Inflammation is a natural phase of the wound healing response, which can be harnessed for the in situ tissue engineering of small-diameter blood vessels using instructive, bioresorbable synthetic grafts. This process is dependent on colonization of the graft by host circulating cells and subsequent matrix formation. Typically, vascular regeneration in small animals is governed by transanastomotic cell ingrowth. However, this process is very rare in humans and hence less relevant for clinical translation. Therefore, a novel rat model was developed, in which cell ingrowth from the adjacent tissue is inhibited using Gore-Tex sheathing. Using this model, our aim here was to prove that functional blood vessels can be formed in situ through the host inflammatory response, specifically by blood-borne cells. The model was validated by implanting sex-mismatched aortic segments on either anastomoses of an electrospun poly(ε-caprolactone) (PCL) graft, filled with fibrin gel, into the rat abdominal aorta. Fluorescent in situ hybridization analysis revealed that after 1 and 3 months in vivo, over 90% of infiltrating cells originated from the bloodstream, confirming the effective shielding of transanastomotic cell ingrowth. Using the validated model, PCL/fibrin grafts were implanted, either or not loaded with monocyte chemotactic protein-1 (MCP-1), and cell infiltration and tissue development were investigated at various key time points in the healing cascade. A phased healing response was observed, initiated by a rapid influx of inflammatory cells, mediated by the local release of MCP-1. After 3 months in vivo, the grafts consisted of a medial layer with smooth muscle cells in an oriented collagen matrix, an intimal layer with elastin fibers, and confluent endothelium. This study proves the regenerative potential of cells in the circulatory system in the setting of in situ vascular tissue engineering.

Introduction

There is a clear clinical demand for suitable, artificial, small-diameter (Ø < 6 mm) blood vessels, in particular for lower limb peripheral artery disease with an overall prevalence estimated to be in the range of 3–10%. Current artificial small-caliber vessels have poor patency rates compared with their living counterparts, which is inherently due to their nonadaptive constitution. Tissue engineering has been proposed to overcome these shortcomings by delivering living small-diameter grafts. However, traditional tissue engineering approaches comprise lengthy and costly in vitro procedures. In situ tissue engineering is emerging as a promising alternative, in which an acellular bioresorbable synthetic graft is implanted, allowing the body to populate the scaffold with host cells and develop extracellular matrix while the scaffold is degraded. This approach is built on the notion that inflammation is not merely a detrimental response to biomaterials, but when harnessed properly, can be exploited to induce a natural regenerative response. By omitting time-consuming cell expansion and bioreactor phases, this method can deliver cost-effective, off-the-shelf available grafts.

Recent studies following the in situ tissue engineering principle have demonstrated colonization and subsequent remodeling of a biodegradable synthetic graft into functional
neoaorties in various animal models.9–13 However, the cellular and molecular mechanisms underlying this functional regeneration remain elusive.4,14 An inflammation-mediated response has been suggested, comprising an initial rapid recruitment of immune cells to the scaffold, followed by an influx of tissue cells that remodel the graft into a native-like artery.15 Monocyte chemotactic protein-1 (MCP-1) has been identified as one of the key mediators in this process, and local release of exogenous MCP-1 from biodegradable scaffolds was shown to enhance de novo tissue formation.15

To translate preclinical results to human application, it is essential to recognize and predict regenerative processes (cell colonization and tissue development) representative for the human situation. From previous small animal studies employing degradable and nondegradable grafts, it is clear that transmural migration is a predominant source of macrophages as well as α-smooth muscle actin (α-SMA)-expressing cells in rodents.16–18 Furthermore, transanastomotic ingrowth of mature endothelial cells (ECs) and smooth muscle cells (SMCs) was identified as the predominant source of tissue cells in the rodent model.19 However, given the fact that transanastomotic endothelialization in humans is restricted to the immediate perianastomotic region, the question arises how predictive these model-dependent processes are for the clinical situation, especially considering that vascular prostheses for humans often require extensive anatomic lengths.20,21 Moreover, this suggests that physiological remodeling of a graft in humans is heavily dependent on the contribution of circulating cells.

