In Vitro and In Vivo Effects of Deoxyribonucleic Acid–Based Coatings Functionalized with Vascular Endothelial Growth Factor

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ABSTRACT

Vascularization is important in wound healing and essential for tissue ingrowth into porous tissue-engineering matrices. Furthermore, peri-implant tissue vascularization is known to be important for the functionality of subcutaneously implanted biosensors (e.g., glucose sensors). As a first exploration of the use of deoxyribonucleic acid (DNA)-based coatings for the optimization of biosensor functionality, this study focused on the effect of DNA-based coatings functionalized with vascular endothelial growth factor (VEGF) on in vitro endothelial cell behavior and vascularization of the peri-implant tissue in vivo. To that end, DNA-based coatings consisting of poly-D-lysine and DNA were functionalized with different amounts of VEGF (25 and 250 ng) and compared to non-coated controls and non-functionalized DNA-based coatings. The results demonstrated the superiority of VEGF-functionalized DNA-based coatings in increasing endothelial cell proliferation and migration in vitro over non-coated controls and non-functionalized DNA-based coatings. In vivo, a significant increase in vascularization of the peri-implant area was observed for VEGF-functionalized DNA-based coatings. Because no dosage-dependent effects were observed, future experiments should focus on optimizing VEGF concentration for this purpose. Additionally, the administration of VEGF in combination with other (pro-angiogenic) factors should be considered.

INTRODUCTION

A n important physiological aspect of healing around biomaterials is vascular development. For tissue-engineering matrices, particularly those that are pre-seeded with cells, fast development of a vascular network is required for tissue ingrowth and cell survival. Also, for solid devices such as subcutaneously implantable biosensors, the development of a contiguous vascular network is important for their functionality.1,2 For instance, previous studies have demonstrated that the formation of a largely avascular fibrous tissue capsule and the resulting increased diffusion distance between sensor and blood supply to the sensor surrounding tissue diminishes the responsiveness of implanted biosensors to fluctuations in analytes in the interstitial fluid.1,2 Furthermore, Gerritsen et al. showed that inflammatory cells and low-molecular-weight serum components influence the performance of amperometric glucose sensors.3 The basis for these findings was the secretion of enzymes by stimulated granulocytes, which consumed the hydrogen peroxide formed by the oxidation of glucose, and the formation of a diffusional barrier for glucose and the inhibition of glucose oxidase activity by serum components. Consequently, the authors of these publications suggested

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that biosensor functionality is likely to improve if the vascularity in the direct vicinity of the device is increased and inflammatory responses to the device are minimal.

The development of the vasculature depends on two processes (vasculogenesis and angiogenesis), which are responsible for de novo vessel formation and sprouting from already existing vessels, respectively. Therapeutical modulation of vascular supply, however, is limited to angiogenesis, because vasculogenesis is believed to occur only during early embryogenesis. Angiogenesis is a complex process that involves subtle interactions between cells, extracellular matrix components, and regulatory molecules. Decisive steps in this process are the localized proteolytic degradation of the subendothelial basement membrane, migration of endothelial cells, the formation of sprouts, remodeling of extracellular matrix, and the formation of anastomoses, which together precede the establishment of blood flow. The delivery of growth factors with pro-angiogenic action is a potentially powerful tool in control over vascular development. A widely used pro-angiogenic growth factor in tissue engineering is vascular endothelial growth factor (VEGF). VEGF affects endothelial cells through positive effects on proliferation, migration, tubule formation, survival, and integrin expression.

To increase vascularity around subcutaneously implanted biosensors, we hypothesized that VEGF functionalization of deoxyribonucleic acid (DNA)-based coatings could be beneficial. It has been demonstrated that these coatings, whose generation is based on the layer-by-layer deposition of cationic poly-D-lysine (PDL) or poly(allylamine hydrochloride) with anionic DNA, are fully histocompatible upon subcutaneous implantation, showing no adverse reactions in terms of inflammation and wound healing. Furthermore, it was shown that these DNA-based coatings are suitable for functionalization with bone morphogenetic protein 2 (BMP-2), an osteo-inductive factor, using different loading modalities and that this factor retained its biological activity. Another important feature of the fabrication method of these DNA-based coatings is its simplicity, which allows a wide variability of substrate materials (e.g., ceramics, metals, and polymers) with almost unrestricted substrate geometry. In view of the described disadvantageous effects of lack of vascularity on biosensor functionality, we hypothesized that VEGF functionalization of DNA-based coatings would be beneficial for the vascularization of the peri-implant tissue. Consequently, the aim of this study was to evaluate the effect of VEGF-functionalized DNA-based coatings on endothelial cell behavior in vitro and vascular development in vivo. Therefore, [PDL/DNA]_5 coatings on glass substrates were functionalized with a small (25 ng) or large (250 ng) amount of VEGF to study dosage-dependent effects. These VEGF-functionalized DNA-based coatings, as well as non-functionalized coatings and non-coated controls, were used in an in vitro experiment to study the proliferation, migration, and morphology of endothelial cells. Additionally, these four types of substrates were implanted in the backs of rats in an in vivo animal model to study vascular development in the direct vicinity of the implants.

