Bioengineering a pre-vascularized pouch for subsequent islet transplantation using VEGF-loaded polylactide capsules†

Naresh Kasoju, †a Alžběta Pátíková,b Edyta Wawrzynska,a Alžběta Vojtíšková,b Tomáš Sedačík,b †a Marta Kumorek,a Ognen Pop-Georgievski,a †a Eva Sticová,c Jan Kříž*b and Dana Kubies †a*†

The effectiveness of cell transplantation can be improved by optimization of the transplantation site. For some types of cells that form highly oxygen-demanding tissue, e.g., pancreatic islets, a successful engraftment depends on immediate and sufficient blood supply. This critical point can be avoided when cells are transplanted into a bioengineered pre-vascularized cavity which can be formed using a polymer scaffold. In our study, we tested surface-modified poly(lactide-co-caprolactone) (PLCL) capsular scaffolds containing the pro-angiogenic factor VEGF. After each modification step (i.e., amination and heparinization), the surface properties and morphology of scaffolds were characterized by ATR-FTIR and XPS spectroscopy, and by SEM and AFM. All modifications preserved the gross capsule morphology and maintained the open pore structure. Optimized aminolysis conditions decreased the $M_w$ of PLCL only up to 10% while generating a sufficient number of NH$_2$ groups required for the covalent immobilization of heparin. The heparin layer served as a VEGF reservoir with an in vitro VEGF release for at least four weeks. In vivo studies revealed that to obtain highly vascularized PLCL capsules (a) the optimal VEGF dose for the capsule was 50 µg and (b) the implantation time was four weeks when implanted into the greater omentum of Lewis rats; dense fibrous tissue accompanied by vessels completely infiltrated the scaffold and created sparse granulation tissue within the internal cavity of the capsule. The prepared pre-vascularized pouch enabled the islet graft survival and functioning for at least 50 days after islet transplantation. The proposed construct can be used to create a reliable pre-vascularized pouch for cell transplantation.

1. Introduction

The replacement of failed cells by transplantation is one of the most promising approaches that have been already tested and applied in clinical practice. Over the last several decades the transplantations of parathyroid glands,۱ thyroid gland,۲ adrenal cells,۳ thymus۴ and pancreatic islets۵ have been introduced and tested under experimental and clinical conditions. In addition, cellular transplantation can be theoretically used for the therapy of haemophilia, liver failure, Parkinson’s disease, transversal spinal cord injuries and others.۶—۹ Outcomes of these procedures still vary widely but it seems likely that the procedure’s effectiveness can be considerably improved via optimization of the transplantation site and the procedure itself. The generally accepted requirement for an adequate environment for an engraftment of transplanted cells inspires many groups to pay great attention to the characterization and subsequent development of the optimal transplantation site. Some approaches focused on cell protection against inflammation and more specific rejection using isolation of the graft in microcapsules.۱۰ This method can effectively serve in the case of cells releasing hormones slowly and evenly with very small variations in time. Another approach focused on the creation of microencapsulation using a cell pouch with a fully finished healing in order to protect the cells from inflammation and to concentrate them into a small volume thus allowing the subsequent excision and a detailed examination.۱۱,۱۲ The finished healing provides a hospitable environment but suffers from poor blood perfusion.۱۳ This feature would be detrimental for the graft survival particularly in the case of cells serving...
as a sensor of the metabolic state in addition to hormone secretion. For example, pancreatic islets are extremely oxygen demanding and even a short time disruption of blood perfusion in combination with a high concentration of glucose can cause their destruction. This crucial limitation known for successful islet transplantation can be improved when the pouch providing an adequate blood perfusion in the time of transplantation is created in advance.

Fig. 1 Schematic of the implantation, capsule preparation and functionalization: (a) a schematic of the proposed approach to bioengineer a pre-vascularized chamber for potential cell transplantation using a PLCL capsular scaffold with a Teflon lid. (b) The capsule preparation was performed following a TIPS principle by using a custom-made setup having a reservoir for a dry ice–ethanol mixture (R) that was connected to a conducting block (C) and a template bar (T) (for details see ref. 13). (c) The capsule functionalization was performed in three stages including aminolysis, heparin crosslinking by carbodiimide chemistry and then VEGF loading by physical adsorption.
Several device-based approaches, including those based on a groin flow-through chamber model,14,15 have been described to create a transplantation site which provides sufficient blood supply for transplanted islets. Although these approaches may be desirable because they allow immediate access to the existing vasculature, none was transferred to clinical practice likely due to the extent of surgical manipulation needed in order to accommodate a sufficient number of islets. Alternatively, devices based on polymeric porous scaffolds are found to be simple in terms of surgical manipulation, offer an adequate volume for islet loading, provide mechanical graft protection and may be easily retrievable in the case of graft infection.16,17

In our study we have focused on the creation of a pre-vascularized pouch with an adequate inner volume, without considerable inflammation in the time of transplantation and rich in a vascular/capillary network on the internal surface. We believe that such induced pre-vascularization could significantly shorten the “hypoxia” time when islets are dependent on oxygen diffusion. A quick islet reconnection to the host vascular network, resulting in a rapid re-perfusion, improves the subsequent islet graft survival and its function. The concept is presented in Fig. 1a. Based on previously published experiments where the number of transplanted islets was considerably higher than the number used in clinical programs,16,17 we have focused on the preparation of a microporous device, which can direct fibrous tissue penetration and gradually release vascular growth factors stimulating ingrowth of vessels together with connective tissue. However, with this type of device, the scaffold porosity is a key parameter for an optimal recipient tissue response and, subsequently, the transplantation outcome. We have previously shown that in comparison with a typical scaffold usually with an isotropic pore morphology, a scaffold consisting of anisotropic channelled pores showed the desired guided cell infiltration.20,21 The proposed capsule-shaped scaffolds have been fabricated from an FDA approved polylactide-based polymer; the scaffolds possess channelled pores oriented into an internal capsule lumen that could serve as a cavity for subsequent cellular, or specifically islet, transplantation (Fig. 1a and 2).

However, a major drawback of the use of synthetic polymers is the fact that they do not usually provide biochemical cues, e.g., molecules acting as proangiogenic signals that are highly required to trigger an efficient vascularization process in tissue engineering applications.22 Although the natural healing process attempts to vascularize a polymer scaffold, our earlier attempts showed that this process was relatively slow and that the resulting tissue was often mostly fibrotic.20,21 To this end, here we undertook functionalization of a poly(lactide-co-caprolactone) (PLCL, \( M_w = 316 \,000 \, g \, mol^{-1} \) and \( M_n = 120 \,000 \, g \, mol^{-1} \), 14 wt% of \( \varepsilon \)-CL) was synthesized according to the protocol described by Kubies et al.24

2. Materials and methods

All materials, chemicals, and reagents were used as received unless stated otherwise. The detailed synthetic procedures, characterization methods, and implantation protocols are provided in the ESL† In the following, a short summary of the sample preparation, the characterization and implantation methods is presented.

