A Subretinal Cell Delivery Method via Suprachoroidal Access in Minipigs: Safety and Surgical Outcomes

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PURPOSE. This study evaluated a new subretinal method for delivery of human or pig umbilical tissue–derived cells (hUTC or pUTC, respectively) using a novel subretinal injection cannula and suprachoroidal approach in Göttingen minipig eyes. hUTC (palucorcel) are currently under development for treating geographic atrophy in humans.

METHODS. Twenty-four Göttingen minipigs (divided into eight groups) were subretinally administered palucorcel, pUTC, or vehicle. In some cases, fluorescently labeled cells and vehicle were administered. Conjunctival cutdown and sclerotomy were performed, then a flexible cannula containing a microneedle was inserted and advanced into the suprachoroidal space. The microneedle was deployed and visualized; 50 μL cells (target concentration, 1.2 × 10⁶ cells/mL [560,000 cells/eye]) or vehicle was injected subretinally. Safety outcomes were evaluated.

RESULTS. For all animals, cells and vehicle were successfully administered. Labeled cells or fluorescent vehicle were contained in the subretinal bleb, without leakage into the vitreous. No retinal detachment or vitreous traction band was identified by ophthalmologic examination. At all time points, observed microscopic changes were attributable to experimental procedures. On histopathology immediately after injection, localized retinal detachments were seen, along with focal retinal, choroidal, and/or scleral discontinuities. A moderate inflammatory response was seen in a limited number of animals. In the allogeneic setting, no antibody responses were detectable. Anti-human UTC antibodies were detected in the xenogeneic setting.

CONCLUSIONS. Palucorcel, pUTC, and vehicle were successfully administered to Göttingen minipigs using a novel subretinal injection cannula via a suprachoroidal surgical approach, with no significant adverse events; therefore, this technique appears to be feasible for further clinical development.

Keywords: subretinal cell delivery, suprachoroidal access, palucorcel, umbilical cells

AMD is a multifactorial, degenerative disease that is a leading cause of blindness in industrialized countries in people 55 years of age or older. The early and intermediate stages of the disease are characterized by the accumulation of drusen and a slow loss of the choriocapillaris. Initially, both stages are associated with minimal impairment of visual acuity. However, in advanced AMD, loss of vision can be caused by either exudative neovascular ("wet") AMD or geographic atrophy (GA). GA, which comprises 35% to 40% of all cases of advanced AMD, is a slowly progressive disease, characterized by loss of the RPE cells, retinal photoreceptors, and choriocapillaris, and is the cause of 20% of all cases of legal blindness in the United States. The appearance and growth of drusen deposits are prognostic indicators for GA, and the median time to develop central GA after any GA diagnosis is 2.5 years. The Age Related Eye Disease Study (AREDS) and Beaver Dam Eye Study demonstrated that the highest likelihood of developing GA is found in eyes with multiple large drusen (>125 μm) and in those with soft indistinct drusen, particularly when associated with macular pigmentary abnormalities. Despite the high prevalence and burden of dry AMD, there are no therapies currently available that are effective in delaying or preventing vision loss related to GA.

Palucorcel (CNTO 2476) is currently being evaluated as a novel cell-based therapy for the treatment of GA secondary to AMD. Palucorcel is composed of human umbilical cord tissue–derived cells (hUTC), and the nature and characterization of these cells has been reported previously. The cells are supplied in a proprietary cryopreserved formulation that is thawed shortly before use according to the manufacturer’s instructions. Subretinal administration of palucorcel was associated with preservation of visual function in a rat model of retinal degeneration. Subsequent in vitro studies showed that palucorcel was capable of rescuing RPE phagocytic dysfunction, promoting excitatory synaptic connectivity, enhancing neuronal survival, and enhancing neurite outgrowth via paracrine signaling pathways. A recent phase 1/2a study evaluated the administration of palucorcel in 35 adults, 50 years of age and older, with bilateral GA secondary to AMD. A microcatheter delivery system (iTrack 275; iScience, Inc., Menlo Park, CA, USA) was used in conjunction with an ab externo surgical approach to access the subretinal space. Palucorcel was well tolerated when delivered without retinal...
perforation. There was no clinical evidence of an immune response, cell rejection, or tumor formation.21,22 Although palucorcel was well tolerated in the phase 1/2a study, the ab externo surgical approach required to access the subretinal space with the microcatheter delivery system was associated with a high rate of retinal perforations (13/35 operated subjects) and retinal detachments (6/35 operated subjects).18 In another phase 1 study in subjects with retinitis pigmentosa, inadvertent seeding into the vitreous cavity, while injecting palucorcel into the subretinal space via a transvitreal approach, was associated with retinal detachment, as well as the formation of epiretinal membranes.23 The occurrence of retinal detachments and perforations in both early studies highlighted the need to develop a safer targeted surgical technique to deliver palucorcel into the subretinal space.

