Patient-derived endothelial progenitor cells improve vascular graft patency in a rodent model

J.D. Stroncek,1, L.C. Ren,1, B. Klitzman,2, W.M. Reichert1, *

1 Department of Biomedical Engineering, Duke University, Durham, NC 27708, USA
2 Kenan Plastic Surgery Research Labs, Duke University, Durham, NC 27708, USA

**Abstract**

Late outgrowth endothelial progenitor cells (EPCs) derived from the peripheral blood of patients with significant coronary artery disease were sodded into the lumens of small diameter expanded polytetrafluoroethylene (ePTFE) vascular grafts. Grafts (1 mm inner diameter) were denucleated and sodded either with native EPCs or with EPCs transfected with an adenoviral vector containing the gene for human thrombomodulin (EPC + AdTM). EPC + AdTM was shown to increase the in vitro rate of graft activated protein C (APC) production 4-fold over grafts sodded with untransfected EPCs (p < 0.05). Unsodded control and EPC-sodded and EPC + AdTM-sodded grafts were implanted bilaterally into the femoral arteries of athymic rats for 7 or 28 days. Unsodded control grafts, both with and without denucleation treatment, each exhibited 7 day patency rates of 25%. Unsodded grafts showed extensive thrombosis and were not tested for patency over 28 days. In contrast, grafts sodded with untransfected EPCs or EPC + AdTM both had 7 day patency rates of 88–89% and 28 day patency rates of 75–88%. Intimal hyperplasia was observed near both the proximal and distal anastomoses in all sodded graft conditions but did not appear to be the primary occlusive failure event. This in vivo study suggests autologous EPCs derived from the peripheral blood of patients with coronary artery disease may improve the performance of synthetic vascular grafts, although no differences were observed between untransfected EPCs and TM transfected EPCs.

**Keywords:** Synthetic vascular graft, ePTFE, Endothelial progenitor cells, Thrombomodulin, Coronary artery disease

1. Introduction

In the US, about 500,000 patients undergo coronary artery bypass grafting (CABG) each year [1]. It is estimated that 15% of CABG patients are unable to undergo repeat surgeries because they lack sufficient blood vessels suitable for grafting [2,3]. This problem is compounded by the lack of clinically suitable small diameter (<6 mm) synthetic vascular grafts, primarily due to unacceptable patency rates caused by luminal thrombosis and intimal hyperplasia.

The presence of a functional and intact endothelium has been shown to improve the patency of synthetic small diameter grafts [4,5]. However, endothelialized synthetic grafts have not received widespread adoption, in part due to difficulties with the invasive autologous sourcing of endothelial cells (ECs) [6]. EC-like cells with high proliferation potential can be isolated non-invasively from peripheral blood. Here, we have adopted the term “late outgrowth endothelial progenitor cells” (EPCs), while others have used “endothelial colony forming cells” [7,8].

It has been shown previously that blood-derived EPCs isolated from patients with significant coronary artery disease (CAD) exhibit many hallmarks of healthy endothelial cells. They can be expanded to higher densities with minimal contamination by other cell types, can maintain firm adhesion to the underlying substrate, and can reduce clot formation by releasing anti-thrombotic factors [9–11]. While short-term proof-of-principle studies have indicated that animal-derived EPCs can improve the patency rates of synthetic small diameter vascular grafts [12–14], there has been limited longer-term in vivo testing of patient-derived EPCs.

A key intrinsic mediator of vascular thromboresistance is thrombomodulin (TM), a membrane-bound glycoprotein that is central to the protein C anti-coagulation pathway [15]. TM binds free thrombin to change its conformation and inhibit its pro-coagulant potential. The TM–thrombin complex then acts as a cofactor that binds with protein C, catalyzing its conversion to activated protein C (APC). APC in turn inhibits clotting by binding protein S and proteolytically cleaving two coagulation factors responsible for generating thrombin: Factor Va and Factor VIIIa [15,16].