**MATERIALS & METHODS**

**Materials**

Polyanionic salmon DNA (±300 bp/molecules; sodium salt) was kindly provided by Nichiro Corporation (Kawasaki, Japan). Potential protein impurities in the DNA were checked using the bicinchoninic acid (BCA) protein assay (Pierce, Rockford, IL) and were measured to be below 0.20% w/w (data not shown). Polycationic polyelectrolyte PDL (mw 30,000–70,000) was purchased from Sigma (Sigma-Aldrich Chemie B.V., Zwijndrecht, the Netherlands). All materials were used without further purification.

**Substrate preparation and cleaning**

Disc-shaped glass substrates (12 mm diameter; 1.5 mm thickness; Waldemar-Knittel Glasbearbeitung GmbH, Braunschweig, Germany) were cleaned in Piranha solution (hydrogen peroxide (H₂O₂, 30% aqueous solution)/sulfuric acid 3:7 v/v) to completely remove all traces of organic materials at the surface and to increase the wettability of the surface. Subsequently, the substrates were thoroughly rinsed with ultra-pure water from a Millipore system (resistivity ≥ 18.2 MW·cm; Millipore B.V., Amsterdam, The Netherlands) and dried using a pressurized air stream.

**Generation of multilayered DNA coatings**

DNA coatings were generated using layer-by-layer deposition, as described previously. Briefly, the substrates were immersed in an aqueous solution of PDL (0.1 mg/mL) for 30 min, allowing sufficient time for the adsorption of the first cationic polyelectrolyte layer onto the substrates. Subsequently, substrates were washed in ultra-pure water (5 min; continuous water flow) and dried using a pressurized air stream. Thereafter, substrates were alternately immersed in an anionic aqueous DNA solution (1 mg/mL) and the PDL solution for 7 min each, with intermediate washing in ultra-pure water and subsequent drying using a pressurized air stream to obtain final coating architecture of [PDL/DNA]_5. (The number indicates the number of double layers.)

**Experimental groups**

Non-coated substrates and [PDL/DNA]_5-coated substrates were sterilized using an ultraviolet treatment (254 nm; 4 h). Subsequently, VEGF functionalization of [PDL/DNA]_5-coated substrates was carried out as described below. The experimental groups used in this study were controls (non-coated glass substrates), [PDL/DNA]_5 (glass substrates with a [PDL/DNA]_5-coating), [PDL/DNA]_5-VEGF25 ( [PDL/DNA]_5-coated glass substrates with 25 ng VEGF), and [PDL/DNA]_5-VEGF250 ( [PDL/DNA]_5-coated glass substrates with 250 ng VEGF).
**Functionalization with VEGF**

Recombinant rat VEGF (rrVEGF164; 164 amino acids; Department of Biochemistry, Radboud University Nijmegen Medical Center, Nijmegen, The Netherlands) was serially diluted in phosphate-buffered saline (PBS), after which 25 or 250 ng of VEGF was applied in 10-μl aliquots to the sterilized [PDL/DNA]5-coated substrates. Because of the iso-electric point of VEGF (pI 8.5), it is positively charged at neutral pH and shows similar interaction with the negatively charged DNA top-layer in the multilayered structure as observed previously for BMP-2. The solvent was evaporated during incubation at 37°C.

