL. Similar behavior was observed for HAEEs and HUVEEs. HAEEs showed 2-fold higher uptake of EPCR-NL than P-NL after both 2-hour and 3-hour incubations, whereas a 4-fold higher uptake of EPCR-NL was observed compared to B-NL. HUVEEs showed a more than a 3-fold higher MFI for EPCR-NL compared to the P-NL formulation (2-hour and 3-hour incubations) and >5.5-fold higher compared with the B-NL one (Fig. 3B). All results showed statistical significance with P values generally below 0.01.

The strong interaction between EPCR-targeting liposomes and the HRECs could also be visualized by confocal laser scanning microscopy. From Z-stack projections, we were able to detect strong fluorescent signal from the EPCR-targeting liposomes, mainly in the intracellular compartment of the HRECs (Fig. 4). In agreement with the flow cytometry results, the signal from the HRECs that were incubated with the P-NL formulation was much weaker (Fig. 4B) and in the case of the B-NL, no signal was detected (Fig. 4C).

**Reduction of ILs by Corticosteroid-Loaded EPCR-Targeting Liposomes**

To assess whether the enhanced uptake of EPCR-targeting liposomes by endothelial cells resulted in an improved therapeutic efficacy, we exposed HRECs and HAECs to corticosteroid-loaded liposomes, PH loaded EPCR-NL1 and dual-loaded (PP and PH) EPCR-NL2, and measured changes in IL levels. We first stimulated increased IL expression in HRECs and HAECs by culturing them in high glucose conditions. We then tested the changes in the secretion levels of 3 different inflammatory mediators with or without corticosteroid treatment. IL-6 and IL-8 levels were significantly increased under normal glucose conditions than P-NL1 in HRECs. This likely reflects the better uptake of P-NL by HAECs than HRECs (Fig. 3). We did not observe similar behavior for IL-1β and were not able to detect any difference in IL-1β secretion levels for any of the 3 different conditions for HRECs or HAECs (data not shown).

**Inhibition of In Vitro Angiogenesis by Corticosteroid-Loaded EPCR-targeting liposomes**

In another attempt to explore the potential efficacy of targeting EPCR, we tested the ability of EPCR-NL1 to inhibit angiogenesis in vitro in an endothelial cell tube formation assay. Before the efficacy studies, we tested the uptake of our liposomes by HRECs in tube-like formations on Matrigel matrix. The results were similar to what we previously observed from the uptake studies in monolayer culture conditions: more specifically, we detected strong fluorescent signal from EPCR-NL1, but weaker signal from P-NL and no signal from B-NL (Fig. 6A–D). The ability of HRECs to form tube-like structures was estimated by calculating the number of segments and the total length of the tube formations from images taken 8 hours after seeding the cells on Matrigel. Incubating the cells with EPCR-NL1 and EPCR-NL2 liposomes significantly inhibited tube-like formations compared to untreated cells, as shown by the decrease in the total tube length ($P < 0.05$) and the number of segments ($P < 0.05$ and $P < 0.01$) (Fig. 6E, 6F). Using the PH release profiles (Fig. 1B) at 8 hours, we extrapolated that 7.8% of PH has been released by ECPR-NL, corresponding to 3.9 μM. This is significantly lower than the 50 μM of free PH required to induce a similar effect on both tube length and the number of segments (Fig. 6E, 6F). We did not observe any reduction in
FIGURE 3. Flow cytometry evaluation of liposome uptake in endothelial cells. (A) Flow cytometry evaluation of the association between EPCR-targeting liposomes and endothelial cells. HRECs and HAECs were incubated with either EPCR-targeting liposomes (EPCR-NL), isotype IgG functionalized liposomes (Iso-NL), or with EPCR-specific antibody (blocked EPCR) before adding the EPCR-targeting liposomes in order to block the receptor. All MFI values are normalized to the MFI of the EPCR-NL. (B) Flow cytometry assessment of the liposome uptake by endothelial cells. HRECs, HAECs, and HUVECs were incubated with either EPCR-NL, P-NL, or B-NL liposomes for 3 different incubation times: 1 hour, 2 hour, or 3 hour. **P < 0.01, ***P < 0.001, ****P < 0.0001.
tube length or the number of segments at 25 μM of free PH (data not shown). Furthermore, P-NL1 treatment did not induce any significant reduction in tube length or the number of segments, even though PH release is similar to EPCR-NL (7.9%). This indicates that the enhanced uptake of EPCR-targeting liposomes leads to the improved efficacy of prednisolone.

**DISCUSSION**

In this study, we showed for the first time that EPCR is expressed by HRECs. It has been previously shown through histologic studies in humans and baboons that all large vessels and arteries express high levels of EPCR. In the brain microvasculature, which can be used as direct comparison to the retina, the expression was lower. The latter observation provides a strong argument that EPCR should be expressed in the human retinal endothelium as well. In an attempt to realize the level of EPCR expression, we performed a comparative study between HRECs and 2 other endothelial cell types, HEACs and HUVECs, which have been shown to heavily express the receptor in vitro. The flow cytometry and immunoblotting assays indicated similar levels of EPCR expression in all 3 different cell types, leading to the conclusion that HRECs strongly express EPCR in vitro.