2.1 Materials

High molecular weight poly(\( \varepsilon \)-lactide-co-\( \varepsilon \)-caprolactone) (PLCL, \( M_w = 316 \,000 \, g \, mol^{-1} \) and \( M_n = 120 \,000 \, g \, mol^{-1} \), 14 wt% of \( \varepsilon \)-CL) was synthesized according to the protocol described by Kubies et al.24

2.2 Fabrication of samples

Macroporous 3D samples: PLCL capsule-shaped scaffolds were prepared using a thermally induced phase separation (TIPS) approach according to the protocol reported earlier by our group.20,21,23 The protocol is schematically depicted in Fig. 1b. Briefly, the metal template connected to the cooling platform was dipped into a PLCL solution (5% w/v in 1,4-dioxane). The system was cooled to \(-80\, ^\circ\text{C}\) and the phase separation proceeded for 30 s followed by coarsening for 30 s. Then the template with the sample was dipped into 1,4-dioxane for another 30 s to avoid any skin effect, left to coarsen for 30 s and subjected to freeze-drying.

Model 2D samples: to properly analyse the surface properties such as topography and the surface elemental composition of PLCL surfaces before and after functionalization, model PLCL films were used instead of the porous scaffolds. The model PLCL films were prepared by the solvent casting procedure from a 10% PLCL solution (w/v in 1,4-dioxane).

2.3 Functionalization of samples

2.3.1 Introduction of amine groups. The optimization study was performed using the capsular PLCL scaffolds or flat model films. Briefly, the samples were incubated in a hexa-
methylenediamine (HMDA) solution in isopropanol (5% w/v) for various incubation times (5, 15, 30 and 60 min) at two different temperatures (22 and 40 °C).

2.3.2 Immobilization of heparin. The optimization study was performed using the capsular PLCL scaffolds or flat PLCL films, both aminolysed with HMDA (5% w/v in isopropanol) at 22 °C for 30 min. Briefly, heparin was immobilized covalently using EDC/NHS chemistry for 1, 6 or 15 h. As a control, physical adsorption was performed using a 0.25% w/v heparin solution in 0.1 M MES buffer (pH 5.5) for 15 h.

2.3.3 VEGF loading and in vitro release. The screening study evaluating VEGF loading and release was performed using the capsular PLCL scaffolds (an average weight of 25 ± 1.5 mg) that had been aminolysed with HMDA at 22 °C for 30 min and contained a covalently immobilized heparin layer (0.25% w/v in MES, using EDC-NHS, 6 h).

VEGF loading was performed from a VEGF solution in PBS (pH 7.4) supplemented with 0.1% w/v BSA. The model tested concentrations were 0.08 µg VEGF per mg of scaffold (i.e., 2 µg VEGF/scaffold, the sample denoted as heparin/VEGF_2) and 0.4 µg VEGF per mg of scaffold (i.e., 10 µg VEGF/scaffold, the sample denoted as heparin/VEGF_10). The release of VEGF from the scaffolds into PBS supplemented with BSA at 37 °C within four weeks was investigated by the full immersion method.

2.4 Characterization studies

2.4.1 Common methods. The surface morphology was assessed by scanning electron microscopy (SEM, Vega, Tescan).

The surface topography was assessed using an atomic force microscope in tapping mode (AFM, Nanoscope IIIa, Digital Instruments).

Surface wettability (CA) was determined by the sessile drop method using a goniometer system (OCA 20, Dataphysics).

Surface chemistry was analysed by X-ray photoelectron spectroscopy (XPS) by Fourier transform infrared spectroscopy in attenuated total reflectance mode (ATR-FTIR).

For SEM, ATR-FTIR and CA analyses the porous scaffolds were used, while for AFM and XPS analyses, the model flat films were used.

2.4.2 Specific characterization of aminolysis. The presence of the introduced amine groups was qualitatively verified using confocal laser scanning microscopy. The loss in physical mass from the scaffolds and the changes in the molecular weight of PLCL after aminolysis treatment were evaluated gravimetrically and by gel permeation chromatography.

2.4.3 Specific characterization of heparinization. The efficiency of heparinization was assessed by toluidine blue assay. For qualitative assessment, the scaffolds after the assay were visualized using a stereo zoom light microscope (Olympus Stereo Zoom Microscope SZ61). For quantitative assessment, absorbance at 631 nm of a 200 µl aliquot of the toluidine blue-water layer was determined.

2.4.4 Specific characterization of VEGF loading and release. Qualitative assessment of VEGF loading onto the scaffolds was performed by immunohistochemical (IHC) staining with anti-VEGF antibody followed by imaging using light microscopy. Quantitative determination of the amount of VEGF released from the scaffolds was assessed by an ELISA method using a commercial kit (R&D Systems, USA).

To deduce the mode of VEGF release from the scaffolds, the release kinetics data were fitted to four mathematical models, viz., zero-order, first-order, Higuchi and power law models.25–28

2.5 In vivo studies

2.5.1 Scaffold preparation for implantation. All the capsules were subjected to sterilization by treating with 70% ethanol for 1 h. All subsequent modifications, i.e., aminolysis, heparin functionalization and VEGF loading, were performed under sterile conditions using sterile solutions/buffers. The capsular PLCL scaffolds (2 cm-long, with an average weight of 25 ± 1.5 mg) for in vivo evaluation were aminolysed at 22 °C for 30 min (see paragraph 2.3.1). Then, coating with a crosslinked heparin layer was performed for 6 h (see paragraph 2.3.2 and the ESI†). Finally, the scaffolds were transferred to PBS and loaded with VEGF in PBS/0.1% w/v BSA (see paragraph 2.3.3) at doses 0.4, 2 or 4 µg per mg of the capsule, i.e., 10, 50 and 100 µg per the capsule, respectively. For better clarity, the samples were denoted as heparin/VEGF_10, heparin/VEGF_50, and heparin/VEGF_100. Heparinized scaffolds without VEGF loading were used as control samples.