**In vitro experiments**

**Cell culture and seeding.** Human umbilical vein endothelial cells (HUVECs; PromoCell GmbH, Heidelberg, Germany) were expanded in growth medium containing basic fibroblast growth factor (1 ng/mL), endothelial cell growth supplement/heparin (0.4% v/v), epidermal growth factor (0.1 ng/mL), hydrocortisone (1 μg/mL), phenol red (0.62 ng/mL), and fetal calf serum (FCS; 2% v/v) (PromoCell GmbH). To increase cell attachment, before cell seeding the experimental substrates were incubated for 1 h at 37°C. To increase cell attachment, before cell seeding the experimental substrates were incubated for 1 h at 37°C in assay medium (AM; alpha-minimal essential medium; Gibco, Invitrogen Corporation, Breda, The Netherlands), supplemented with 10% v/v FCS and gentamycin (50 μg/mL). For each experimental run, cells were seeded at a density of 20 x 10^3 cells/cm² and maintained in AM to exclude potential effects of factors supplemented in growth medium. Medium was refreshed at 1 day after cell seeding and thereafter 3 times per week.

**Proliferation of endothelial cells.** At selected times after cell seeding (1, 4, 7, and 10 days post-seeding), cells were trypsinized and counted using a Coulter counter (Beckman Coulter Inc., Fullerton, CA, USA). Data were obtained from 2 independent experimental runs with triplicate samples for each condition and time point (n = 3).

**Morphology of endothelial cells.** Parallel to the proliferation assays, 2 independent runs to study endothelial cell morphology were conducted. In each of these runs, 2 experimental samples for each condition at each time point were seeded with HUVECs at a density of 20 x 10^3 cells/cm² and incubated for 3 and 10 days. After these culture periods, cell layers were washed twice with PBS, fixed with 2% glutaraldehyde, and dehydrated in a graded series of ethanol. Finally, cell layers were dried using tetramethylsilane, sputter-coated with gold, and morphologically examined using a scanning electron microscope (JEOL 6310, JEOL Europe BV, Nieuw-Vennep, The Netherlands).

**Migration of endothelial cells.** The migrational behavior of endothelial cells (HUVECs) was assessed in 1 experimental run using a commercially available in vitro endothelial cell migration assay platform (BD BioCoat Angiogenesis System: Endothelial Cell Migration; BD Biosciences, Bedford MA), according to the manufacturer’s instructions. The assay measures the migration of cells (located in an insert) through a microporous membrane toward medium (with or without supplement) in the well. Briefly, [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 experimental substrates (n = 3) were placed in a 24-well plate and incubated with 1 mL AM for 1, 4, 7, 10, and 14 days. At each time point, the medium was collected, after which fresh medium was added to each experimental substrate for subsequent VEGF release. Collected medium was stored at −80°C until use in the migration kit. For the assay, 225 μL of collected medium was added in duplicate in a 96-well plate. Subsequently, HUVECs were added to each insert (25 x 10^3 cells in 75 μL of AM 2% bovine serum albumin (BSA)), whose content was separated from the well volume using a fluorescence-blocking microporous (3 μm) poly(ethylene terephthalate) (PET) membrane. The migration of HUVECs was quantified using post-labeling of the cells with calcine-AM and subsequent fluorescence measurements (excitation 485 nm, emission 530 nm) of the PET membrane after 22 h of incubation. For comparison, the migration of endothelial cells toward medium with known amounts of VEGF (standards: range 0–250 ng/mL) was measured. Data were corrected for fluorescence values of blanks (inserts without endothelial cells) and expressed as fold migration, which is set at 1 for the lowest VEGF concentration of the standard.

**In vivo experiment**

**Animals and implantation procedure.** For implantation, 200- to 250-g male Wistar rats were used. The animals were housed in the central animal facility at the Radboud University Nijmegen Medical Center (Nijmegen, The Netherlands), observing all national guidelines for care and use of laboratory animals. The animals received chow and water ad libitum. Before the implantation procedure, approval was obtained from the ethical committee for animal experiments at the Radboud University Nijmegen (KUNDEC #2003-14).