In order to investigate the potential of EPCR as a nanomedicine target, we functionalized EPCR-specific antibody onto the surface of liposomes to study the uptake of these liposomes by HRECs, HAECs, and HUVECs in comparison to 2 nontargeting formulations. All of our formulations used saturated lipids (DPPC and DPPE-Atto), which, due to higher transition temperatures compared to unsaturated lipids, tend to exhibit better stability. In our case with P-NL formulations, we obtained a sustained PH release for up to 5 days, which was not the case for unsaturated formulations that exhibited burst release within the first 24 hours (data not shown). We also incorporated cholesterol into all of our formulations, as it has been previously shown that adding 30 to 40 mol% of cholesterol can significantly improve the stability of liposomes. Functionalizing the surface of the liposomes with PEG is a widely used technique in order to not only improve the circulation/residency time of nanocarriers but also to improve nanocarrier mobility in the vitreous. We chose to PEGylate our EPCR-targeting formulation due to the benefits of PEG, but we also used a formulation with no surface

**Figure 4.** Confocal laser scanning microscopy of EPCR-targeting liposomes and HRECs: (A) HRECs incubated with EPCR-NL liposomes, (B) HRECs incubated with P-NL liposomes, (C) HRECs incubated with B-NL liposomes. The first column shows actin staining (Phalloidin-TRITC), the second column shows nucleus staining (TO-PRO-3), the third column shows liposomes (Atto 488), and the fourth column shows columns 1–3 merged. Scale bar: 25 μm.
functionalization (B-NL) to investigate how PEGylation would affect the uptake of the liposomes by the cells.

In all the experiments, the EPCR-targeting liposomes displayed significantly higher uptake than the control formulations, even for incubation times as short as 1 hour. In order to establish that it was actually EPCR targeting that led to the enhanced uptake, EPCR was blocked before incubating with the EPCR-targeting liposomes. This blockage lead to a 5-fold inhibition of the uptake of the carrier with both HRECs and HAECs (\( P < 0.0001 \)), clearly indicating that EPCR targeting drove the enhanced uptake.

The delivery of corticosteroid drugs to the posterior part of the eye is a challenge because systemic administration is associated with undesirable systemic side effects and eye drop formulations have poor penetration. In contrast, injection of corticosteroid into the vitreous is very effective in controlling inflammation and macular edema secondary to diabetic retinopathy and retinal vein occlusion. Diabetic retinopathy is closely linked to hyperglycemia, low-grade chronic inflammation, and elevated reactive oxygen species and is dominated by the development of microaneurysms, hemorrhage, vascular leakage, macular edema, and preretinal neovascularization at the back of the eye. These findings have led to the emergence of several novel treatment modalities based on the administration of antivascular endothelial growth factor agents and corticosteroids to the diseased retina. Corticosteroids are an attractive option due to their broad range of actions including anti-inflammatory and antiangiogenic effects. Corticosteroid-loaded liposomes have been reported for several decades, but drug loading efficacy and release rates have presented significant challenges. The passive loading of hydrophilic corticosteroids in liposomes initially resulted in very low encapsulation efficiencies (<5%). Low loading can be overcome by choosing amphipathic weak acid corticosteroids and using a remote-loading approach. It has been shown that using a calcium acetate ion gradient can lead to high loading efficiencies of corticosteroid succinates, with 80% encapsulation efficiency (EE) for betamethasone hemisuccinate, 100% EE for hydrocortisone hemisuccinate, and 95% EE for methylprednisolone hemisuccinate. In our case, we achieved a 70% EE for PH that is lower compared to the other corticosteroids, and this could likely be improved by optimizing the loading parameters. Furthermore, palmitate corticosteroid derivatives have also been studied since the late 1970s, with cortisol palmitate, prednisolone palmitate, and dexamethasone palmitate having been loaded into liposomes. In our case, we loaded 5% molar concentration of PP into our liposomes with a
45% EE, which is low compared to what has previously been reported (up to 99% EE). However, these high EEs are predominantly associated with large multilamellar liposomes and not small unilamellar liposomes where extrusion results in the structural rearrangement of the liposome bilayer, leading to drug loss. In order to compensate for the loss of corticosteroid, we developed a dual-loaded liposome (EPCR-NL2) to see if this resulted in an improved anti-inflammatory effect.