2.5.2 Surgery and scaffold implantation. Experiments were conducted according to the guidelines for the Care and Use of Laboratory Animals of the Institute for Clinical and Experimental Medicine (ICEM), Prague, Czech Republic. The protocol was approved by the Animal Care Committee of the ICEM and the Ministry of Health, Czech Republic. The capsular PLCL scaffolds were implanted in the greater omentum of Lewis rats (230–270 g, age: 2–3 months, purchased from Charles River, Germany). The surgery was performed using a previously described technique.29

For the dose-optimization experiment, scaffolds supplemented with 10, 50, or 100 µg VEGF (the samples denoted as heparin/VEGF_10, heparin/VEGF_50 and heparin/VEGF_100) were implanted into the greater omentum of the experimental animals for 2 weeks. For time-optimization experiments, scaffolds loaded with 50 µg VEGF (i.e., heparin/VEGF_50) were implanted for 2, 3, or 4 weeks. After the selected time period, the abdominal cavity was again opened under general anaesthesia, and the scaffold with the surrounding tissues was excised and fixed; the animal was then euthanized.

2.5.3 Pancreatic islet isolation. Pancreatic islets were isolated from Lewis male rats (300–400 g) according to the standard protocol (see the ESI†). The isolated islets were cultured overnight at 37 °C and 5% CO2 in CMRL-1066 medium (Sigma-Aldrich, USA) supplemented with 5% HEPES buffer, 10% fetal bovine serum, and 1% penicillin/streptomycin.

2.5.4. Diabetes induction, islet transplantation and islet survival monitoring. Male Lewis rats (n = 8) were injected with
65 mg kg$^{-1}$ of streptozotocin (Sigma Aldrich, USA) intraperitoneally. Animals with confirmed diabetes mellitus were considered as recipients ($n = 6$) of pancreatic islets.

The scaffolds loaded with 50 µg VEGF (i.e., heparin/VEGF$_{50}$) were immersed in platelet enriched plasma and were implanted within the greater omentum of recipients (see paragraph 2.5.2. and the ESI†). In the meantime, insulin pellets were implanted subcutaneously in order to control the blood glucose levels prior to islet transplantation.

Four weeks after the scaffold implantation a cranial midline laparotomy was performed and the scaffold was placed on wet gauze. A small incision in the omentum was performed to allow a Teflon bar removal and islets were then injected into the created cavity; finally, the omental incision and abdominal wall were closed. Two weeks after islet transplantation, insulin pellets were removed.

The blood glucose level was monitored twice a week and body weight once a week for 50 days. Glycemia under 10 mmol l$^{-1}$ was considered as a sign of a functional graft, and glycemia over 11.1 measured at least three times was considered as a marker of graft failure. An intravenous glucose tolerance test (0.5 g of glucose per kg) was performed 40 days after islet transplantation. At the end of the experiment, each scaffold with the islet graft was harvested, fixed and provided for immunohistological analysis.

2.5.5 Histochemical and immunohistochemical analysis. Each scaffold was fixed after excision, and processed using standard techniques for histochemical and immunohistochemical staining. For evaluation of tissue ingrowth and creation of new capillaries within the scaffold wall, the sections were stained with haematoxylin and eosin to identify a basic structure of the tissue, with Masson’s trichrome to display extracellular matrix components, particularly collagen, and with anti-CD31 antibody specific for endothelial cells.

The islet structure and the presence of blood capillaries in the graft were evaluated using double-staining with anti-insulin and anti-glucagon antibodies and with anti-CD31 antibody specific for endothelial cells.

3. Results and discussion

3.1 Typical morphological features

The pore properties of a scaffold, both at the macro-scale and micro-scale, greatly influence the fate of a scaffold in cell and tissue engineering; therefore, the morphological characteristics of the prepared 3D microporous PLCL capsules were studied in detail. The capsule lumen has a diameter of 4 mm and a length of 20 mm (Fig. 2a) and can accommodate 250 µl of the cell suspension. The pore size of various sections of the capsule was analysed by SEM. A representative cross-section of the capsule (Fig. 2b) shows the channelled pores and the interconnected character of the pore network; the thickness of the capsule’s wall ranged between 600 and 650 µm. The outer surface exhibited pores 50–70 µm in diameter, and the inner surface revealed pores 5–10 µm in diameter (Fig. 2c and d).

The capsule possessed a unique radially oriented, channelled, interconnected and anisotropic pore morphology as a result of the Dip TIPS approach that we employed. The use of isotropic porous scaffolds often results in the formation of a necrotic core due to random cell infiltration and associated constraints on nutrient diffusion, whereas anisotropic porous scaffolds inherently promote guided cell infiltration without affecting nutrient diffusion and therefore yield fully functional viable tissue.

3.2 Aminolysis

Polylactide-based surfaces possess only a few carboxyl and hydroxyl groups on the chain ends resulting from the mechan-ism of lactide polymerization in the presence of tin(II) octoate. To introduce functional NH$_2$ groups for covalent immobilization of heparin on the complex surface of the 3D PLCL scaffold, the pristine capsules were subjected to aminolysis. Aminolysis is considered one of the most convenient procedures for surface modifications and can also be used in the surface hydrophilization of polymers. Since the reaction takes place only at the polymer surface and does not change the polymer’s bulk properties, this procedure is especially favourable for the modification of post-formatted scaffolds and membranes. Highly reactive amine groups break some of the ester groups of polyester chains and covalently bind to a single carboxyl group. Subsequently, the remaining amine end-group can be used for immobilization of various functional biomacromolecules. However, the ester bond cleavage results in a reduction of the molecular weight of the sample. Zhu and co-workers have extensively investigated the functionalization of poly(e-caprolactone) and poly(lactic acid) films with different diamines. The group studied the influence of the reaction time, temperature, solvent type, and the length and concentration of the diamine on the mechanical properties of aminolysed membranes. It was found that the reaction of 1,6-hexanediame (HMDA) proceeds very mildly.
We adjusted the aminolysis conditions to achieve minimal degradation of polymer chains and to preserve the morphology of the complex 3D PLCL capsules. The reaction with HMDA was conducted in isopropanol (a non-solvent for PLCL) to avoid any damage to the bulk of the capsule. First-hand visual confirmation of successful amination was obtained by FITC staining followed by CLSM imaging. In agreement with earlier reports, we found that pristine PLCL capsules showed no signal, whereas the aminated capsules showed intense green fluorescence after staining with FITC (Fig. 3A).