Surgery was performed under general inhalation anesthesia with nitrous oxide, oxygen, and isoflurane. The animals were immobilized and placed in a ventral position. Subsequently, the dorsal skin of the animals was shaved, washed, and disinfected with povidone-iodine. On each side of the vertebral column, 2 paravertebral incisions were made (~10 mm), after which a subcutaneous pocket was created using blunt dissection with scissors lateral to each incision. The insertion of substrates was performed such that the VEGF-functionalized side faced the dorsal skin. After insertion of an implant, the skin was closed using skin staples (Agraves; InstruVet C.V., Cuijk, The Netherlands). A total of 64 implants (16 implants of each group) were distributed in 16 rats (4 implants per rat) according to a Latin square randomization scheme.
Implant retrieval and histological preparations. At the end of each implantation period (1 or 3 weeks), 8 rats were killed using CO2-suffocation. Subsequently, tissue-covered specimens (n = 8 for each implant type and implantation period) were collected, the implants excised from the tissue capsule and checked for potential tissue remains using scanning electron microscopy as described previously, and the implant-surrounding tissue processed for embedding in paraffin. Paraffin sections were analyzed histologically (hematoxylin/eosin staining) to obtain complete descriptions of the observed thin sections.

Immunohistochemistry. In parallel with the hematoxylin/eosin staining, paraffin sections were used for immunohistochemical staining using an anti-alpha smooth muscle actin (anti-αSMA) antibody. Among other tissues, α-SMA is present in blood vessel walls and myofibroblasts, and its expression was used to determine the total number of blood vessels in the peri-implant area. Deparaffinated sections were treated with 3% H2O2 in PBS for 20 min to block endogenous peroxidase, fixed with 4% formalin, blocked with 5% BSA (Sigma), incubated with monoclonal immunoglobulin G2a mouse anti-αSMA (1:1600; Sigma), and incubated with biotinylated donkey anti-mouse secondary antibody (1:500; Jackson Laboratories, West Grove, PA). Staining was performed with a biotin-streptavidin detection system (Vectra Elite kit; Vector Laboratories, Burlingame, CA) for 45 min. Positive controls (rat brain tissue sections with positive staining in previous immunohistochemical procedures) and negative controls (1% BSA in PBS) were included.

Histological and histomorphometrical evaluation. A light microscope (Leica Microsystems AG, Wetzlar, Germany) was used for histological evaluation of the hematoxylin/eosin-stained sections. Immunohistochemically (anti-αSMA) stained sections were used for histomorphometrical evaluation of the number of blood vessels per area. Computer-based image analysis techniques (Leica Qwin Pro-image analysis software; Leica, Wetzlar, Germany) were used to determine the following parameters in digital images of immunohistochemically stained sections:

- region of interest (ROI): the connective tissue area at the dorsal site of the implant (excluding muscle and fat tissue) in a digital image of approximately $700 \times 560 \mu m$ (original magnification $20\times$; Fig. 1).
- number of blood vessels (BV): number of positively stained vascular structures (blood vessels) within the ROI.

The values obtained with these parameters were used to determine the vascularization (vessel density, VD):

$$VD = \frac{BV}{ROI}$$

For each implant-surrounding tissue section, VD was determined by averaging the numbers of counted vascular structures per ROI of 3 digital images (at $20\times$ magnification) per tissue section. Eight average VDs were obtained for each group at each time period.

Statistical analysis

All measurements were statistically evaluated using GraphPad Instat software (version 3.05; GraphPad Instat, San Diego, CA). Statistical analysis was performed using a one-way analysis of variance with a post hoc Tukey-Kramer multiple comparison test for cell proliferation, migration, and vascularization at individual implantation periods. Vascularization for individual groups and pooled groups at different implantation periods was assessed using...
Proliferation of endothelial cells. The 2 independent runs of endothelial cell proliferation showed comparable results. The results of one of these runs are presented in Figure 2. Initial cell numbers (on day 1) showed that comparable numbers of endothelial cells attached to the experimental substrates after cell seeding. From day 4, 7, and 10, however, endothelial cells on [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 showed statistically significantly more cells than controls and [PDL/DNA]5 (p < 0.05, p < 0.01, and p < 0.001 for days 4, 7, and 10, respectively). Whereas cell numbers on [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 increased over time, those on controls and [PDL/DNA]5 were constant over time and did not differ statistically from each other (p > 0.05). Furthermore, no statistically significant differences were found between [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 at individual time points (p > 0.05).

Morphology of endothelial cells. The morphological appearance of the endothelial cells (HUVECs) on the experimental substrates was evaluated using scanning electron microscopy. In Figure 3, representative images of endothelial cells cultured for 4 days on control and [PDL/DNA]5-VEGF250 substrates are depicted. Endothelial cells showed a similar morphology on all types of experimental substrates. The cells exhibited a flat appearance and had several cellular extensions. After 4 and 10 days of cell culture, a visibly greater number of cells was observed on [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 than on controls and [PDL/DNA]5.