Therefore, our aim was to prove that functional blood vessels can be formed in situ through the host inflammatory response solely by circulating cells. To isolate the role of blood-borne cells in the regenerative process, we employed a more predictive rat model, in which the graft is isolated from surrounding tissues, both transmurally and transanastomotically. To preclude cell ingrowth from the adjacent tissue, highly porous electrospun poly(e-caprolactone) (PCL) tubes were shielded by impenetrable Gore-Tex sheet material on both anastomoses and enclosing the adventitial side of the PCL scaffold over the entire length. To enhance rapid cell recruitment from the circulation, we incorporated fast-releasing MCP-1 into our scaffold using fibrin gel. Grafts were implanted as abdominal aortic interposition grafts in rats and were explanted at different time points in the remodeling cascade. We hypothesized that (i) extracellular matrix-producing cells can be recruited from the circulation and that (ii) increased early monocyte recruitment to the graft by MCP-1 has a positive effect on long-term tissue remodeling and regeneration.

Materials and Methods

Experimental animals

Seventy-six healthy Sprague-Dawley rats, 250–300 g, were used, purchased from Harlan Laboratories. Each animal was fed ad libitum and was housed in groups in an environment maintained at room temperature for 24 h a day and with a 12-h light–12-h dark cycle. Two studies were performed; Study 1 was dedicated to validate the model, and Study 2 was dedicated to investigate the effect of MCP-1 in the scaffold. In Study 1, Gore-Tex shielded grafts (n = 12) were compared with grafts without shielding (n = 12) (Fig. 1A–C). To study general cell infiltration in both groups over time, grafts were explanted at 1 day (n = 2), 3 days (n = 2), 1 week (n = 2), 2 weeks (n = 2), and 3 months (n = 4). To specifically study the efficacy of the model in transanastomotic shielding, a sex-mismatch experiment was performed, in which sections of male rat aorta were interpositioned proximally and distally to the graft in female rats, either shielded or unshielded with Gore-Tex (n = 12 per group) (Fig. 1D, E). These grafts were explanted after 1 or 3 months (n = 3 per group, per time point) for fluorescent in situ hybridization (FISH) analysis.

Study 2 was dedicated to study the effect of MCP-1, comparing fibrin-seeded grafts with incorporated MCP-1 (n = 20) versus fibrin-seeded controls (n = 20). All grafts in this study group were shielded with Gore-Tex. Grafts were explanted at 1 and 3 days, 1 and 2 weeks, and 3 months (n = 4 per time point). All procedures were performed in accordance with and approved by the Institutional Animal Care and Use Committee of the University of Utrecht, The Netherlands.

Scaffold fabrication

Electrospun (PCL) tubular scaffolds were manufactured and provided by Xeltis BV. In brief, PCL (Purasorb; Purac Biochem) was dissolved in chloroform at 20% (w/w) and driven through a horizontally fixed nozzle at high voltage (15 kV) toward a grounded rotating cylindrical copper target (Ø 2 mm) at 15 cm distance. The resulting tube diameter, wall thickness, and average fiber diameter were determined with scanning electron microscopy (SEM; Quanta 600F, Fei). The materials were sterilized by gamma irradiation. Before implantation, the PCL tube was cut to size and the vessel wall was impregnated with fibrin gel (5 U/mL thrombin+5 mg/mL fibrinogen; Sigma). Depending on the test group, MCP-1 (4 μg/mL; Chemicon) was added to the fibrin solution. To measure the release rate of the MCP-1 from the graft, a cumulative leakage experiment was performed in vitro. For this, MCP-1-loaded scaffold samples were incubated in culture medium (RPMI-1640; Sigma) supplemented with 10% (v/v) fetal bovine serum (FBS Gold; PAA Laboratories) over a period of 7 days. At each time point (0, 0.75, 1.5, 3, 8, 24, 48, 72, 120, 168 h), all supernatant was collected, replaced with fresh medium, and samples were stored at −80°C until further analysis. MCP-1 content was determined cumulatively using a human ELISA kit, according to the manufacturer’s protocol (RayBiotech) (n = 3).