TM expression by ECs is decreased in patients with diabetes and atherosclerosis, and these patients might benefit from up-regulating their TM expression [17–19]. Autologous vein grafts with ECs overexpressing TM have previously been shown to reduce thrombosis [20,21] and decrease intimal hyperplasia [21]; however, the ability of TM overexpression to improve the performance of small diameter synthetic vascular grafts remains unknown. The purpose

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* Corresponding author. Tel.: +1 919 660 5151; fax: +1 919 660 5362.
E-mail address: reichert@duke.edu (W.M. Reichert).

1 Co-first authors.
of this study was to test whether EPCs isolated from patients with CAD could reduce thrombus formation of small diameter ePTFE grafts, and whether overexpressing TM by EPCs would further increase the thromboreistance.

2. Materials and methods

2.1. EPC isolation and cell culture

All cells designated as “EPCs” in the current study were late outgrowth EPCs isolated and expanded from 50 ml of peripheral blood drawn from patients undergoing cardiac catheterization in the Duke University Medical Center who had advanced coronary artery disease diagnosed by angiography. The Duke University Institutional Review Board approved the protocol for collection and use of human blood employed in the study.

EPCs were isolated following a protocol for isolation from cord blood [22]. Briefly, mononuclear cells were separated and plated onto collagen I coated plates and cultured in complete endothelial basal medium (EBM-2) plus endothelial growth medium (EGM-2) SingleQuots (Lonza, Walkersville, MD) with 10% fetal bovine serum (Thermo Scientific HyClone, Waltham, MA) and 1% antibiotic/antimycotic solution (Gibco, Carsbad, CA). After 24 h of culture, nonadherent cells were removed and complete EGM-2 medium was added to each well. Medium was changed daily for 7 days and then every other day. EPCs were characterized previously as “EC-like” if they were positive for the EC markers CD31 and CD105 and negative for CD133, CD14 and CD45 [11]. EPCs were isolated from five patients and used at passages 5–10 (three males and two females with an average age of 56 years, range 44–69). Patient demographics have been reported previously [11].

2.2. Adenovirus vectors

Two different adenoviral vectors were used in this study and have been described previously [19]. Replication deficient β-galactosidase adenovirus “control vector” (AdCV) was purchased (Eton Bioscience, San Diego, CA). Replication deficient adenoviral expressing human thrombomodulin (AdTM) was a gift from Dawn Bowles, Duke University. EPCs were transfected with AdTM or AdCV for 4 h at 100 MOI (multiplicity of infection) 1 day before graft sodding.

2.3. Vascular graft conditions

Small diameter ePTFE vascular grafts with 1 mm inner diameter and 30 μm nominal internodal distance (International Polymer Engineering, Tempe, AZ) were used in all studies. Five vascular graft conditions were tested: (i) bare off-the-shelf ePTFE vascular grafts; (ii) denuded grafts (grafts having air removed by immersing in 100% ethanol and drawing a vacuum until no bubbles were released from the graft); (iii) grafts sodded with native EPCs; (iv) grafts sodded with EPCs transfected with AdCV; and (v) grafts sodded with EPCs transfected with AdTM. Grafts were denuded in all conditions that involved cell sodding.

2.4. Graft sodding

The graft endothelialization method used in this study was similar to that of Kidd et al. [23], where cells are “sodded” onto the porous graft lumen under positive pressure. ePTFE vascular grafts were placed in vacutainers containing 2.5 mg ml−1 tridodecylmethylammonium chloride (TDMAC; Polysciences, Warrington, PA), a cationic surfactant, in ethanol at room temperature. A vacuum was drawn to “denude” grafts by removing residual air.

Denudation was halted when air bubbles were no longer released from the graft and the graft became translucent and sank to the bottom of the solution. Denudation reduces material thrombogenicity [24] and TDMAC increases fibronectin adsorption on hydrophobic materials such as ePTFE [25]. The grafts were washed three times in Dulbecco’s phosphate-buffered saline and incubated in a solution of human fibronectin (20 μg ml−1) at 37°C for approximately 1 h.