Migration of endothelial cells. An in vitro migration assay was used to study the effect of VEGF release from VEGF-functionalized DNA-based coatings on endothelial cell migration. The migration of endothelial cells toward a concentration gradient of VEGF standards (rVEGF164 in AM; range 0–250 ng/mL) is depicted in Figure 4 (inset) and was used to obtain information on the VEGF release from VEGF-functionalized DNA-based coatings. The results show that VEGF significantly increased endothelial cell migration at concentrations ranging from 1.25 to 100 ng/mL (p < 0.05). In contrast, a VEGF concentration of 250 ng/mL had no effect on endothelial cell migration.

Collected medium from the incubation of [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 experimental substrates showed that these media also increased endothelial cell migration (Fig. 4). The increased effect on endothelial cell migration was observed in medium that had been collected from 1 to at least 14 days after initiation of incubation. No statistically significant differences were observed between media collected from [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250 experimental substrates (p > 0.05).

In vivo experiment

General observations. All 16 rats in the animal experiment remained in good health during the implantation periods.
and did not show any wound complications after surgery. At 1 week after implantation, a total of 32 implants were retrieved. At retrieval, a thin fibrous capsule surrounded all subcutaneous implants. None of the implant sites showed macroscopic signs of inflammation or adverse tissue reaction. Similarly, at 3 weeks after implantation, another 32 implants were retrieved, all of which were surrounded by a thin fibrous capsule and devoid of any macroscopic signs of inflammation or adverse tissue reactions.

**Descriptive histological evaluation.** Scanning electron microscopic examination of the excised implants showed that implants were sparsely covered with proteinaceous-like deposits, which were not identifiable as cellular material (data not shown). Gross evaluation of the hematoxylin/eosin-stained sections of the implant-surrounding tissue capsules after 1 week of subcutaneous implantation revealed a relatively uniform tissue response to all implants (Fig. 5). Tissue at the dorsal side of the implants consisted of a fibrous capsule, connective tissue, muscle tissue, and the dermis. Occasionally, fat tissue was observed in the vicinity of the implants. The fibrous capsule was immature, reflected by the absence of flat fibrocytes at the implant side of the capsule. Few or no inflammatory cells could be observed at the implant–capsule interface or in the direct vicinity of the implants.

After 3 weeks of subcutaneous implantation, a tissue response similar to that after 1 week was observed for all implants. However, the fibrous tissue capsules had matured, as indicated by the presence of fibrocytes with a flat appearance within the tissue capsules (Fig. 6A, B). The thickness of the fibrous capsules was within the range of 5 to 30 fibroblasts, independent of implant type. No gross differences in tissue response were observed, depending on implant type.

**Histomorphometrical evaluation of vascularization at the dorsal site of the implant.** For the evaluation of the vascularization of the connective tissue at the dorsal site of the implants, paraffin sections were subjected to immunohistochemical techniques using a monoclonal antibody against αSMA. Representative images of the stained sections are presented in Figure 6C and D. Positive staining was observed for vascular structures and fibrous capsules. A difference was observed in immunohistochemical staining of fibrous capsules after 1 and 3 weeks of implantation, with greater staining in the more-mature fibrous capsules at 3 weeks after implantation.

Histomorphometrically, the vascularity in the tissue at the dorsal site of the implant was calculated by determining the ROI and the number of positively stained vascular structures (blood vessels) within this ROI. The histomorphometrical results are presented in Table 1. The average ROI of the digitally imaged area in all experimental groups ranged from 0.25 to 0.28 mm². Vascularization (VD; # vascular structures/mm² ROI) of the analyzed areas showed a tendency to increase in both groups of VEGF-functionalized DNA-based coatings. However, no statistically significant differences between the individual experimental groups at individual implantation periods were found (p > 0.05). On the other hand, pooled data of VEGF-functionalized experimental implants ([PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250) and non-functionalized implants (controls + [PDL/DNA]5) showed significantly greater vascularization in the peri-implant tissue of VEGF-functionalized implants (p < 0.05). Vascularization in the peri-implant tissue was in the range of 102 to 155 and 66 to 142 vascular structures per mm² at 1 and 3 weeks after subcutaneous implantation, respectively. No statistically significant differences were found for individual experimental groups at different implantation periods, except for [PDL/DNA]5, which showed less vascularization after 3 weeks than after 1 week.