Figure 6. Liposome influence on HREC tube formation. (A) HRECs incubated with EPCR liposomes, (B) HRECs incubated with P-NL liposomes, (C) HRECs incubated with B-NL, and (D) nontreated cells. Nuclei were stained with TO-PRO-3 and liposomes labeled with Atto 488. Scale bar: 250 μm. Quantification of tube formation by assessment of (E) the total tubule length and (F) the number of segments 8 hours after seeding the cells on Matrigel matrix. At this timepoint ~4 μM of PH has been released from the liposomes. *P < 0.05, **P < 0.01, ***P < 0.001.)
Cells cultured in high glucose conditions have previously shown to have an inflammatory response.\(^{57-59}\) Corticosteroids have repeatedly demonstrated inhibition of proinflammatory cytokines in endothelial cells,\(^{58,60-62}\) and so we compared the effect of prednisolone-loaded EPCR-targeting liposomes (EPCR-NL1 and EPCR-NL2), nontargeting liposomes, and free drug on endothelial cells. The outcome of these studies indicated that EPCR-targeting liposomes and free drug inhibited the secretion of cytokines, with the greatest impact being associated with EPCR-NL2 for both HRECs and HAECs. These findings are in accordance with older studies that have shown that \(\mu\)M concentrations of dexamethasone can induce and approximately 50\% decrease in the secretion of IL-6 and a 43\% decrease in IL-8 by stimulated endothelial cells,\(^{64,65}\) whereas cortisol was reported to inhibit IL-6 by approximately 40\%.\(^{64}\) Moreover, the performance of the EPCR-targeting liposomes was superior to nontargeting liposomes, which could be explained by the significantly higher uptake by HRECs and HAECs. Glucocorticoid receptors are found mainly in the cytoplasm\(^{65}\) and delivering drug payloads into cells should maximize drug-receptor binding. We observed that EPCR-targeting liposomes with significantly lower PH concentrations (~5 \(\mu\)M per day) resulted in similar reductions in IL-6 and IL-8 expression as a single dose of free PH (25 \(\mu\)M). This observation assumes that all the liposomes were endocytosed by cells, which is unlikely. The concentrations of PH released from the liposomes are therefore likely to be lower. Treating stimulated HRECs with a single dose of free PH at 10 \(\mu\)M did not induce any effect in the secretion of IL-6 or IL-8 over 48 hours (see Supplementary Fig. S4). This indicates that it was the continuous intracellular release of PH which lead to an improved effect with a 5-fold lower drug concentration. It has been previously shown that corticosteroids remain active in vitro for several days,\(^{67}\) can induce cytokine inhibition to stimulated cells within 2 hours of exposure, and retain the effect for up to 48 hours.\(^{64,65}\) This likely explains the sustained effect observed for free PH. Moreover, EPCR-NL2 exhibited significantly better performance compared to EPCR-NL1 and free drug, which is interesting considering the low concentration of PP (~2.4 \(\mu\)M) and its poor water solubility. PP would have to partition out of the liposomes, either through interactions with proteins and/or phospholipid bilayers in cells and vesicles, before readily reaching glucocorticoid receptors. We did not observe differences in IL-1\(\beta\) secretion levels for any of our culture conditions and treatments, even though it has been previously reported.\(^{50,59}\)

It has been shown that \(\mu\)M concentrations of triaminolone acetonide and cortisol as well as \(\mu\)M concentrations of dexamethasone can induce up to a 50\% inhibition in endothelial cell tube formation in vitro.\(^{68-70}\) We observed no inhibition in tube length or the number of segments for HRECs when treated with free PH at 25 \(\mu\)M (data not shown), but did observe inhibition of tube length and segment numbers at 50 \(\mu\)M (Fig. 6). EPCR-NL2 reduced tube length (20\% decrease) and significantly reduced the number of segments (50\% decrease) 8 hours after seeding HRECs on Matrigel. EPCR-NL1 did not perform as well and was comparable with free PH. No significant differences were observed amongst the EPCR-targeting liposomes and free PH. Based on the PH release profile of EPCR-NL, by the time of our measurements (i.e., 8 hours), less than 8\% of the drug had been released from the liposomes, which means that with less than 4 \(\mu\)M of PH, we induced a similar effect as 50 \(\mu\)M of free drug. The nontargeting liposomes that exhibited a similar release profile as the EPCR-targeting liposomes did not induce any effect in cell tube formation. We believe that this enhanced effect occurs because prednisolone is released intracellularly at higher concentrations for EPCR-NL than P-NL, maximizing the probability of drug-glucocorticoid receptor interactions. This results in a lower therapeutic concentration of prednisolone required compared to free prednisolone and a greater effect for EPCR-NL due to the improved endocytosis compared to P-NL. EPCR-NL liposomes showed a more profound effect on the endothelial cell tube formation assay despite the increased complexity of it compared to the two-dimensional cell monolayer assay.

In conclusion, we demonstrated that HRECs strongly express EPCR and that this receptor holds promise as a target for enhanced drug delivery to endothelial cells due to the significant increased uptake of EPCR-targeting liposomes in endothelial cells compared to nontargeting liposomes. EPCR-targeting liposomes showed significant reduction in IL-8 and IL-6 secretion from HRECs and HAECs, as well as suppressed HREC tube length and segment formation. Furthermore, EPCR-targeting liposomes loaded with prednisolone in both the bilayer and aqueous core showed greater anti-inflammatory potential than the equivalent liposomes loaded with prednisolone in the core only or free drug. Further experiments are required to determine if EPCR expression at the retina endothelium is significant in vivo, whether EPCR is evenly distributed or localized in the retina, and whether EPCR can be used as a nanomedicine target in diseased retinas.

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### References