Transfected and native EPCs were detached using 0.025% trypsin (Lonza) for approximately 10 min and resuspended in 5 ml medium to achieve an effective graft luminal seeding density of 1.5 × 10⁶ cells cm⁻². The graft was cannulated with a 19-gauge needle and the distal end of the graft was clamped with titanium vascular clips (Teleflex medical, Research Triangle Park, NC). The cell suspension was infused into the graft at a rate of 0.5 ml min⁻¹ using a programmable syringe pump (Harvard Apparatus, Holliston, MA). The cell solution was flushed through the graft and cells were trapped in the graft interstices. The effluent was collected into a separate container and showed few cells, indicating that the majority of the cells were present on the graft. Following sodding, the cannula and vascular clamps were removed and the graft was placed into a 6-well plate containing EBM-2 complete medium and cultured for 2 days before in vitro or in vivo testing.

2.5. EPC adhesion to the graft lumen

Cell coverage on the graft was assessed after exposure to surgical clamping and super-physiological flow. Fresh EBM-2 complete medium was used to cover the graft while in the 6-well plate. Microvascular clamps (S&T AG, Neuhauen, Switzerland) were placed at each end of the graft. Every 5 min, the position of the clamps were adjusted along the length of the graft to simulate the suturing movements. This procedure was performed for 30 min and the 6-well plate was maintained at 37°C using a heated stage (Pathology Devices, Westminster, MD). After 30 min, the graft was either fixed with 3.7% paraformaldehyde (Clamped + Static) or exposed for 5 min to a laminar flow at 100 dynes cm⁻² with a peristaltic flow setup as described above and fixed (Clamped + Flow). En face scanning electron microscopy (SEM) images (1000×) were taken at four randomly selected visual fields of the graft luminal surface. Cell coverage was determined by overlaying a transparent sheet with a 10 × 13 uniform grid onto the micrograph and counting the percentage of grid intersection points overlying cells (n = 4).

2.6. APC production

The biological activity of EPCs sodded onto vascular grafts was assessed by measuring the rate at which protein C was cleaved by the TM–thrombin complex to form APC. Grafts (length 7 mm) were sodded with EPCs as described above, placed into wells of a 24-well plate and incubated with 5 mM thrombin (Haematologic Technologies Inc (HTI), Essex Junction, VT) and 400 mM human protein C (HTI). The rate of APC production over 1 h was assayed in a microtiter plate reader (BioTek Instruments, Winooski, VT) after the addition of Spectrozyme PCa (American Diagnostica, Stamford, CT), as described previously [19] (n = 4).

2.7. Implantation

The experimental protocol was approved by the Duke University Institutional Animal Care and Use Committee. Thirty-three male athymic rats, weighing 180–280 g (Charles River Laboratories, Wilmington, MA), underwent bilateral femoral artery interposition grafting with ePTFE grafts. One of two identical grafts...
was implanted into each side of the animal.

Animals were anesthetized with isoflurane in oxygen (4% induction, then maintained with 2% to effect) and subcutaneous flunixin meglumine (2 mg kg\(^{-1}\)). The inguinal area of the rat had the hair clipped and the skin prepared for aseptic surgery. The skin was incised, bupivacaine administered topically to the incision, the femoral sheath was exposed and the artery dissected from the vein. The femoral artery was clamped and approximately 3 mm of the artery was excised. A 7 mm length of ePTFE graft was anastomosed interpositionally using standard microsurgical technique with 10–0 nylon interrupted sutures (Ethicon, Somerville, NJ) using an operating microscope (Olympus, OME-8000, Tokyo, Japan). The graft and surgical site was irrigated with heparinized saline (12.5 U ml\(^{-1}\)) throughout the procedure. Once the sutures were in place the vascular clamps were released and hemostasis was achieved within 2–3 min. Blood flow through the graft was monitored using a laser Doppler flowmeter (Laserflo BPM\(^{2}\) Blood Perfusion Monitor, Vasa-medics, St. Paul, MN) for 30 min following suturing. Patency was additionally confirmed distal to the graft using a “milking test”, whereby the artery distal to the graft was clamped and a section of graft further downstream was gently squeezed (i.e. milked) with loosely closed forceps to push the remaining blood distally. If the distal artery filled with blood when the proximal clamp (closest to the graft) was removed, then the graft was considered patent. No anticoagulants were administered post-operatively and pain was managed by flunixin for two additional post-operative days.