The chemical composition of pristine and aminated PLCL surfaces was studied by ATR-FTIR and XPS analyses. As presented in Fig. 3B, pristine PLCL showed characteristic peaks associated with ester linkages (C=O, C–O and long alkyl chain-related) at 1000–1300 cm⁻¹, carbonyl stretching (C=O related) at 1750 cm⁻¹ and CH₂ stretching (alkyl group-related) at 2945 and 2995 cm⁻¹. Yuan et al. reported that aminolysis of poly(ε-caprolactone) (PCL) surfaces resulted in additional, tiny peaks at 3321, 1650 and 1550 cm⁻¹; these peaks were attributed to the successful introduction of amine groups. However, in our experiments, the FTIR spectra of aminated PLCL did not exhibit detectable peaks of this type, possibly due to the low degree of amination, the limitations of the instrument, or both. The surface composition of the samples was further analysed by XPS (Fig. 3C). The high-resolution C 1s spectrum of pristine PLCL was characterized by aliphatic C–C and C–H contributions at 285.0 eV, C–O contribution at approximately 286.9 ± 0.2 eV and ester C(=O)
—O contribution at 289.0 ± 0.1 eV. Aminolysis of the PLCL capsules resulted in an increase in the contributions at approximately 286.3 ± 0.2 eV due to the additional presence of C–N moieties that overlap with the initially observed C–O contributions. Consistent with the observations from the C 1s spectrum (ESI Fig. 1†), the high-resolution N 1s spectrum of the aminolysed PLCL capsules was characterized by a dominant amide and amine contribution at 400.0 ± 0.1 eV and a minor charged amide contribution at 401.9 ± 0.2 eV. The obtained results are in agreement with the results reported by Zhu et al.35 The nitrogen content of the PLCL surfaces reached 0.5 atomic % after 60 min of aminolysis irrespective of the reaction temperature. This observation is consistent with previous work on the surface composition of aminolysed PLA and PCL surfaces, in which a nitrogen content in the range of 0.5–1.3 atomic% was reported.33,35,41 The surface coverage with amine active species obtained herein should be sufficient for the purpose of further functionalization.33 This assumption was confirmed in subsequent heparin immobilization experiments.

Proof of successful amination was also sought from contact angle measurements based on the fact that the incorporation of amine groups decreases the hydrophobicity of the capsule surface. As shown in Fig. 3D, the CA of pristine PLCL was 144 ± 4°, thus representing a highly hydrophobic surface. The observed high CA value was not typical of PLA-based flat surfaces and resulted from the porous open structure of the aminolysed capsule surface. This surface feature was described in more detail in our previous work.23 After aminolysis for 30 min, the CA of the PLCL 3D surfaces decreased significantly to 101 ± 4°. This significant reduction of 30% clearly demonstrates that effective surface hydrophilization occurred due to the presence of the introduced amine groups. Previously, several groups reported an overall reduction in contact angles between 15 and 26% for flat non-porous PCL surfaces aminolysed with an HMDA solution.34,35,43 On the other hand, Croll et al. reported a decrease of only 2% when PLGA surfaces were treated with 0.05 M N-(2-aminoethyl)-1,3-propanediamine13 and Sharkawi et al. observed no significant changes in the wettability of HMDA-aminolysed PDLA films with a thickness of 0.5 mm.42 During aminolysis, some of the ester bonds of the PLCL chains break apart, and HMDA forms amide bonds by nucleophilic substitution. In principle, aminolysis is destructive to the polymer chain; too intensive scission of the polymer chains will result in changes in their mechanical properties and thereby in changes in the integrity of the initial 3D scaffold structure that can affect the scaffold’s stability and thus its possible practical applications. Therefore, the changes in the material properties after the aminolysis are a matter of serious concern. We analysed the changes in the molecular weight (M_w) and the physical mass of the scaffolds as a function of the duration and temperature of aminolysis treatment (Fig. 4A). There was an increasing trend in the loss of physical mass as a function of time and temperature of aminolysis. Nevertheless, the mass loss was less than 3% under all tested conditions. In contrast, the changes in M_w were more prominent. At 22 °C, the M_w decreased from 6.5 to 20.4% as the duration of aminolysis increased from 5 to 60 min. A similar increasing trend was observed at 40 °C. In addition, the changes in M_w were temperature dependent; after 30 min of aminolysis, the decrease in M_w was 11.4% at 22 °C, whereas it was 28.8% at 40 °C. Only a few groups have investigated the effects of reaction conditions on changes in molecular characteristics. Effects similar to those observed in our study were observed by Zhu et al. during the aminolysis of PCL films in the presence of 5% HMDA at 30 °C for 30 min with a resulting M_w decrease of 11.6%,35 by Hong et al. when PLA microspheres underwent aminolysis in 6% HMDA/m-propanol solution at 60 °C for 60 min, resulting in a mass loss of 30%,43 and by Collins et al. in the case of poly(ethylene terephthalate) (PET) films aminolysed in the presence of 40% aqueous methyamine at 20 °C for 1 h, for which a M_w decrease of 26.8% was reported.44

To obtain information on the overall structure of the aminolysed capsules, we analysed the samples by SEM. Importantly, there were no visible defects in the gross morphology or in the character of the pore structure before or after aminolysis (ESI Fig. 2f and Fig. 4B). We also tracked the possible changes in the topography of the model solvent-cast PCL films by AFM (Fig. 5). Notably, the pristine and aminolysed PLCL surfaces exhibited very similar overall topographies with lamellar structures as a result of the crystallization of the PLCL chains during the film processing (Fig. 5A and B). The slight increase in the surface roughness, and subtle changes in the character of the lamellae, can be considered to be a result of the etching procedure. Similar effects of surface etching on the surface topography have been reported by Zhu et al.,34,35 and Yuan et al. for aminolysed PCL substrates and by Drobeta et al. for PET substrates.45

To summarize, the aminolysis performed at 22 °C for 30 min resulted in sufficient surface modification with an acceptable decrease in the molecular weight while preserving the gross morphology of the modified scaffolds. These conditions were selected for subsequent modification studies with heparin and for the preparation of the samples for in vivo studies.