Surgical procedure

Before implantation, graft composites were prepared depending on the study group. To shield the electrospun grafts, an end-to-end anastomosis was made to a 4 × 10 mm² impenetrable Gore-Tex strip (Preclude Pericardial Membrane; Gore Medical) using 10-0 interrupted sutures distally and proximally of the electrospun tube. Additionally, Gore-Tex was wrapped around the PCL in samples, creating an impenetrable outer layer (Fig. 1A–C). For the FISH experiments, sex-mismatched segments of aorta were interpositioned on the proximal and distal anastomoses between the native aorta and the graft (Control group, n = 3 per time point; Fig. 1D) or Gore-Tex (Test group, n = 3 per time point; Fig. 1E). Animals were anesthetized using isoflurane gas. Once adequate
anesthesia was achieved, a midline laparotomy was performed and the abdominal viscera were lateralized for exposure of the abdominal aorta and the inferior vena cava. After separation of the aorta from the inferior vena cava and surrounding tissue, the segment of the abdominal aorta between the renal arteries and the aortic bifurcation was occluded with microvascular clamps. The aorta was transected and the graft composite was introduced with end-to-end anastomosis performed at both the proximal and distal ends using interrupted sutures. After removing the vascular clamps, the pulsatile flow was confirmed in the aorta distal to the graft (Fig. 1A). The abdomen was closed in two layers. Immediately after implantation, all scaffolds were exposed to arterial hemodynamic conditions. There was no heparin administration during or after surgery. Animals recovered on warmed pads to promote blood flow through the grafts. Before rats returned to their cages, they were assessed for evidence of hind limb paralysis or acute graft thrombosis. Postoperative buprenorphine was given intraperitoneally.

At termination, animals were anesthetized using isoflurane. Tubes were dissected free from surrounding tissue. The Gore-Tex on the outer layer was opened and the PCL graft was carefully explanted. All explants were examined with immunohistochemistry. In Study 2, two explants per time point were additionally analyzed with quantitative polymerase chain reaction (qPCR) to determine the expression of genes involved in the inflammatory response. Furthermore, blood samples were collected and analyzed with ELISA to determine systemic protein levels of MCP-1 in the blood serum (Supplementary Data).

### Immunohistochemistry

Explants were fixed in formalin 10% before pre-embedding in 1% (w/v) agar (Eurogentec), followed by embedding in paraffin. Consecutive 4-µm sections were stained with Mayer’s hematoxylin and eosin, Verhoeff van Gieson, and 0.1% (w/v) Sirius red F3B (Gurr BDH) in saturated picric acid solution, respectively. Immunohistochemical stainings were performed on 4-µm sections after deparaffinization and dehydration. Sections were stained for myeloperoxidase (A398, 1:2000, MPO; Dako), CD68 (1:400, MCA341GA; Serotec), CD163 (1:75, MCA342GA; Serotec), CCR7 (Cell Applications, CG1678, 1:15000), CD34 (R&D Systems, AF4117, 1:100), (α-SMA; Sigma, A2547, 1:32,000), and Von Willebrand factor (vWF; Dako, A0082, 1:1600). All antibodies were diluted in PBS/BSA/Azide. Following deparaffinization, antigen retrieval was performed in citrate (For CD68, CCR7, and MPO), pepsin (CD163), or EDTA (vWF) before blocking with 10% normal goat serum (α-SMA and CCR7) or Ultra V block (MPO; Thermo Scientific). Sections were blocked for endogenous peroxidase when a horseradish peroxidase (HRP)-conjugated secondary antibody was used. Appropriate BrightVision signal amplification was used for labeling of primary antibody binding sites: poly AP anti-mouse for α-SMA, poly HRP anti-mouse (CD68 and CD163), and poly HRP anti-rabbit (CCR7, MPO, and vWF) (Immunologic). Slides were subsequently incubated with liquid permanent red substrate-chromogen (Dako) or diaminobenzidine substrate-chromogen (Sigma) and counterstained with Mayer’s hematoxylin. Slides were dehydrated in alcohol changes and