2.8 Implant retrieval

After 7 or 28 days, grafts were explanted. Rats were placed under general anesthesia, the inguinal skin was incised, the grafts were surgically exposed and the graft patency was evaluated by laser Doppler flowmetry and the milking test. No differences in animal behavior were observed prior to explant between animals having patent grants and those having non-patent ones. Grafts were explanted and washed in heparinized saline for 5 min, placed in 10% formalin overnight and transferred to 70% ethanol prior to paraffin embedding.

2.9 Histology

Tissue samples were embedded in paraffin and sectioned for histological evaluation. A series of 5 µm cross-sections were obtained so that each 7 mm graft was visualized by at least one cross-section from the proximal, middle and distal regions. All sections were removed of paraffin using Sub-X (Surgipath Medical, Richmond, IL) and stained with hematoxylin and eosin. Additionally, Masson’s trichrome staining was used to evaluate the neointimal hyperplasia of 28 day graft samples.

Sections were stained for endothelial cells after antigen retrieval with a rabbit anti-human von Willebrand Factor (VWF, 1:12,000 dilution) (Dako, Carpinteria, CA) antibody. VWF sections were incubated with a biotinylated horse anti-rabbit secondary antibody for peroxidase staining (1:200 dilution) (Vector Labs, Burlingame, CA). Human cells were distinguished after antigen retrieval through staining with mouse anti-human major histocompatibility complex class I (MHC-I) antibody (1:50 dilution) (Santa Cruz Biotechnology, Santa Cruz, CA). MHC-I staining was visualized with Alexafluor 488 goat anti-mouse secondary antibody (1:500 dilution) (Invitrogen). Sections with no primary antibody incubation served as negative controls.

Sections were imaged with a Nikon TE2000U microscope and NIS Elements software (Nikon). The luminal coverage of thrombosis and intimal hyperplasia in a given histological section were calculated, respectively, by measuring the cross-sectional area of the graft lumen occupied by the clot or intimal hyperplasia and then dividing by the total luminal cross-sectional area of the graft.

2.10 Statistics

Statistical differences in graft patency were assessed with Fisher’s exact test. All other statistical analysis was performed with a one-way analysis of variance followed by a Tukey–Kramer post hoc test. All error bars represent the standard error of the mean. Values of \( p \) below 0.05 were considered sufficient to indicate a statistically significant difference.

3. Results

3.1. Graft EPC coverage

EPCs formed a confluent cell layer uniformly covering the lumen of the small diameter vascular grafts within 2 days after sodding (Fig. 1). SEM imaging was used to characterize EPC graft coverage after grafts were exposed to conditions that mimic the mechanical manipulation during surgical implantation. EPCs were sodded onto the grafts, cultured for 2 days and observed under three conditions: after no mechanical disruption (static control), after exposure to laminar flow for 5 min (flow) and after clamping along the length of the graft for 30 min followed by laminar flow (clamping + flow). Flow produced a mean shear stress of 100 dynes cm\(^{-2}\), a magnitude 3- to 10-fold higher than the graft would typically experience in a rat femoral artery [26].

Table 1 shows that EPCs sodded onto vascular grafts had approximately 90% graft coverage in the static control. Exposure to flow caused no significant drop in overall cell coverage vs. static conditions (\( p = 0.35 \)). Clamping along the length of the graft with vascular clips followed by exposure to flow reduced EPC graft coverage by approximately 5%. No difference in cell coverage was observed between native and transfected EPCs (\( p = 0.78 \)).

3.2. Activated protein C production in vitro

EPC sodding improved the anti-thrombotic bioactivity of small diameter vascular grafts as measured by the in vitro rate of graft APC production (Table 2). Grafts sodded with EPCs transfected with TM (EPC + AdTM) exhibited APC production that was greater than 10-fold higher than bare control grafts (Bare control) and 4-fold higher than both grafts sodded with native EPCs (EPC) and EPCs transfected with the control vector (EPC + AdCV). The rate of APC production between EPC and EPC + AdCV was similar, showing no detrimental effect of the adenoviral control vector on APC production.