3.3 Heparinization

To further functionalize the capsule with heparin, a biomacromolecule that has a strong affinity for VEGF and thereby increases its loading amount and structural stability, aminolysed PLCL capsules were incubated with a heparin solution in the presence or absence of crosslinking agents. During physical adsorption (PA), the negatively charged sulfate and carboxyl groups of heparin participate in electrostatic interactions with the positively charged amine groups of aminated PLCL. This surface modification was considered as a control, whereas, during the reaction with crosslinking agents (CL), heparin was first activated in the presence of EDC and NHS and then crosslinked to the amine groups of aminated PLCL via covalent amide bonds. In both cases, first-hand visual confirmation of
successful amination was obtained by toluidine blue staining. The aminated PLCL control samples remained white, whereas the heparinized PLCL test samples showed a distinct purple colour (Fig. 4C), in good agreement with the earlier reports.\textsuperscript{38} A high intensity of the purple colour in the chemically cross-linked heparinized PLCL sample demonstrated the effectiveness of the covalent heparin grafting in comparison with the physical adsorption method.
The chemical surface composition of the physically adsorbed (PA-Hep-PLCL) and chemically crosslinked (CL-Hep-PLCL) heparinized PLCL surfaces was analysed by ATR-FTIR and XPS. The ATR-FTIR spectra (Fig. 3B) of pure heparin showed heparin’s characteristic peaks for asymmetrical stretching of $\text{S} = \text{O}$ in $\text{CH}_2\text{SO}_3^-$ at 1010 cm$^{-1}$, symmetrical stretching of $\text{S} = \text{O}$ in $\text{NH}_2\text{SO}_3^-$ at 1220 cm$^{-1}$, N–H bending at 1420 cm$^{-1}$, C=O stretching at 1610 cm$^{-1}$ and OH stretching at 3400 cm$^{-1}$. Most of these peaks were found in both the PA-H-PLCL and CL-H-PLCL samples; however, in contrast to PA-H-PLCL, the chemically crosslinked CL-H-PLCL exhibited a new peak at 1640 cm$^{-1}$ that was characteristic of the amide bond created as a result of the covalent crosslinking between heparin and aminated PLCL. Our data are consistent with an earlier report by Singh et al., who observed a similar peak after chemical crosslinking of heparin and PCL. The chemical crosslinking of heparin and PCL.38

Further evidence supporting the successful heparinization of the PLCL capsules was obtained from XPS analysis. The physical adsorption of heparin on the Am-PLCL surfaces induced only minor changes in the C 1s spectrum. At the same time, the covalent binding of heparin caused an increase in the amide contribution at 287.6 ± 0.2 eV. Notably, both PA-Hep-PLCL and CL-Hep-PLCL (Fig. 3C) samples exhibited a new contribution in the high-resolution S 2p spectrum at 168.4 ± 0.2 eV (with a separation between the $2p_{3/2}$-$2p_{1/2}$ spin-split doublet of approximately 0.6 eV) that corresponds to the $\text{SO}_4\text{H}$ and $\text{NH}_2\text{SO}_3\text{H}$ moieties of heparin. Such a peak was also observed by Kolar et al. in heparin-functionalized PET tubes47 and by Sharkawi et al. in the heparin-functionalized PDLLA films.42

The successful heparinization of the PLCL capsules led to changes in their surface wettability. The heparinization resulted in surface hydrophilization. As shown in Fig. 3D, the contact angles of PA-Hep-PLCL and CL-Hep-PLCL were 81 ± 3 and 53 ± 5°, respectively; this corresponds to a reduction by 20 and 47%, respectively, compared to the aminated Am-PLCL (101°), and by 44 and 63%, respectively, compared to pristine PLCL (145°). Similar changes in surface wettability were reported by Khandwekar et al., who found a 43% reduction in the advancing contact angles of heparinized PCL surfaces compared to those of pristine PCL surfaces.46 The observed profound difference in the contact angles of the PA-Hep-PLCL and CL-Hep-PLCL surfaces also indirectly showed that the chemical crosslinking approach made it possible to immobilize a higher (and/or more stable) amount of heparin on the polymer surface than was accomplished via the physical adsorption method. The heparin content of the PLCL capsules quantified by the toluidine blue assay was 0.33, 1.08, 2.10 and 2.39 µg per mg of the scaffold in the 15 hPA-Hep-PLCL, 1 hCL-Hep-PLCL, 6 hCL-Hep-PLCL and 15 hCL-Hep-PLCL samples, respectively (Fig. 4C). Consistent with the qualitative results, the efficiency of heparin immobilization by physical adsorption was lower than that achieved by chemical crosslinking. Furthermore, the heparin content increased with crosslinking time. Our observation agrees with the findings of Wei et al., who reported a similar difference in the adsorbed vs. crosslinked heparin content of chitosan/graphene oxide hydrogels,49 and with those of Singh et al., who reported 1.5 and 7.5 µg heparin per PCL scaffold (discs of 6 × 2 mm) after a crosslinking time of 1 and 15 h, respectively. However, although the heparin content and crosslinking time appeared to be directly correlated, as shown in Fig. 5C, the relationship was not linear. The data suggest that the rate of heparin immobilization may be linear during the first few hours but that it may then reach a plateau due to a lack of sufficient binding sites. Since a detailed kinetic study was beyond the scope of the current investigation, based on our observations we used a 6 h crosslinking time as the optimum reaction time for all following experiments.

The changes in the surface topography of the scaffolds before and after heparinization were investigated by AFM using model solvent-cast films. Fig. 5 shows typical topographies and distinct differences between physically and covalently immobilized heparin layers. The topography of the physically adsorbed Hep surface closely follows the topography of the original aminolysed surface (Fig. 5C) reflecting the low amount of Hep adsorbed as evaluated by the toluidine blue assay (Fig. 4C). On the other hand, we observed spherically shaped objects (hills) in the samples that had reacted for 6 h to produce covalently immobilized Hep layers (Fig. 5D); in these samples, an amount of heparin greater than one order of magnitude was deposited. There was also a significant increase in the mean square roughness, $R_q$. The $R_q$ of Am-PLCL was 2.9 nm, whereas $R_q$ increased to 4.4 nm in the case of 15 hPA-Hep-PLCL and to 41.4 nm in the case of 6 hCL-Hep-PLCL.

On SEM analysis, the aminolysed capsule surface appeared smooth, whereas globular structures were clearly seen in the Hep-PLCL capsules crosslinked for 6 and 15 h (Fig. 6); in the case of CL-Hep-PLCL capsules, we can consider that a gradual increase in the granule size and granule surface density occurred with increasing crosslinking time. Elahi et al. recently reported similar morphological changes in silk fibroin fabric after deposition of polyelectrolyte multilayers (PEMs) followed by EDC/NHS-mediated heparin immobilization; the authors speculated that these changes reflect the non-uniform deposition and distribution of PEMs and heparin on the fabric surface.50,51 Based on our findings and on their similarity to the findings of Elahi et al., we conclude that the observed morphological changes were the result of EDC/NHS-mediated heparin immobilization.