In the present study, we evaluated a novel method for the subretinal delivery of balanced salt solution (BSS+; Alcon Laboratories, Inc., Fort Worth, TX, USA), as well as both palucorcel and pUTC, using a novel subretinal injection cannula via the suprachoroidal space in minipigs. Features of this device include a flexible cannula with an advanceable microneedle for accessing the subretinal space (Fig. 1). This approach was designed to make use of the suprachoroidal space to deliver palucorcel at a specific site in the subretinal space with minimal trauma to ocular tissues while minimizing the egress of cells. The objectives of this study included the evaluation and identification of potential adverse effects associated with the use of the novel subretinal injection cannula; an assessment of the suprachoroidal surgical approach to deliver palucorcel subretinally; and the documentation of the cell injection procedure, device, and surgical performance in the preclinical setting. Additional objectives included an evaluation of the surgical healing time; characterization of postdosing disposition of administered fluorescent vehicle and carboxylfluorescein diacetate succinimidyl ester (CFSE)-labeled cells; and the identification of potential humoral immune responses as induced by the subretinal placement of either palucorcel or the pig analog, pUTC.

**METHODS**

This study was performed in accordance with the Organization for Economic Cooperation and Development Principles of Good Laboratory Practice and as accepted by local regulatory authorities. This study was conducted at the Charles River Laboratories Montreal ULC Senneville Site (Senneville, QC, Canada). The Test Facility Quality Assurance Program monitored the study to assure the facilities, equipment, personnel, methods, practices, records, and controls conformed to Good Laboratory Practice regulations. Animal use was in adherence with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research, and the protocol was reviewed by an animal care and use committee.

**Subretinal Injection Cannula**

The subretinal injection cannula used in this study has a number of unique design features to improve the safety of subretinal cell delivery via a suprachoroidal approach compared to previous surgical approaches or devices used to access the subretinal space. The flexible cannula conforms to the curvature of the eye and is intended to cannulate the suprachoroidal space, reducing the risk of choroidal hemorrhage when entering the suprachoroidal space. The advanceable microneedle housed within the cannula is intended to penetrate the choroid to provide access to the subretinal space without disruption or penetration of the retina. A knob on the handle allows for controlled advancement of the microneedle, and a third arm-positioning tool is used to stabilize the cannula during insertion into the suprachoroidal space. During the surgical procedure, a BSS+ entry bleb is created prior to cell delivery. The device fluid path allows for BSS+ and cell delivery without mixing, and the BSS+ entry bleb allows for the procedure to be aborted before cells are delivered if retinal penetration is observed.

**Test Articles**

Palucorcel was derived from umbilical cord tissue of a single donor as described previously.17,18 Palucorcel was developed in compliance with all relevant regulations, including donor consent. Briefly, after digestion with enzymes, a homogenous subpopulation of umbilical cord tissue cells was isolated, cultured, and used to derive a Master Cell Bank. These cells were then prepared as a frozen suspension in CryoStorSCO 4B/BSS (CS-SCO 4B/BSS, proprietary formulation, Janssen R&D, Spring House, PA, USA). Palucorcel was stored in liquid nitrogen vapor phase at temperatures below −120°C until ready for use.

The pUTC were isolated and expanded in a manner similar to palucorcel from pigs that were recognized by the United States Department of Agriculture as free of brucellosis, pseudorabies, transmissible gastroenteritis, and porcine reproductive and respiratory syndrome. Piglets used for deriving pUTC were vaccinated against *Mycoplasma hyopneumoniae*, *Pasteurella multocida*, *Bordetella bronchiseptica*, and *Erysipelothrix rhusiopathiae*. Adult pigs were vaccinated against *E. rhusiopathiae*, *Leptospira*, and parvovirus. Cells were formulated in 90% (vol/vol) fetal bovine serum (FBS; GE Healthcare, Logan, UT, USA) and 10% (vol/vol) dimethylsulfoxide (Sigma-Aldrich Corp., St. Louis, MO, USA) and cryopreserved in liquid nitrogen vapor phase at temperatures below −120°C.