3.3. In vivo testing

Bare off-the-shelf grafts (Bare Control), bare denuded grafts (Denucleated), or denuded grafts sodded with EPC, EPC + AdCV or EPC + AdTM were implanted bilaterally into the femoral arteries of athymic rats as described above. Upon removal of the vascular clamps, bleeding was largely limited to the anastomosis site for 1–2 min before hemostasis was achieved, with minimal weeping through the graft wall. Grafts were observed for approximately 30 min before closing of the surgical site to ensure successful anastomoses.

After 7 days the dominant histological feature was thrombus. Fig. 2 shows representative mid-graft histological sections of 7 day implants with extensive thrombus in bare and denuded grafts, but limited thrombus in grafts sodded with native or transfected EPCs. No intimal hyperplasia was observed after 7 days.
Fig. 3 shows the percentage of luminal cross-sectional thrombosis of grafts implanted for 7 and 28 days. The average area of thrombosis was evaluated along multiple sections for each patent graft. The presence of thrombosis was largely diminished at 28 days, with the graft response dominated instead by the presence of intimal hyperplasia. Comparing the EPC conditions, there were no significant differences in thrombus coverage at 7 days and at 28 days ($p = 0.32$); however, grafts sodded with EPC + AdTM had slightly less thrombus present at 7 days than native EPCs or EPC + AdCV.

3.4. Graft patency

Table 3 lists graft 7 day and 28 day patency rates. All unsodded grafts exhibited poor patency after 7 days and were not tested for 28 days. Sodding with native EPCs or EPCs transfected with AdTM resulted in significantly improved 7 day patency compared to bare or denucleated grafts ($p < 0.05$). Native EPCs and EPC + AdTM exhibited patency rates of 88% and 89%, respectively, after 7 days, and 75% and 88% after 28 days. EPC + AdCV had patency rates of 63% after 7 days and 75% after 28 days. No significant differences were observed in 28 day patency rates between EPC and EP + AdCV or EP + AdTM ($p = 0.91$). Within the same animal, graft patency in one leg did not appear to affect the patency in the other leg.

3.5. Graft intimal hyperplasia

At 28 days the dominant histological feature of all grafts was intimal hyperplasia. Fig. 4 shows representative hematoxylin and eosin (H&E) staining of the proximal, middle and distal regions of native and transfected EPC-sodded grafts after 28 days. All grafts showed little thrombus formation and there were no significant differences in graft cross-sectional thrombus between regions of the graft. Staining for smooth muscle cell actin revealed that the hyperplasia response consisted of smooth muscle cells (see Supplemental Fig. 1). Fig. 5 shows the percent area of 28 day graft lumens occupied by intimal hyperplasia at proximal, middle and distal positions. Hyperplasia was significantly reduced in the proximal and mid-graft regions for EPC and EPC + AdTM sodded grafts vs.
EPC + AdCV grafts ($p < 0.05$). The distal regions of the grafts had the greatest hyperplasia response followed by the proximal region.

### Table 3

<table>
<thead>
<tr>
<th></th>
<th>Bare control</th>
<th>Denucleated</th>
<th>EPC</th>
<th>EPC + AdCV</th>
<th>EPC + AdTM</th>
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<tr>
<td>7 day patency rate</td>
<td>25% (2/8)</td>
<td>25% (2/8)</td>
<td>89% (<em>8/9)</em></td>
<td>63% (5/8)</td>
<td>88% (7/8)</td>
</tr>
<tr>
<td>28 day patency rate</td>
<td>NA</td>
<td>NA</td>
<td>88% (7/8)</td>
<td>78% (6/8)</td>
<td>75% (6/8)</td>
</tr>
</tbody>
</table>

*p < 0.05 vs. Bare and Denucleated grafts.

### 3.6. EC staining

Fig. 6 shows that confluent layers ECs remained present on the lumen of EPC + AdTM sodded grafts after 7 and 28 days as visualized through vWF staining (see arrows). Native femoral artery also stained positively for vWF, whereas denucleated grafts showed no positive staining. Sodded grafts had vWF staining on the graft lumen and within the graft wall after both 7 and 28 days.

Finally, bare grafts were negative for human MHC-I staining (Fig. 7A) while grafts sodded with human EPCs showed positive staining for human MHC-I within the graft lumen after 7 (Fig. 7B) and 28 days (Fig. 7C and D). The 28 day images showed that the neointimal area was likely made up of rat-derived cells as human MHC-I staining was negative in the area of hyperplasia (Fig. 7D).