### 3.4 VEGF immobilization

VEGF is a potent proangiogenic molecule that has been shown to promote infiltration by and proliferation of endothelial cells leading to efficient vascularization of scaffolds.52 Taking advantage of the strong bioaffinity between VEGF and heparin, we immobilized VEGF on the heparinized surfaces of PLCL capsules without using any crosslinkers, i.e., by physical adsorption. To study the capability of modification, VEGF was loaded at two model concentrations, i.e., 0.08 µg mg$^{-1}$ of scaffold (i.e., in total 2 µg per capsular scaffold;
heparin/VEGF_2) and 0.4 µg mg⁻¹ of scaffold (i.e., in total 10 µg per capsular scaffold; heparin/VEGF_10). The success of the VEGF loading was qualitatively determined by IHC staining using an antibody against VEGF. As presented in Fig. 7A, staining was negative in the control sample that lacked VEGF and was positive in the samples loaded with VEGF. The staining intensity was lower in the heparin/VEGF_2 sample than in the heparin/VEGF_10 sample, confirming the difference in VEGF loaded in these two test samples. The amount of VEGF loaded onto the sample evaluated by ELISA was approximately 970 ng in the case of heparin/VEGF_2 and approximately 4200 ng in the case of heparin/VEGF_10; therefore, the loading yield was 97% and 87%, respectively. The loading efficiency appeared to be inversely proportional to the concentration of VEGF. An inverse relationship was also observed by Singh et al. who obtained loadings of approximately 60% and 45% at initial concentrations of 1 and 10 µg per sample, respectively.³⁸ This effect may be due to well-known ligand–receptor binding-related issues such as saturation, competitive binding, and other factors. Nevertheless, the loading obtained in the current study was relatively high compared to the values reported by

Fig. 6  Morphological changes after heparinization: SEM images suggested that the deposition of heparin was achieved by both cross-linking (CL) and physical adsorption (PA) methods. However, the heparin content was higher in the CL approach with a direct correlation with the CL reaction time. The changes in the surface morphology were clearly visible at a higher magnification, yet the gross morphology of the samples remained largely unaltered. The sample details: (a) the pristine PLCL sample, (b–d) the PLCL samples heparinized via CL for 1, 6 and 15 h respectively, and (e) the sample heparinized via PA for 15 h.
Singh et al., suggesting that the conditions explored in the current study are optimal for loading.

The in vitro release profile of VEGF under physiological conditions was investigated. For both heparin/VEGF_2 and heparin/VEGF_10 samples, a typical two-phase release profile with a relatively short burst release phase followed by a slow and sustained release phase was observed (Fig. 7B). The actual (absolute) amount of VEGF released varied depending on the amount that was initially loaded; however, the release profile was comparable in both cases. The cumulative release was ~25% at 3 h. A high initial burst of VEGF release was also reported by Murphy et al. for VEGF-loaded PLGA scaffolds and by Singh et al. for VEGF-loaded PCL scaffolds and can be explained by the rapid release of weakly bound VEGF molecules especially from the top of the heparin layer. Then the release decreased resulting in the cumulative release of ~60% after 120 h. Finally, VEGF continued to be released slowly and steadily until the end of the study, i.e., at four weeks, during which ~25% VEGF retention was observed. The obtained profile also suggests that additional continuous release of a low amount of VEGF may occur for more than 28 days with the cumulative release of approximately 80%. This can be of potential benefit for efficient support of capsule vascularization when the capsule is implanted in vivo. Furthermore, the drug release kinetics data were found to fit the power law model (Fig. 7B). This indicates that VEGF bound to heparinized-PLCL by physical adsorption was released by diffusion.

Based on the results obtained from all optimization studies, the scaffolds for in vivo experiments were first aminolysed at 22 °C for 30 min, then a heparin layer was covalently immobilized via EDC-NHS chemistry for 6 h and finally VEGF was loaded overnight at 4 °C.

3.5 In vivo response

An unlimited supply of oxygen is crucial for beta cell function as a glycaemic sensor and for adequate insulin release. It is also beneficial for islet engraftment. Most studies in which an artificially created pouch is described emphasize the detrimental effect of inflammation caused by the surgery itself and by reaction to the foreign body. Therefore, the maturation of the cavity was optimized according to fully finished healing after scaffold implantation. Some papers reported promising support of islets with oxygen releasing substances, but, despite this enhanced oxygen supply, a large graft to correct glycaemia had to be used.

In previous studies, we have tested the in vivo behaviour of pristine PLCL capsules and found that the capsules were infiltrated mostly by fibroblasts; preliminary experiments also showed no islet survival and functioning when transplanted in such a cavity (ESI Fig. 3†). Then we also tested the PLCL capsules supplemented with VEGF that should additionally support the formation of a capillary network on the internal surface of the artificial cavity. When the VEGF-loaded scaffolds were implanted subcutaneously in rats, fibrous tissue growth was
insufficient and slow, which resulted in the deficiency of tissue to be stimulated by VEGF present within the scaffold. Therefore, the engraftment and long-term survival of transplanted islets was not improved in that model (data not presented). However, the high specific surface of the PLCL scaffolds\(^\text{21}\) in combination with the attached VEGF holds promise for achieving much better tissue stimulation than that achieved in our previous experiments. Thus, in this study, another site of implantation that is predominantly more compliant to incorporate the scaffold, i.e., in the greater omentum, was tested. As shown previously,\(^\text{20}\) the greater omentum provides a fast-growing tissue. In addition to that, the drainage of blood coming from the graft to the portal vein is much more physiological; it provides a natural drainage of the graft blood towards a portal vein allowing the first pass effect for insulin. This feature is important especially due to the first pass effect of the liver removing approximately 50% of insulin from blood flowing through.

Scaffold implantation was followed by prompt recovery and did not cause any side effects in experimental animals either immediately after the surgery or during the 28-day post-implantation period. The scaffolds maintained their physical integrity and were effectively engrafted to the host tissues throughout the duration of the experiment (ESI Fig. 4a and b†). A typical sound of a vacuum release generated while removing the Teflon bar from the scaffold lumen indicated the airtightness of the cavity within the scaffold and showed that it had been fully penetrated by connective tissue.