Palucorcel and pUTC dose formulations were prepared using aseptic technique. Vials of cryopreserved cells were thawed in a water bath at 37°C. Shortly after, palucorcel was administered directly to the subretinal space. For pUTC, cells...
were washed twice in BSS and resuspended in CS-SCO 4B/BSS. Cell counts were then obtained on transferred aliquots of palucorcel and pUTC to ensure that the viable cell concentrations were ≥70% in order to meet dosage level requirements for administration.

Some vials of palucorcel and pUTC were labeled with cCFSE using a kit ( Vybrant CFDA SE Cell Tracker Kit; Molecular Probes, Inc., Eugene, OR, USA). Briefly, cells were thawed at 37°C in a water bath for 2 minutes. The pUTC were washed twice in BSS and resuspended in CS-SCO 4B/BSS. Palucorcel was used directly after thaw. Each cell type (330,000 cells/mL) was combined with 670 μL 37°C 1X Dulbecco’s phosphate buffered saline (DPBS; Life Technologies, Carlsbad, CA, USA). Cells were subsequently incubated with 0.5 μL 5 mM CFSE for 15 minutes in the dark. The reaction was quenched by adding 1 mL 37°C FBS. Cells were washed twice with 1X DPBS using centrifugation. Cellular pellets were resuspended in 10% FBS prepared in 1X DPBS such that the target concentration was 11.2 × 10^6 cells/mL (560,000 cells/eye).

**Animals and Treatment Administration**

Male Göttingen minipigs were obtained from Marshall Bio-Resources (North Rose, NY, USA). Animals were between 9 and 10.5 months old and weighed 12.9 to 17.6 kg at the start of the study. Animals were assigned to treatment groups by a stratified randomization scheme designed to achieve similar group body weights. Animals were single-housed in stainless steel cages with an automatic watering valve and had access to a standard certified commercial laboratory diet (Harlan Teklad Miniswine Diet 8753; Harlan Laboratories, Inc., Madison, WI, USA) and the position and size of the position of retaining sutures (6-0 Vicryl sutures; Ethicon, Inc., Somerville, NJ, USA) were used to facilitate comparison of potential effects in the xenogeneic and allogeneic setting.

All surgeries were performed by a single vitreoretinal surgeon. The conjunctiva was opened, and the underlying sclera was freed from any overlying connective tissue. A marker was used to indicate the position of retaining sutures (6-0 Vicryl sutures; Ethicon, Inc., Somerville, NJ, USA) and the position and size of a sclerotomy, parallel but located 8 to 9 mm posteriorly to the limbus. The sclerotomy was created using a crescent knife, and the remaining scleral fibers were removed with a Sinsky hook. The diameter was adjusted to allow for passage of the cannula while providing a snug fit. The subretinal injection cannula was passed under the retaining sutures and advanced tangential to the eye through the sclerotomy, approximately 3 to 5 mm into the suprachoroidal space. Once in position, the cannula was sutured to the surface of the sclera to prevent unwanted movement of the cannula. Using indirect ophthalmoscopy to visualize the cannula tip, the microneedle was carefully advanced into the subretinal space. A small BSS+ entry bleb was created to confirm that the microneedle tip was in the subretinal space. This was followed by the administration of cells (560,000 cells/eye) or vehicle in a 50-μL volume forming a suitable pupil dilation. Animals received intravenous lactated Ringer’s solution (10 mL/kg/h) during the dosing procedure to aid in recovery from anesthesia.

On day 1, a single subretinal injection of vehicle (CS-SCO 4B/BSS; groups 3 and 6), palucorcel (target concentration, 11.2 × 10^6 cells/mL [targeted dose, 5.6 × 10^5 cells] in CS-SCO 4B/BSS; groups 1, 4, and 7), or pUTC (target concentration 11.2 × 10^6 cells/mL [targeted dose, 5.6 × 10^5 cells]) in CS-SCO 4B/BSS; groups 2, 5, and 8) was administered in a volume of 50 μL in the right (treated) eye (Table 1). These injections were preceded by injection of a small volume of BSS+ for bleb formation. For groups 1 and 2, palucorcel and pUTC, respectively, were fluorescently labeled with CFSE before administration. Animals in group 2 received vehicle CS-SCO 4B/BSS containing fluorescein (FA) and indocyanine green (ICG) in the left eye. For all other groups, the left eye was untreated and did not undergo surgical procedures. The dose of palucorcel used in this study was based on the highest dose used in prior clinical studies. An equivalent dose of pUTC was used to facilitate comparison of potential effects in the xenogeneic and allogeneic setting.