### 4. Discussion

Vascular graft endothelialization is well known to improve the patency of small diameter vascular grafts [4,5]. These and other studies, however, rely on harvesting autologous blood vessels or microvascular tissue, an invasive procedure that introduces donor site morbidity [4,5,27]. In contrast, EPCs can be isolated non-invas-
sively from samples of peripheral blood drawn from patients with documented CAD [11,28].

Previous studies using animal derived cells indicate that EPCs can improve the performance of synthetic vascular grafts [12–14]. To date, however, only one study has tested the use of human blood-derived EPCs from healthy volunteers on vascular grafts in vivo [29]. Of course, healthy volunteers are not the target population for this therapy; rather, it is patients with significant CAD. The goal of the present study was to assess the in vivo patency rates of small diameter vascular grafts endothelialized with EPCs derived from peripheral blood of patients with CAD.

Recently published in vitro testing showed that patient-derived EPCs overexpressing human TM exhibited increased APC production, reduced platelet adhesion and prolonged clotting times compared to untransfected EPCs [19]. Therefore, this study also assessed whether enhanced graft patency was achieved by transfecting patient derived EPCs with the human TM gene ex vivo prior to graft sodding.

EPCs transfected with AdTM expressed TM that was biologically active and increased the in vitro rate of graft APC production 4-fold relative to grafts endothelialized with native EPCs. APC production per graft luminal area was approximately 30% lower than APC production than on flat TCPS [19]. The 4-fold increase in APC production rate due to TM transfection was similar in magnitude to previous studies with TM-transfected vein graft segments [20,21,30,31].

Bare off-the-shelf and bare denucleated ePTFE positive controls exhibited the expected unacceptably low patency rates after 7 days (2/8 remained patent in both cases). Although removal of trapped air from vascular graft materials by denucleation can decrease clot formation in both small and large animals [24,32] and enhance cell adhesion in vivo [33], no difference in patency rates was observed between the two control groups after the 7 day testing. Control trials out to 28 days were not attempted.

Sodding the lumen of denucleated grafts with EPCs dramatically increased graft patency over the control grafts after 7 days and also maintained improved patency after 28 days. Only one graft failed in each of the experimental trials for native EPCs after 7 and 28 days, and for EPCs transfected with AdTM after 7 days, whereas two grafts failed in 28 day trials for EPCs transfected with AdTM or AdCV. Thrombus was the dominant luminal histological feature after 7 days, while intimal hyperplasia was the dominant feature after 28 days, particularly at the proximal and distal peri-anastomotic regions.

Evidence that sodded human cells remained adherent 28 days after implantation was obtained from graft endothelium that

Fig. 6. H&E and vWF staining of the native rat femoral artery, denucleated graft and EPC + AdTM sodded graft at 7 and 28 days after implantation. Positive vWF staining (brown) is highlighted with arrows (scale bar 100 μm).

Fig. 7. Staining of ePTFE grafts for human MHC-I (green) and cell nuclei (blue). Bare grafts had cell nuclei lining the outer walls of the graft and within the clot but showed no human MHC-I staining (A). Grafts sodded with native EPCs showed positive MHC-I staining near the graft lumen 7 days after implantation and nuclei were observed on the lumen and within the graft wall (B). After 28 days, sodded human EPCs remained on the graft original ePTFE lumen but not on the region of hyperplasia (C and D).

stained positive for vWF – which was not specific for human or rat – and also stained positive for human specific MHC-I. However, regions of intimal hyperplasia that stained positively for vWF were negative for human MHC-I. We speculate that ECs in the hyperplasia regions may be of rat origin rather than the originally sodded human ECs. The migration of ECs from the anastomosis is commonly observed in vivo, and is more pronounced in animal models than in humans [34]. It unlikely that the rat ECs were derived from circulating EPCs [35], or from ECs from migrating through the graft wall as the ePTFE graft material utilized in the present study was low porosity [36].

Overall, native and TM-transfected EPC sodded grafts had lower levels of thrombus formation and intimal hyperplasia compared to EPCs transfected with the control vector; however, there were no significant differences in either case between the native and TM-transfected EPCs groups. This observation differed from previous in vivo studies where autologous vein graft over-expression of TM reduced thrombosis [20,21] and intimal hyperplasia [21].