Generally, after 4-week implantation, haematoxylin and eosin (H&E) staining, Masson’s trichrome (TRI) staining and special immunohistochemical techniques showed that the fully matured scaffolds were surrounded by a very thin layer of fibrous tissue. Beneath the fibrous capsule, a foreign body-type granulomatous reaction in the presence of mononuclear and multinucleate macrophages was discernible. There were no signs of adhesion or other complications. The connective tissue penetrating the walls of the scaffolds was composed of cellular components and the extracellular matrix. The cellular components consisted predominantly of fibroblasts and myofibroblasts, sporadic inflammatory cells and vascular endothelial cells (ESI Fig. 4c, d and e†). The neovessels were detectable as either thin non-luminized strands of endothelial cells or fully mature branching vascular structures with patent lumina containing red blood cells, lined by CD31 positive endothelial cells and a thin layer of smooth muscle cells. A mild inflammatory reaction involving predominantly small lymphocytes and histiocytes was slightly accentuated along the outer surface of the scaffolds. Importantly, no accumulation of neutrophils and/or eosinophilic leukocytes indicating purulent or hypersensitivity reaction was observed in any animal. The non-cellular component of the connective tissue was composed of small amounts of the ground substance and strands of collagen fibres oriented predominantly radially around the internal cavity with the highest fibre density along the outer surface of each scaffold, reflecting the orientation and size of the pores. Interestingly, no elastin fibres were discernible in the walls of the scaffolds.

### 3.5.1 Dose-optimization study.

To evaluate the quantity of VEGF that is adequate to sufficiently stimulate the formation of a capillary network for this \textit{in vivo} model, we tested the scaffolds with three doses of VEGF immobilized, i.e., 0.4, 2 and 4 µg mg\(^{-1}\) of scaffold (in total 10, 50 and 100 µg per scaffold). The samples were evaluated after 2-week implantation. The histochemical (H&E and TRI) and immuno-histochemical (using anti-CD31 antibody (CD31)) analyses of cross-sections are presented in ESI Fig. 5† (as an overview of the cross-sections) and in Fig. 8 (at higher magnification). Two weeks after the implantation of the heparin-coated scaffold, the connective tissue poor in capillaries reached approximately 20–25% of the wall thickness from the outer surface (Fig. 8a–c). In some parts of the scaffolds (representing less than 10% of the scaffold circumference), the tissue ingrowth was more pronounced and reached the middle part of the wall (ESI Fig. 5†). Coating of the scaffolds with 10 µg of VEGF (i.e., heparin/VEGF\(_{10}\)) did not trigger any stimulation of tissue growth that was detectable macroscopically or by histological techniques (Fig. 8d and e). However, coating with 50 µg of VEGF (i.e., heparin/VEGF\(_{50}\)) stimulated tissue growth considerably and at two weeks after implantation the connective tissue extended nearly to the inner surface of more than 50% of the scaffold circumference (ESI Fig. 4f and Fig. 8g–i). Some vessels were located preferentially around the external surface and appeared to follow the fibrous tissue towards the wall of the scaffold. Vessels were also clearly detected inside the scaffold (Fig. 8i). Infrequent and small vessels were also clearly detected inside the scaffold wall (Fig. 8i). Nevertheless, no vessels had reached the internal surface of the scaffold. An increase in the dose to 100 µg of VEGF (i.e., heparin/VEGF\(_{100}\)) did not have any additional effects on tissue growth or density (Fig. 8j–l). Therefore, we decided to use the scaffold coating with 50 µg of VEGF (i.e., heparin/VEGF\(_{50}\)) for subsequent time-optimization experiments.

### 3.5.2 Time-optimization study.

Time-optimization experiments were performed in order to evaluate the optimal time for sufficient engraftment of the implanted heparin/VEGF\(_{50}\) scaffold with the surrounding tissue of the recipient animal. The scaffolds were excised 2, 3 or 4 weeks after implantation. The experiment did not reveal any significant macroscopically noticeable differences among heparin scaffolds and heparin/VEGF\(_{50}\) scaffolds. The results of H&E, TRI and CD31 analyses after particular time intervals are presented in ESI Fig. 6, 7 and 8† as an overview of the cross-sections. As we anticipated, the depth of connective tissue infiltration and the extent of the microvascular network increased with time. After two weeks, clearly visible tissue reaching nearly the inner surface of the scaffold was present around more than 50% of the scaffold circumference in the VEGF group, as was observed in the dose-optimization experiment (Fig. 8g, h and i). The heparin-coated scaffolds were engrafted incompletely with connective tissue that infiltrated 20–25% of the wall thickness from the outer surface. After three weeks, the soft tissue penetrated the scaffold almost completely around the whole perimeter in the VEGF group. Neovessels were detectable over the whole thick-
ness of the wall; multiple patent vascular structures were present within the outer half of the wall, whereas only scarce mostly non-luminized vascular structures were observed close to the inner surface. The engraftment of the heparin-coated scaffold was much slower, and the connective tissue infiltrated only half of the wall. After four weeks, dense fibrous tissue accompanied by vessels of small calibre completely infiltrated the scaffolds and created sparse granulation tissue

Fig. 8 Dose-dependent experiment (evaluation of the VEGF quantity for sufficient stimulation of cell ingrowth and formation of new vessels two weeks after implantation into the greater omentum): histochemical and immuno-histochemical analyses of heparin–PLCL (a–c), heparin/VEGF_10 (d–f; 0.4 µg VEGF per mg scaffold), heparin/VEGF_50 (g–i, 2 µg VEGF per mg scaffold) and heparin/VEGF_100 (j–l, 4 µg VEGF per mg scaffold) capsules. H&E, TRI and anti-CD31 (black arrows indicate formation of vessels in the scaffold wall) staining; cross-sections. * indicates the internal cavity of the capsule, original magnification: 200×.
within the internal cavity of the scaffold in the VEGF group. The capillary network developed close to the surface of the cavity and inside the cavity, as clearly shown in Fig. 9 at higher magnification. We consider this kind of cavity a potential pouch for a subsequent cellular transplantation. In contrast, in the heparin-coated scaffolds, the connective tissue and occasionally some endothelial structures reached the inner surface of the wall over up to 30% of the scaffold’s circumference.

To summarize, both dose- and time-optimization studies showed that the pre-vascularized pouch can be obtained after a 4-week integration of the heparin/VEGF_50 scaffolds with the host tissue.

3.5.3. Monitoring of islet graft function and structure. The chosen concept meets conditions providing a transplantation cavity surface that is rich in vessels immediately before the transplantation of islets. Until reconnection of islets to the recipient bloodstream, the diffusion distance for oxygen and nutrients is minimal and one can expect a much better islet engraftment than in previously proposed models. This means that islets can be inserted in a convenient environment without the need for additional manipulation such as in the case of scaffolds that are pre-loaded with islets in vitro prior to transplantation.\textsuperscript{54–56} To test the real potential of the VEGF-loaded PLCL capsules for creation of a suitable cavity for pancreatic islets, the pre-vascularized pouch formed by the heparin/VEGF_50/PLCL scaffold after four-week preimplantation in the greater omentum was used. Parameters for monitoring of the graft survival were as follows: blood glucose levels, a continuous increase in body weight, and an intravenous glucose challenge test 40 days after transplantation. The results are presented in ESI Fig. 9† and Fig. 10.