Animals in groups 1 and 2 were terminated on the day of surgery; those in groups 3 to 5 were terminated at month 1 (day 30); and those in groups 6 to 8 were terminated at month 5 (day 92; Table 1).

**Suprachoroidal Surgical Approach and Subretinal Injection Cannula**

All surgeries were performed by a single vitreoretinal surgeon. The conjunctiva was opened, and the underlying sclera was freed from any overlying connective tissue. A marker was used to indicate the position of retaining sutures (6-0 Vicryl sutures; Ethicon, Inc., Somerville, NJ, USA) and the position and size of a sclerotomy, parallel but located 8 to 9 mm posteriorly to the limbus. The sclerotomy was created using a crescent knife, and the remaining scleral fibers were removed with a Sinsky hook. The diameter was adjusted to allow for passage of the cannula while providing a snug fit. The subretinal injection cannula was passed under the retaining sutures and advanced tangential to the eye through the sclerotomy, approximately 3 to 5 mm into the suprachoroidal space. Once in position, the cannula was sutured to the surface of the sclera to prevent unwanted movement of the cannula. Using indirect ophthalmoscopy to visualize the cannula tip, the microneedle was carefully advanced into the subretinal space. A small BSS+ entry bleb was created to confirm that the microneedle tip was in the subretinal space. This was followed by the administration of cells (560,000 cells/eye) or vehicle in a 50-μL volume forming a

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**Table 1. Dosing Groups**

<table>
<thead>
<tr>
<th>Group</th>
<th>Left Eye</th>
<th>Right Eye</th>
<th>Scheduled Euthanasia</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Untreated control</td>
<td>11.2 × 10^6 palucorcel labeled with CFSE† Day 1</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Vehicle‡ labeled with FA (1 mg/mL) and ICG (0.05 mg/mL)</td>
<td>11.2 × 10^6 pUTC labeled with CFSE† Day 1</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Untreated control</td>
<td>Vehicle‡ Month 1 (day 30)</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Untreated control</td>
<td>11.2 × 10^6 palucorcel† Month 1 (day 30)</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Untreated control</td>
<td>11.2 × 10^6 pUTC† Month 1 (day 30)</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Untreated control</td>
<td>Vehicle‡ Month 3 (day 92)</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Untreated control</td>
<td>11.2 × 10^6 palucorcel† Month 3 (day 92)</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Untreated control</td>
<td>11.2 × 10^6 pUTC† Month 3 (day 92)</td>
<td></td>
</tr>
</tbody>
</table>

† Target dose, 5.6 × 10^5 cells/eye.
‡ CS-SCO 4B/BSS.

ICG, indocyanine green.

* All doses were administered as a single dose of 50 μL per eye.

‡ Target dose, 5.6 × 10^5 cells/eye.
‡ CS-SCO 4B/BSS.
small bleb around the microneedle tip (Fig. 2). The microneedle was then retracted, and the cannula was removed. The sclerotomy and conjunctiva were closed using Vicryl sutures.

**Study Assessments**

Minipigs first underwent routine ophthalmic examinations using a handheld slit lamp. Eyes were then dilated with 1% tropicamide and examined using an indirect ophthalmoscope. All ophthalmic examinations were performed by a trained ophthalmic veterinarian. FA and ICG angiography using a scanning laser ophthalmoscope (Spectralis; Heidelberg Engineering, Heidelberg, Germany) and electroretinography (ERG) (LKC UTAS-E-4000 with EMWin software) were performed at predetermined time points, as described below. In eyes that had received CFSE-labeled cells or vehicle with a fluorescent dye, fluorescence was measured using the blue filter on the Spectralis imaging system on the day of injection. Ophthalmic examinations were performed once before treatment; on days 3, 7, and 15; approximately 1 month after surgery; and monthly thereafter. Fundus imaging was performed once before and immediately following the surgical procedure on day 1 for animals who received fluorescently labeled cells or vehicle with fluorescent dye. FA and ICG angiography were performed once before treatment, at 1 month after dosing, and before necropsy. No postdose angiography was performed on animals euthanized on day 1. ERG was performed once before treatment and before necropsy for all animals except those euthanized on day 1. Following necropsy, histopathology was performed on each eye by a board-certified veterinary pathologist.