**FIG. 1.** Implantation procedure and experimental groups. (A) Camera microscopy images of the graft implantation with, in bright white, the Gore-Tex sheet. The aorta before (i), after transection (ii), after implantation of the graft (iii), and directly after removing the vascular clamps (iv). (B–E) Schematic representation of the experimental groups in Study 1, with the shielded grafts (C) versus unshielded control (B) and the sex-mismatched shielded graft (E) versus the unshielded control (D). Color images available online at www.liebertpub.com/tea
coverslipped using pertex. Double-label immunohistochemistry for αSMA and vWF was performed after EDTA antigen retrieval by incubating sections for 1 h at room temperature with a mix of 1:32,000 mouse anti- αSMA and 1:1600 rabbit anti-vWF antibodies. The sections were then incubated with anti-mouse Alexa Fluor 488 and anti-rabbit Alexa Fluor 555-conjugated secondary antibodies (1:200; Invitrogen) for 30 min. Cell nuclei were counterstained with 4′,6-diamidino-2-phenylindole (DAPI).

Quantitative histologic and immunohistological analysis

Two investigators, who were blinded to the investigated groups and the time point of explantation, independently conducted the analysis. Sections were photographed using a Nikon E800 microscope with ACT-1 software. Wall thickness and lumen diameter were measured manually under a 2× objective lens. Wall thickness was measured in two sections of each graft and in four locations per section. The lumen diameter was measured in two sections of each graft and in two directions per section. Cellularity was studied under high-power magnification (high-power field [hpf], 40× objective lens; area comprising 0.034 mm²), with cell sizes being comparable between groups. The total number of cells in four random hpf areas per PCL tube was manually counted using ImageJ software. Quantification of aligned collagen was performed with picrosirius red stains and circularly polarized light. Stained sections were digitally photographed and converted into gray value images and regions of interest (ROIs) were drawn lining the graft. The average gray value per ROI was calculated using ImageJ software. A quantitative analysis of the immunopositive cells of each stained slide was performed at four random hpfs. The portion of area staining positively for the marker within the four hpfs was counted and summed for each image. The mean of the sums for four high-power images was then calculated for each sample. Area of positive staining was measured as a percentage of the total from binarized images.

Fluorescent in situ hybridization

To identify X- and Y-chromosomes, FISH was performed on fresh, sectioned paraffin sections using Rat idetect Chr X FISH Probe Green and Chr Y FISH Paint Probe Red (ID Labs, IDRF1067 and IDRR1070, respectively). Following paraffinization, slides were pretreated with 0.2N HCl for 20 min at room temperature, citrate buffer (pH 6.0) for 20 min at 100°C, and digested with 0.1% proteinase K for 10 min at 37°C. After dehydration in graded ethanol, 7.5 µL of probe mixture was added. Sections were then denatured for 5 min at 69°C, cooled on ice for 5 min, and hybridized overnight at 37°C in a ThermoBrite. The next day, slides were washed in 0.4× SSC/0.3% NP-40 at 73°C, 2× SSC/0.1% NP-40 at room temperature, and 2× SSC for 2, 1, and 5 min, respectively. After washing, the sections were counterstained with DAPI, dehydrated, and mounted in Vectashield (Vectorlabs, H-1000). For quantification of the FISH results, at least 30 images per sample were analyzed by two investigators blinded to the experimental group, sample location, and time of implantation.

Statistical analysis

Statistical differences were determined using two-way ANOVA with time and test factor as the variables, with Bonferroni post-tests to compare the means per time point using GraphPad Prism software. p < 0.05 was considered statistically significant.

Results

Grafts are constructed from porous electrospun PCL and dense Gore-Tex

SEM analysis showed the dense microstructure of the Gore-Tex, with an intermodal distance <1 μm. In contrast, the PCL graft was highly porous with interconnected pores and an average fiber diameter of 9 μm (range 8.4–9.7 μm) (Fig. 2A, B). The inner diameter of the graft was ~2 mm with an average wall thickness of 275 μm (range 210–340 μm). Grafts were seeded with a fibrin gel with or without MCP-1. In vitro leakage experiments revealed a burst release of MCP-1 within the first 3 h, after which the release gradually leveled off (Fig. 2C).