In spite of in vitro evidence to the contrary, it was surprising that TM-transfected EPCs had no further benefit than native EPCs in increasing patency and reducing both thrombus formation and intimal hyperplasia. We offer the following possible explanations for this unexpected finding.

First, vector concentration was optimized in vitro for human EPCs, balancing cell death with robust TM expression; however, our previous studies also showed that adenovirus-transfected EPCs had lowered proliferation potential after transfection [19].

Second, in vitro characterization studies showed that TM up-regulation lasted for only 1 week before returning to the basal level [19], indicating that longer durations of TM expression may be necessary to reduce intimal hyperplasia.

Third, TM-mediated anti-thrombotic activity may have been suppressed in vivo due to the differences in human TM activity compared to the rat coagulation substrates. Human TM is capable of binding rat thrombin and accelerating the conversion of human protein C to APC (data not shown). It remains unknown, however, whether human TM bound to rat thrombin can accelerate the activation of rat protein C. Pig TM, for example, activates human protein C at only 10% of the rate of human TM [37]. Roussel et al. [37] suggested that the positioning and conformation of the pig TM active site may not be optimal for interacting with bound human protein C. Additional studies using human TM in rats showed that soluble human TM was capable of binding thrombin but was unable to activate rat protein C [38]. This study was limited by the lack of commercial availability for rat protein C. That notwithstanding, it is important to note that human TM overexpression has been used in rat [31] and rabbit [20,21] studies, and showed a reduction in thrombus [20,21,31] and intimal hyperplasia [21].

Fourth, blood flow dynamics may have contributed to intimal hyperplasia that countered the effect of TM over expression. As blood travels from the native femoral artery into a graft of larger diameter, the expansion in lumen diameter decreases blood velocity, which can result in eddy formation and recirculating flow. This disturbed blood flow is often found at the vascular anastomosis, and is widely acknowledged to contribute to intimal hyperplasia (reviewed by Chiu and Chien [39]). Furthermore, laminar fluid flow increases the expression of the TM transcription factor KLF-2 [40], whereas disturbed flow suppresses KLF-2 expression and other anti-thrombotic and anti-cell proliferation molecules such as endothelial nitric oxide synthase [41,42].

Finally, intimal hyperplasia was observed at both proximal and distal sites. While hyperplasia is most common at the distal anastomosis, hyperplasia at the proximal anastomosis may have been due to recirculating flow. Computational simulations with an idealized geometry (detailed in the Supplement Information) indicated the presence of proximal recirculating flow during the diastole portion of the heart cycle. In agreement with prior studies modeling microvascular anastomoses, recirculation may have been a major contributor to the observed proximal hyperplasia [43].

The athymic rat femoral artery model was employed to allow the testing of human EPCs in a rodent model with relatively straightforward microsurgical access. The model provided lowered flow rates than an abdominal aortic interpositional graft, and therefore is a more stringent test of patency. Future studies require in vivo testing in a large animal model that can more closely mimic the geometry and flow rates of bypass grafts used clinically.

5. Conclusions

In this study we have shown that patient-derived endothelial cells, called EPCs, can be isolated noninvasively from small samples of peripheral blood, expanded to higher cell densities, and sodded onto the lumen of small diameter grafts to enhance graft patency in a thrombotic animal model. While TM overexpression improved graft performance vs. EPCs transfected with a control vector, there was no observed difference between native untransfected and TM-transfected EPCs. Thus, untransfected EPCs may be sufficient to improve the patency of small diameter vascular grafts. The use of untransfected EPCs also limits the risk of pre-existing immunity and the development of adaptive immune responses against viral proteins produced due to the adenoviral vector [43]. These results should encourage testing of EPCs in larger animal models with more physiologically relevant geometries and flow profiles.

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Appendix A. Figures with essential colour discrimination

Certain figures in this article, particularly Figures 2 and 4–7, are difficult to interpret in black and white. The full colour images can be found in the on-line version, at doi:10.1016/j.actbio.2011.09.002.

Appendix B. Supplementary data


References