Intraoperative assessments included the device’s ability to access the suprachoroidal space, the ease of insertion and advancement into this space, the microneedle’s ability and ease of accessing the subretinal space, and the microneedle’s ability to deliver the product. These outcomes were recorded during or directly following the procedure using a formalized questionnaire. Any complications, such as a hemorrhage in any ocular compartment, retinal perforation and/or delivery of vehicle or cells to the vitreous, leakage of injected formulation after microneedle withdrawal, or collapse of the bleb following the removal of the cannula, were also recorded. Routine monitoring and safety assessments, including body weights, food consumption, and clinical pathology parameters (hematology, coagulation, clinical chemistry) were evaluated based on clinical observations and blood samples. Serum was collected once before treatment (on all animals); during weeks 1 and 2; and at months 1, 2, and 3 (depending on scheduled euthanasia) and was tested for the presence of IgG antibodies against palucorcel or IgG antibodies against pUTC using flow cytometry.

Detection of Anti-hUTC or Anti-pUTC IgG Antibodies in Serum

Serum samples were analyzed for the detection of anti-hUTC or anti-pUTC IgG antibodies using a flow cytometry method. Briefly, hUTC or pUTC were cultured to confluence and subsequently activated by addition of 25 ng/ml human IFN-γ (Invitrogen, Carlsbad, CA, USA) or 80 ng/ml porcine IFN-γ (R&D Systems, Inc., Minneapolis, MN, USA), respectively, for 48 hours. Cells were harvested and then treated with normal minipig serum (NMS), undiluted test samples, or undiluted positive controls (porcine serum obtained from pigs immunized with pUTC or hUTC). Fluorochrome-labeled antibodies against pig IgG antibodies were used as a secondary reagent to allow detection of pig anti-pUTC or pig anti-hUTC IgG antibodies by flow cytometry. Negative controls (secondary antibody only) were used to set the flow cytometer (FACSCalibur; Beckton-Dickinson, Franklin Lakes, NJ, USA) voltages by setting the peak geometric mean fluorescence (GMF) between 100 and 101 (log histogram), while the NMS control represented background signal. A sample was considered positive if that sample’s GMFSample/GMFNegative ratio was above the set cut-point. The pre- and all postdose samples from each animal were analyzed in the same experiment on IFN-γ-activated hUTC and pUTC cells. The cut-point was established to be 1.013 for anti-hUTC IgG and 0.960 for anti-pUTC IgG. Postdose samples that were found to be positive based on the cut-point were further evaluated by comparing their GMF against the GMF of their respective predose sample (GMFPredose/GMFPostdose) to see if an increase in anti-hUTC or anti-
pUTC IgG was observed during the dosing phase when compared to predose levels. A postdose sample that showed an increase greater than 30% (GMFPostdose /GMFPredose ratio > 1.3) compared to the corresponding predose sample was considered as potentially containing anti-hUTC or anti-pUTC IgG antibodies.

**Terminal Procedures**

Animals were subjected to a complete necropsy examination. Eyes and optic nerves were collected and preserved in Davidson’s fixative. The eyelids, extraocular muscle, and nictitating membrane (bilateral) were preserved in 10% neutral buffered formalin. Tissues were embedded in paraffin, sectioned, mounted on glass slides, and stained with hematoxylin and eosin. For each eye, at least two slides were prepared for sagittal sections from each of the following locations: the sclerotomy incision site, the catheter tract, the bleb areas (if visible), and the optic nerves. Histopathological evaluation was performed by a board-certified veterinary pathologist at Charles River Labs in Montreal. Select images were captured at the discretion of the study pathologist with the Aperio Digital Image Capture System (Leica Biosystems, Buffalo Grove, IL, USA).

**RESULTS**

**Palucorcel Exposure and Fundus Imaging**

Palucorcel, pUTC, and vehicle were successfully administered using the subretinal injection cannula and suprachoroidal surgical procedure in all animals. The targeted cell concentration of palucorcel and pUTC across all dose groups was $11.2 \times 10^6$ cells/mL (560,000 cells/eye). Actual concentrations of palucorcel delivered ranged from $7.93 \times 10^6$ to $11.98 \times 10^6$ cells/mL. Actual concentrations of pUTC delivered ranged from $6.88 \times 10^6$ to $14.68 \times 10^6$ cells/mL.

Immediately post dose, CFSE-labeled palucorcel was clearly visible as multifocal pinpoints of hyperfluorescence located in the bleb in autofluorescence fundus images (example shown in Fig. 3A). The fluorescent vehicle was clearly visible as a discrete area of hyperfluorescence in the bleb area during fundus imaging (example shown in Fig. 3B). Based on fundus imaging, no leakage of cells or vehicle into the vitreous was observed.