Gore-Tex shielding led to a significant reduction of cell ingrowth from neighboring tissues

All grafts remained patent and there were no signs of stenosis or thrombosis in all groups studied. Lumen diameter and wall thickness remained constant throughout the whole study (1.8 mm ± 0.1 and 385 μm ± 108, respectively), with no signs of intimal hyperplasia or aneurysms.Macroscopically, granulation tissue was seen on the adventitial side of Gore-Tex sleeves (Fig. 3A, B, D, E). No cell infiltration into the dense adventitial sleeve was detected at any time point. Adventitial Gore-Tex shielding led to a significant reduction in total cellularity at all intervals compared with unshielded controls, with the largest difference at day 1 and day 14 (Fig. 3C).

For the FISH experiments, sections were taken from five different locations in the graft as well as the native (female) aorta and the implanted male segment (as indicated in Fig. 4) and analyzed for the presence of Y-chromosome using FISH. In the Control group, the average fraction of male cells detected in the graft was 37.4% ± 12.1% and 43.1% ± 9.8% after 1 and 3 months, respectively. Moreover, after 3 months of implantation, the distribution of male cells was homogenous throughout all sections of the Control group, with an average of 44.1% ± 14.5% (Fig. 4A). In contrast, in the Test group, the fraction of male cells detected in the graft was severely limited, with an average of 9.4% ± 3.2% and 10.1% ± 7.7% after 1 and 3 months, respectively. There were no marked differences between the different locations in the graft (Fig. 4B). After 3 months, the native (female) aorta contained on average 38.1% ± 9.3% male cells in this group, indicating a comparable migration of male cells to unshielded tissues compared with the Control group.

These results demonstrate effective isolation of the graft using Gore-Tex sheathing in this rat model, both transmurally and transanastomotically.

MCP-1 recruits immune cells in the early inflammatory phase

Cells penetrated immediately after implantation through the entire thickness of the PCL in all grafts (Fig. 5A–C). Directly after implantation, mainly neutrophilic granulocytes infiltrated all grafts, with a significant increase in
overall cell infiltration in the MCP-1 group compared with the control group at day 1 (Fig. 5D). There was a marked increase in CD68<sup>+</sup> macrophage infiltrates in the MCP-1 and control group up to day 7 and day 14, respectively, with an even distribution throughout the graft. At day 7, the concentration of macrophages was significantly higher in the MCP-1 group compared with the control group. After the first 2 weeks, macrophage numbers gradually decreased in both groups with a significantly higher macrophage concentration in the MCP-1 group after 3 months (Fig. 5E). In both groups, macrophage giant cells were seen. Macrophage subtypes were characterized by CCR7 (M1) and CD163 (M2) expression. Surprisingly, CD163<sup>+</sup> cells were detected very sparsely (<1%). CCR7 expression was detected, but only a small fraction of the CCR7 staining double-stained positive for the pan-macrophage marker, CD68.

**Initial MCP-1 release results in enhanced tissue formation in the healing phase**

A progressive increase in CD34<sup>+</sup> cells was observed in both groups, with no significant differences between groups up to day 14 (Fig. 6A, B). At day 84, the expression of CD34 was significantly higher in the MCP-1 group compared with

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**FIG. 2.** Graft composition. Schematic representation of the monocyte chemotactic protein-1 (MCP-1)-loaded graft (A) with scanning electron microscopy images (B) showing a transection (i), the luminal surface of the Gore-Tex (ii), the Gore-Tex anastomosed to the poly(ε-caprolactone) (PCL) (iii), and the luminal surface (iv) and transection of the porous PCL scaffold (v). (C) MCP-1 release curve as determined in vitro. Scale bars represent 500 μm (i and v), 40 μm (ii and iv), and 100 μm (iii).