**DISCUSSION**

This study was initiated to study the effect of functionalized experimental substrates (poly-D-lysine/deoxyribonucleic acid/ [PDL/DNA]5-VEGF25 and [PDL/DNA]5-VEGF250). Inset shows migration effects of different concentrations of VEGF (rVEGF164; range 0–250 ng/mL; *p < 0.05 compared with 0 ng/mL).
proliferation and migration than of non-coated controls and non-functionalized DNA-base coatings. Furthermore, greater vascularization of the peri-implant tissue surrounding VEGF-functionalized DNA-based coatings was observed in vivo.

In an attempt to optimize the functionality of biosensors, this study was a first exploration of the use of DNA-based coatings for this purpose in view of the feasibility of their being functionalized with desired factors. Consequently, this study focused on functionalization of DNA-based coatings with the pro-angiogenic factor VEGF, and experiments were aimed at elucidating in vitro effects on endothelial cells and in vivo effects on implant site vascularization. The type of DNA-based coating was limited to [PDL/DNA]5, although the fabrication and characterization of different types of DNA-based coatings have been described previously. The choice for a single type of DNA-based coatings is justified, because previous experiments have demonstrated that cellular and histological responses to different types of DNA-based coatings were similar. The amount of VEGF needed for biological activity in vitro and in vivo appears to be unclear. Amounts as small as 10 ng have shown biological effects in vitro, whereas others used micrograms to induce biological effects of VEGF. It is likely that the amount needed for biological effects on vascular development are dependent on factors such as type of VEGF, release profile, and animal model. For our experiments, as a first approach, 2 amounts (25 and 250 ng) were chosen with a 10-fold difference.

The greater proliferation of endothelial cells on VEGF-functionalized DNA-based coatings in the in vitro experiments demonstrates that the pro-angiogenic factor VEGF remains biologically active. Similar results were recently found for the osteo-inductive factor BMP-2, which was superficially loaded onto DNA-based coatings and accelerated the differentiation of osteoblast-like cells. Furthermore, our proliferation experiments demonstrated that the presence of VEGF is a prerequisite for endothelial cells (HUVECs) to proliferate, because cells cultured on controls and DNA-based coatings did not proliferate. Additionally, no negative effects of DNA-based coatings on endothelial cells were observed, which corroborates observations of others using polyelectrolyte multilayers as a substrate for endothelial cells.

The efficacy of biomaterial devices in combination with growth factors depends partly on the availability of the growth factors for sensible cells. For instance, a minimal release of BMP-2 from a porous calcium phosphate scaffold appeared to be sufficient for enhanced bone formation. However, for our purpose, an increasing effect of VEGF on

FIG. 5. Representative histological sections (hematoxylin/eosin staining) of implant-surrounding tissue 1 week after subcutaneous implantation in the back of rats. (A, B) Overview showing the fibrous tissue capsule with connective tissue, a muscle layer, and dermis at the dorsal side of the implant. (C, D) Magnification of A and B showing the thin fibrous capsule and connective tissue at the dorsal side of the implant. Color images available online at www.liebertpub.com/ten.
vascularity in the peri-implant area in our in vivo experiment was likely to be dependent on release of the factor from the DNA-based coating. Additionally, the increased effect on the proliferation of endothelial cells observed on VEGF-functionalized DNA-based coatings does not necessitate factor release, because bound factor might also be available for adherent cells. In light of this, the results of the migration assay indicate a continuous release of VEGF from VEGF-functionalized DNA-based coatings. Although this assay does not provide detailed information on exact amounts of VEGF released at each incubation period, it shows that released amounts are sufficient for the chemotaxis of endothelial cells. On the basis of the release profiles of BMP-2 from [PDL/DNA]5 coatings and the comparable iso-electric point and molecular weight of BMP-2 and VEGF,13 it can be speculated that VEGF release profiles also show an initial burst of release.