**Surgical Observations**

In all operated eyes, the procedure was performed as planned without any device failure. Cannula insertion into the suprachoroidal space was rated as very easy or easy for the majority of operated eyes (23/27). Possible retinal perforations were reported in four of 27 operated eyes, and partial bleb collapse was noted in six of 27 operated eyes. A subretinal hemorrhage was noted in one of the 27 operated eyes, but in no case did a choroidal or subretinal hemorrhage prevent successful delivery (Table 2).

**Clinical Observations**

As expected, there were no unscheduled deaths during the study. Following surgery, there were no clinical signs (e.g., changes in body weight, food consumption, clinical pathology parameters) that were considered related to palucorcel or pUTC. The observed clinical signs were related to the experimental procedures. Ocular clinical signs included ocular discharge, conjunctival hyperemia, and peri orbital swelling, as well as peri orbital skin staining and scabbing. These clinical signs generally resolved within the first month, although conjunctival hyperemia persisted through day 52 in animals euthanized at month 3. Hyperemia was not associated with any signs of intraocular inflammation.

**Ophthalmic Observations**

Ophthalmic observations for eyes administered with CS-SCO vehicle, palucorcel, or pUTC are summarized in Supplementary Table S1. Diffuse hyperemia, hemorrhage, and swelling resulting from the surgery improved significantly over the first few weeks. Minor corneal findings (focal or diffuse opacities
and superficial vascularization) related to the experimental procedures improved or resolved during the initial postoperative month, although slight focal corneal opacities remained present in two eyes at 3 months post dose. Minor retinal changes (focal grayish retinal haze, irregular retina/choroid pigmentation, focal/multifocal hemorrhages, and focal retina/choroid opacities) related to the experimental procedures improved or resolved during the first postoperative month. Slight to moderate focal retinal elevations caused by bleb formation during the dosing procedure were observed at day 3 in all vehicle-treated eyes and resolved within 1 week.

Similar to vehicle, subretinal administration of palucorcel or pUTC caused minor observable changes. There was no sign of ocular inflammation. At the site of injection, the following minor alterations were observed: slight to moderate grayish cell-like retinal opacities suggestive of cellular aggregates that were present in the subretinal space of all palucorcel-treated eyes and most pUTC-treated eyes. These grayish opacities, which were not present in vehicle-treated eyes, resolved within 1 to 2 months after dosing. Slight to moderate focal retinal elevations caused by bleb formation were observed on day 3 in eyes receiving palucorcel or pUTC. These had resolved by 1 month. Similar focal retinal elevations were observed in vehicle-treated eyes, but resolved more rapidly. At the cannula entry site into the suprachoroidal space, a small patch of bare sclera was present that was devoid of choroidal vasculature. This was not associated with any retinal detachments or vitreous traction bands (by ophthalmic observation) with vehicle, palucorcel, or pUTC administration. No abnormalities were noted in the neighboring tissues.

**Angiography**

In four animals that received palucorcel, focal changes were observed on FA and ICG angiography in the area of subretinal delivery. No dye leakage was noted from either the retinal or choroidal vasculature. At the 1-month angiogram, one animal had focal areas of FA hyperfluorescence within the area corresponding to the surgically induced subretinal bleb. Two focal areas of hypofluorescence were visible on ICG angiograms in the same vicinity for that animal. On 3-month angiograms, focal areas of FA hyperfluorescence and/or hyperfluorescence were present in the superior and superotemporal region of the fundus of two animals (both administered with palucorcel) in the area of the subretinal bleb. Focal hypofluorescent areas were present in the same locations on ICG angiograms and may indicate the subretinal injection site. In addition, tortuous small vessels were present at one hypofluorescent site for one of the two animals with focal hypofluorescence and/or hyperfluorescence on FA angiograms at month 3. Similar findings were observed in one animal administered with pUTC at months 1 and 3.

**Electroretinography**

No changes in scotopic or photopic ERG amplitudes and implicit times were observed in eyes that were administered with palucorcel or pUTC as compared with eyes treated with CS-SCO vehicle. At 1 month and 3 months post dose, minor increases or decreases in average amplitudes and implicit times were often observed in all treated eyes compared to their baseline measurements and were considered to be related to individual variability and/or anesthesia levels (Supplementary Table S2).