**FIG. 3.** Efficacy of transmural isolation. Scaffold cellularity of the shielded (A, D) versus nonshielded grafts (B, E). Cellularity decreased significantly in the grafts shielded with Gore-Tex when compared with the control group (98 ± 6 cells per high-power field [hpf] vs. 137 ± 17 cells per hpf, p = 0.03 at 84 days), with the largest differences at day 1 (9 ± 4 cells per hpf vs. 103 ± 12 cells per hpf) and day 14 (82 ± 6 cells per hpf vs. 163 ± 15 cells per hpf) (C). Color images available online at www.liebertpub.com/tea
the control group. For α-SMA expression, no significant differences were detected between groups for all time points (Fig. 6C, D). In both groups, the first α-SMA+ cells were detected at day 7. After 3 months, the myofibroblast population formed a circumferentially oriented neointima layer of SMCs under an endothelium in both groups (Fig. 6C). vWF staining revealed no significant differences between groups, with first evidence of ECs in the grafts at day 7 in both groups. Small patches of vWF-expressing cells lined the lumen, growing more confluent through week 2, before reaching a fully confluent monolayer at 3 months (Fig. 7J–L). This was similar in both the MCP-1 group and the control group. At day 14, the first signs of collagen fibers were detected in both groups. After 3 months, circumferentially orientated bundles of collagen fibers were found lining the endothelial and outer layers of the graft, with more loose fibers toward the center of the wall (Fig. 7A–F). Aligned collagen was markedly increased after 3 months in both groups, with a significantly higher expression of aligned collagen in the MCP-1 group compared with the control group (Fig. 7M). In the MCP-1 group, first signs of elastin were seen in the Verhoeff van Gieson-stained slides in three of four rats after 84 days. In the control group, only one of four rats stained positive (Fig. 7G–I).

**Myofibroblasts do not originate from endothelial-to-mesenchymal transdifferentiation**

All explants were analyzed for occurrence of endothelial-to-mesenchymal transdifferentiation (endoMT) using immunofluorescent double staining of vWF and α-SMA. No colocalization of vWF and α-SMA was detected in any of
the explants of all time points, apart from a single cell showing colocalization of vWF and α-SMA at the 14-day time point, indicating possible endoMT (Fig. 8).

Discussion

In situ tissue engineering of vascular grafts using synthetic scaffolds could deliver an off-the-shelf alternative to current vascular prostheses. In the context of clinical translation, we aimed to achieve an early attraction of circulating cells into a graft releasing MCP-1, using a novel rat model that prevents transanastomotic pannus overgrowth, as well as periadventitial ingrowth. Our results demonstrate that (i) extracellular matrix-producing cells can be recruited from the circulation and that (ii) increased early monocyte recruitment to the graft by MCP-1 has a positive effect on long-term tissue remodeling and regeneration. Grafts underwent extensive cellularization by blood-derived cells, followed by tissue formation throughout

![Image](url)
the entire PCL graft, with all grafts remaining patent up to 3 months follow-up, without signs of aneurysms or intimal hyperplasia. Driven by early MCP-1 release, our remodeled synthetic graft showed several similarities to the native aorta after 3 months in vivo; the intimal layer with ECs lining the lumen, with early signs of a developing elastic lamina, and a medial layer consisting of SMCs in an aligned collagen matrix.

Ingrowth of host cells into vascular implants in rodent models is typically dominated by two mechanisms: transmural (mainly macrophages and α-SMA-expressing cells) and transanastomotic infiltration (ECs and SMCs).16–19 The infiltration of cells through these routes is typically rapid and progressive and may obscure any contribution of fallout healing when using these animal models. However, the consistent observation that transanastomotic cell ingrowth, in particular, is limited in humans, makes that such rodent models may have limited predictive value for the clinical outcome, and the importance of a transanastomotic isolation model has been advocated previously.20,21 Moreover, since recruitment of blood-derived cells represents a valuable target for colonization of acellular grafts, this study was specifically aimed at isolating the role of circulating cells in the regenerative process (fallout healing) of a degradable synthetic graft showed several similarities to the native aorta after 84 days. Collagen (PSR staining, polarized light) content in grafts without MCP-1 significantly higher in the A–G, B, and native rat aorta (C). Endothelial cell (EC) lining (von Willebrand factor) in grafts without MCP-1 (J), grafts with MCP-1 (K), and native rat aorta (L). Collagen alignment was significantly higher in the +MCP-1 group after 84 days (21 ± 2.9 vs. 11.9 ± 1.7 area fraction per hpf) (M). Smooth muscle cell lining in the graft (N) and native aorta (O). Scale bars represent 500 μm (A–C), 100 μm (D–F), 500 μm (N), 200 μm (O), and 20 μm (G–L). Arrows indicate elastin fibers (H). *p < 0.001.
FIG. 6.