![Histological sections of implant-surrounding tissue 3 weeks after subcutaneous implantation.](image)

**FIG. 6.** Representative histological sections of implant-surrounding tissue 3 weeks after subcutaneous implantation in the back of rats. (A, B) Hematoxylin/eosin staining showing the fibrous tissue capsule with connective tissue at the dorsal side of the implant. (C, D) Immunohistochemical staining (anti-alpha smooth muscle actin) showing the blood vessels in the connective tissue at the dorsal side of the implant. Color images available online at www.liebertpub.com/ten.

<table>
<thead>
<tr>
<th>Implantation time (weeks)</th>
<th>Vascularization (mean # vascular structures/mm² ± standard deviation)</th>
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</thead>
<tbody>
<tr>
<td>Control</td>
<td>111 ± 55</td>
</tr>
<tr>
<td>[PDL/DNA]5</td>
<td>111 ± 39</td>
</tr>
<tr>
<td>[PDL/DNA]5-VEGF25</td>
<td>155 ± 46</td>
</tr>
<tr>
<td>[PDL/DNA]5-VEGF250</td>
<td>102 ± 40</td>
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<td>[PDL/DNA]5-VEGF25</td>
<td>142 ± 54</td>
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<tr>
<td>[PDL/DNA]5-VEGF250</td>
<td>136 ± 65</td>
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*p < 0.05.

PDL, poly-D-lysine; DNA, deoxyribonucleic acid; VEGF, vascular endothelial growth factor.
release followed by a sustained release for several weeks. This would indicate that the initial VEGF burst release would approximate 17 and 170 ng for DNA-based coatings functionalized with 25 and 250 ng VEGF, respectively. The subsequent sustained VEGF release would then approximate 0.5 and 5 ng per week, respectively. Although the exact released amounts are not clear, the amounts released in the collected release media are apparently sufficient for an effect on endothelial cell migration.

Several animal models for studying angiogenesis have been described in detail, including dorsal skinfold chambers and ear pinnae; however, these models have limitations regarding implant size, which impedes their use for our implants. Additionally, because the focus of our study was to monitor vascularity effects of the peri-implant tissue of subcutaneous implants, we used a simple subcutaneous implantation model in rats. Inevitably, this simple approach for the detection of effects on vascularization has several drawbacks. For instance, the confinement of the area for evaluation of the effects of VEGF on vascularity was difficult to standardize. In contrast to previous studies, in which vascularity was determined in porous implants, the use of solid implants is associated with concessions in peri-implant area. The excision of the implants, allowing deformations of the fibrous capsule and surrounding tissue during subsequent histological processing further hampered standardization of the peri-implant area. Nevertheless, the approach used in this study allows the comparison of vessel density in the peri-implant area, in which the maximum distance of vessels to the implant surface approximates 600 μm (image height of zSMA-stained sections ≈ 650 μm). The results show no significant effects of individual VEGF-functionalized DNA-based coatings on peri-implant vessel density. However, significantly greater peri-implant vascularity was observed in VEGF-functionalized DNA-based coatings than in controls and non-functionalized DNA-based coatings. Furthermore, a significant decrease in vascularity from 1 to 3 weeks after implantation was observed in tissue surrounding non-functionalized DNA-based coatings. This observation might be related to the effect of DNA-based coatings on fibroblasts, which form the fibrous capsule, and subsequent intercellular signaling between fibroblasts and cells in their surroundings. However, the actual cause for this observation remains unclear and should be addressed in future experiments.

Although the functionalization of DNA-based coatings in this study was restricted to a single angiogenic factor, considerable debate is ongoing on the necessity for combined growth factor delivery for angiogenesis and vessel maturation. Indeed, synergistic effects of combined administration of VEGF and basic fibroblast growth factor have been demonstrated on collateral circulation in a hind limb ischemia model. Additionally, Peatitie et al. demonstrated enhancement of angiogenic responses in a murine ear pinnae model using dual delivery of VEGF and keratinocyte growth factor. Such combined delivery vehicles could also be fabricated using DNA-based coatings, because the mechanism of fabrication has been demonstrated to be applicable for compartmentalization of such polyelectrolyte multilayers, which would allow temporal control over growth factor release.

**CONCLUSION**

This study demonstrates that the functionalization of DNA-based coatings with the pro-angiogenic factor VEGF results in greater proliferation and migration of endothelial cells in vitro, indicating that the VEGF retained its biological activity. In an in vivo implantation experiment in rats, greater vascularity of the peri-implant tissue surrounding VEGF-functionalized DNA-based coatings was observed.

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