**Histopathology**

Histopathologic observations in animals euthanized on day 1, month 1 (day 30), and month 3 (day 92) are summarized in Table 3. At all time points, some microscopic changes attributable to the experimental procedures were observed. On histopathologic examination, animals that were euthanized immediately after surgery characteristically had a peripheral retinal detachment at the site of injection; focal discontinuity of the retina, choroid, and/or sclera; infiltration of the limbus by a mixed population of inflammatory cells; and subretinal, subchoroidal, and/or limbus hemorrhage. The retinal detachment observed immediately post dose was limited to a focal area in the mid to posterior retina and corresponded to the injection site and the retinal elevations observed in ophthalmic observation. In some cases, test item cells were noted in the subretinal space (Fig. 4). Changes in the eyes of all groups of animals euthanized 1 month post dose included inflammatory cell infiltration (choroid, sclera, and/or limbus); fibrosis of the
sclera; pigmented macrophages in the vitreous chamber, choroid, and retina; and disorganization of the retina (mainly the outer nuclear and photoreceptor layers). By 3 months post dose, microscopic findings included inflammatory cell infiltration (retina, choroid, and/or sclera), choroid fibrosis or discontinuity, and disorganization (Fig. 5). Retinal degeneration or atrophy, following focal discontinuity of the retina, remained present. In two of three animals that received palucorcel (xenogeneic) and were euthanized at month 3 (day 92), small focal retinal granulomas were observed (Fig. 5A). No granulomas were observed in eyes that received pUTC (allogeneic) or vehicle (Fig. 5B).

**TABLE 3. Summary of Microscopic Eye Findings**

<table>
<thead>
<tr>
<th>Microscopic Eye Findings</th>
<th>Euthanized Day 1</th>
<th>Euthanized Month 1</th>
<th>Euthanized Month 3</th>
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<tbody>
<tr>
<td></td>
<td>Palucorcel, n = 3</td>
<td>pUTC, n = 3</td>
<td>Vehicle, n = 3</td>
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<tr>
<td>Hemorrhage</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Retina detachment</td>
<td>3</td>
<td>2</td>
<td>2</td>
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<tr>
<td>Choroid discontinuity</td>
<td>2</td>
<td></td>
<td>2</td>
</tr>
<tr>
<td>Retina discontinuity</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sclera discontinuity</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed cell limbus infiltration</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed cell sclera infiltration</td>
<td>2</td>
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<tr>
<td>Mononuclear cell choroid infiltration</td>
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<td>Mononuclear cell retina infiltration</td>
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<td>Choroid fibrosis</td>
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<td>Macrophage aggregation</td>
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<td>Retina degeneration</td>
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<td>Retina atrophy</td>
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<td>Retina granuloma</td>
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**Serum Antibody Titers**

Subretinal administration of palucorcel in CS-SCO 4B/BSS to minipigs (xenogeneic setting) resulted in detectable levels of anti-hUTC IgG antibodies in two of six animals (Supplementary Table S3). One animal (necropsied on day 50) showed elevated anti-hUTC IgG levels at week 2 post dosing. The antibody response continued to persist at month 1. Another animal (necropsied on day 92) showed no change at month 1 and elevated anti-hUTC IgG levels at 2 months post dosing that dropped off at month 3 post dosing. In contrast, subretinal administration of pUTC to minipigs (allogeneic) resulted in no detectable levels of anti-pUTC antibodies (Supplementary Table S4).

**DISCUSSION**

This study evaluated the safety, tolerability, and clinical response of a single, subretinal injection of palucorcel or pUTC through the suprachoroidal space, using a novel injection cannula. The technology and techniques used in this novel subretinal delivery procedure were developed with expert retinal surgeons who helped to create a procedure that builds off previously known skills and devices. Surgical steps,
such as conjunctival peritomy, eye rotation, choroidal exposure via a sclerotomy, and needle puncture of the choroid are all standard techniques used for decades in retinal detachment drainage procedures. With regard to suprachoroidal cannulation, Olsen and colleagues described a microcatheter for (1) insertion into the suprachoroidal space, (2) advancement to the macular region, and (3) delivery of a therapeutic product to the suprachoroidal space. El Rayes and Elbogy used suprachoroidal cannulation to create an internal buckle for retinal detachments by injecting astaining viscoelastic through an olive tip cannula. A number of subretinal injection procedures, particularly in the macular area, have made use of such cannulas, with the cells themselves, would be rapidly cleared out of the eye through emissary channels or would be hindered from reaching the retina by multiple barriers within the choroid, Bruch’s membrane, and RPE. For these reasons, the subretinal space was defined as the target for delivery; and the technique described here was developed to ensure the drug product is physically deposited adjacent to the target retinal cells.