A

CD34

no MCP-1

+ MCP-1

7 days

14 days

84 days

B

CD34

Area Fraction (%)

Time (days)

#

C

αSMA

no MCP-1

+ MCP-1

7 days

14 days

84 days

D

FIG. 7.

Aligned Collagen

Native Aorta

M

Aligned Collagen

Area Fraction (%)

1 day

7 days

14 days

54 days

84 days

+ MCP-1

- MCP-1

#
Macrophage function during inflammation and healing is governed by polarization state. Unexpectedly, we detected only very sparse expression of CD163, a typical surface marker for the M2 macrophage type that was reported in previous findings. However, on the gene level, we did observe indications, which suggest a shift in macrophage polarization in response to MCP-1 (Supplementary Fig. S2). Additionally, peripheral-derived macrophages have been shown to acquire myofibroblast-like characteristics during the foreign body response in rodents. The observed absence of coexpression of vWF and α-SMA suggests that the process of endoMT was virtually absent during the in vivo time frame of the current study, implicating that the vast majority of α-SMA+ myofibroblasts did not originate from recruited ECs through endoMT. Taking into account that transmural and transanastomotic infiltration of mature SMCs was hampered, combined with the apparent absence of endoMT, it is plausible that the infiltrating blood-derived α-SMA+ cells descend directly from transdifferentiated macrophages. In addition to peripheral leukocytes, circulating progenitor cells may contribute to vascular regeneration in terms of endothelialization as well as extracellular matrix production. Blood stream-derived cellularization of vascular grafts was proven in dogs, represented by patchy endothelial coverage with underlying α-SMA+ SMCs. Immunohistochemical characterization of infiltrated cells in our grafts demonstrates an increasing presence of CD34+ cells over time, with a significant upregulation in the MCP-1 group. CD34 is expressed by various circulating progenitor populations, including fibrocytes, which can differentiate into mature myofibroblasts upon activation by transforming growth factor-β and through cross talk with macrophages. On the gene level, this coincided with a trend in upregulation of stromal cell-derived factor-1α (SDF-1α), which is a known attractant for CD34+ progenitor cells (Supplementary Fig. S2). MCP-1 has been shown to mobilize angiogenic monocytes, which may be responsible for the increased SDF-1α expression and secondary recruitment of progenitors. The burst release of exogenous MCP-1 in the present study led to a remarkable effect on downstream events up to 3 months later, ignited by an increased early influx of leukocytes. Although the exact chain of events between the early and late-term observations in this study remains to be clarified, our results suggest that MCP-1-recruited monocytes/macrophages have a dual role; indirectly by creating a biochemical milieu favorable for attraction of secondary progenitor cells and directly through possible contribution of macrophage-to-myofibroblast transdifferentiation.

In conclusion, the results of the current study prove the in situ regenerative potential of circulating cells in rats. In these conditions, relevant for the human situation, a short local burst of exogenous MCP-1 led to improved or at least accelerated neotissue formation and organization, with a layered structure of intimal endothelium and a medial layer of SMCs in an oriented collagen matrix with elastin fibers. These findings represent a valuable contribution to the understanding of the regenerative potential of circulating cells in various clinical settings.

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Disclosure Statement

No competing financial interests exist.

References


by mesenchymal stromal cells seeded in 3D scaffolds and circulating leukocytes in physiological flow. Biomaterials 35, 9100, 2014.


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