In order to achieve our goal, a novel suprachoroidal positioning cannula coupled to a subretinal extension needle was required. The current procedure and device is novel in its ability to combine the positioning of a catheter in the suprachoroidal space with the safe delivery of a biologically active product to the subretinal space. In the process of developing the cannula/needle combination, several parameters were considered. These included the following: allowing the surgeon to safely position the catheter within the suprachoroidal space at a predetermined position (a rectangular design); the safe advancement of a microneedle to pierce through Bruch’s membrane without damaging or perforating the retina (extension from the distal end of the catheter at an appropriate angle); and removing the effect of hand tremor and the need to proceed quickly with the injection.

Dose formulations were successfully administered into the subretinal space of Göttingen minipigs in 27 out of 27 eyes. CFSE fluorescently labeled cells or fluorescent vehicle were immediately visible post dose within the subretinal blebs, and no leakage of cells or vehicle into the vitreous was observed. Intraoperative observations generally indicated that the procedure was easy to perform and was associated with a relatively low rate of retinal perforations in comparison to the previous ab externo microcatheter-based system. The delivery blebs that were formed were closely observed. Most of the blebs remained fully formed when observed immediately after the procedure, with partial collapse observed in only 6 of 27 blebs. These collapses were typically present when the removal of the catheter was more complicated, for example, when the catheter was torqued during removal. These findings indicate that a smooth retraction along the same entry path is needed to prevent bleb collapse. The ophthalmic observation of retinal elevation in the area of the subretinal injection was observed and tracked over time, generally decreasing in severity from moderate at day 3 in some animals to slight or very slight by day 15, with resolution in all animals by 1 month (Supplementary Table S1). These intraoperative and ophthalmic exam findings provide supporting evidence for successful subretinal cell delivery followed by cell media fluid transport out of the subretinal space over time. This result on cell media fluid
transport is in agreement with previously described kinetics of macromolecules injected into the subretinal space.37

Histology performed immediately after dosing showed the presence of the expected retinal detachment. This histopathologic observation of retinal detachment corresponded to the ophthalmologic observation of retinal elevations. In some eyes, cells could be observed in the subretinal space but not in the choroid or suprachoroidal space. At 3 months, in two of three eyes that received palucorcel (xenogeneic cells), histology revealed a focal granuloma. No such cell-mediated immune response was observed in animals administered with pUTC. A humoral immune response was also observed in animals receiving palucorcel. Detectable anti-hUTC IgG antibody levels were observed in two of six animals. Anti-hUTC was observed in one animal after 2 weeks, which persisted at 1 month when this animal was necropsied. Anti-hUTC was observed in a second animal at 2 months post dose. This animal was one of two animals observed with a focal granuloma. There were no detectable antibodies observed in animals receiving pUTC. For this study, in which human-derived cells were injected into minipigs, the observed mild immune response suggests a low likelihood for palucorcel to induce an immune reaction and is consistent with the low immune response to UTC observed in previous studies.18,21

This study provides data on the potential adverse events and other safety sequelae associated with this procedure for subretinal cell delivery via a suprachoroidal access. Data on procedure-related adverse events is generally lacking for other types of cells being evaluated as potential therapies for AMD. Preclinical studies of these products generally focus on the safety and tolerability of the cells, rather than the procedure.38,39 Furthermore, preclinical studies of subretinal cell delivery have reported the occurrence of retinal perforations and hemorrhages, though not generally the rates of these complications.39,41 These findings highlight the need for a procedure and device with improved safety outcomes for subretinal delivery of therapeutic products.

Taken together, these results indicate that utilization of a suprachoroidal approach represents an improvement over previous surgical approaches, including ab externo and transvitreal routes of administration, for which rates of retinal detachments and retinal perforations were high and dependent on surgical expertise.18,23 This approach appears to be less traumatic, having mitigated the risk of retinal detachment, retinal traction, and formation of vitreal membrane-like white opacity observed using previous methods of administration in the minipig eye, and may be more widely adaptable for a broad surgical audience.

The device and its procedure were the result of a large multidisciplinary team, with close interactions throughout the development process. Future efforts are focused on addressing the challenges of transferring this new technology to a wide range of surgeons, and studies assessing the safety and tolerability of this approach in patients with AMD are warranted.

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