A pancreatic islet graft normalized the blood glucose levels (under 10 mmol l\(^{-1}\)) in five of six diabetic recipients, and in four of them the full function of islets was maintained until the end of the study, i.e., 50 days after islet transplantation (Fig. 10a, the left y-axis and ESI Fig. 9†). The graft did not start to function in one animal, and the function failure appeared in the second animal three weeks after transplantation. Importantly, the body weight of recipients increased in time continuously in the animals with the functioning graft (Fig. 10a, the right y-axis). This observation confirms an adequate insulin secretion by the graft, which prevents the massive loss of glucose (a key nutrient) and stimulates the growth of tissues (as insulin acts also as a powerful growth factor). Furthermore, the intravenous glucose challenge test performed 40 days after islet transplantation confirmed a full function of the graft in four animals (Fig. 10b). This means that the basal insulin secretion can be adequately stimulated by the i.v. administration of glucose (0.5 g kg\(^{-1}\)).

Using anti-insulin and anti-glucagon double-staining, large pancreatic islet conglomerates (Fig. 10c and ESI Fig. 10† at lower magnification) rich in vascular networks were detected in all animals. The observed frequency and localization of alpha-cells resemble their distribution in normal islets. This finding indicates that a cellular dedifferentiation of beta-cells resulting from the transplantation procedure likely did not occur. The good function of the grafts indicated that an even
distribution within a hosting tissue (as is typical in portal vein transplantation) is not necessary for adequate insulin secretion and graft survival until the end of the study. However, the good function of the graft is strongly related to its efficient reconnection to the recipient bloodstream (capillaries close to the internal surface of the pouch) early after its transplantation. Indeed, the presence of vessels within the graft was proved by anti-CD31 staining of the samples; Fig. 10d shows the luminized blood vessels together in the islet grafts and surrounding tissues as well.

In the present work, we made a strong emphasis on the characterization of all performed surface modifications to verify that the modification processes described for 2D model surfaces can be successfully adapted for real 3D scaffolds of particular size and morphology. We proved that the pro-angiogenic factor VEGF, when used in an appropriate dose, can adequately stimulate microvascular network infiltration through the micro-channelled porous scaffold following the fibroblast ingrowth under conditions of optimal timing. In vivo studies indeed confirm the successful creation of a realistic pre-vascularized bed for cellular transplantation.14,57,58 Using a rodent model, we have shown that the created pre-vascularized pouch can successfully host transplanted islets; the newly formed vascular network inside the cavity led to the graft survival and functioning for at least 50 days. In addition, the chosen rat animal model used in this study makes possible testing of the islet/mass (i.e., islet per kg of body weight) ratio similar to the ratio required for clinical applications. In mice, the device must be very small, and the surface, which can be provided to islets for engraftment, is inadequately small. In clinical practice, the proposed concept would be a promising approach particularly for islet auto-transplantations, which can be precisely scheduled; therefore, islet transplantation can be performed during the optimal phase of healing. We believe that
this can be achieved using our approach providing an adequate blood supply to islets at the time of transplantation.

Nevertheless, we expect that further optimization of the coatings with respect to faster cell in-growth through the scaffold wall and application of the “cocktail” of factors supporting vascularization would even more increase the efficiency of islet transplantation and a long-term graft function.

4. Conclusions

We studied the feasibility of adapting modification processes described for planar model surfaces to complex three-dimensional porous scaffolds. We described the 3-step surface functionalization of the PLCL capsules prepared by the Dip-TIPS method and its promising application in bioengineering a realistic pre-vascularized pouch for potential cell transplantation applications.

Importantly, all modification steps preserved the anisotropic channelled pore structure of the capsules as confirmed by microscopic analysis. The success of amination and heparinization of the PLCL surface was proven by ATR-FTIR and XPS spectroscopy analysis. Although the decrease in the $M_w$ of the PLCL polymer induced by aminolysis was time- and temperature-dependent and could reach 50%, the chosen conditions resulted in only a 10% $M_w$ decrease and had no effect on the gross capsule morphology. The covalent coupling approach enabled us to vary the amount of heparin immobilized on the capsule surface from 1.08 to 2.39 µg per mg of scaffold. Performing the EDC/NHS cross-linking for 6 h led to sufficiently high heparin content while the open character of inner pores in the capsule lumen was maintained. Heparinization led to a significant decrease in wettability from 144° for the pristine capsule to 53°, a more convenient value for cell ingrowth in general. VEGF release studies showed that sustained release that followed the power law-based diffusion kinetics occurred for at least four weeks.

The results of in vivo studies in Lewis rats revealed that the scaffolds coated with heparin or heparin/VEGF implanted in the greater omentum are fully biocompatible. Although heparin itself slightly decreased the rate of ingrowth of fibrous tissue and endothelial cells, with stimulation by VEGF the infiltration of the formed microvascular network was deeper than that previously found in native scaffolds. When an appropriate dose of VEGF was used, a suitable pre-vascularized pouch was obtained after a 4-week implantation. The prepared pouch provided for the islet graft survival and functioning for 50 days after islet transplantation in rats. The proposed concept of creation of the pre-vascularized pouch represents a promising approach for clinical cell transplantation.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

The study was supported by Czech Health Research Council, Project No. 16-28254A, the Ministry of Education, Youth and Sports within the National Sustainability Program II (Project BIOCEV-FAR LQ1604) and by the BIOCEV project (CZ.1.05/1.1.00/02.0109), the Ministry of Health Czech Republic – DRO (Institute for Clinical and Experimental Medicine IKEM, IN00023001) and by the Charles University, project GA UK No. 100217.

References

1. O. Iorio, V. Petrozza, A. De Gori, M. Bononi, N. Porta, G. De Toma and G. Cavallaro, J. Invest. Surg., 2018, 1–8, DOI: 10.1080/08941939.2018.1441344.
2. T. F. Davies, Thyroid, 2013, 23, 139–141.
3. A. Dworzynska, K. Paduszynska and L. Pomorski, Pol. Przegl. Chir., 2018, 90, 22–28.
4. Y. Fan, A. Tajima, S. K. Goh, X. Geng, G. Gualtierotti, M. Grupillo, A. Coppola, S. Bertera, W. A. Rudert, I. Banerjee, R. Bottino and M. Trucco, Mol. Ther., 2015, 23, 1262–1277.
5. T. Anazawa, H. Okajima, T. Masui and S. Uemoto, Ann. Gastroenterol. Surg., 2019, 3, 34–42.
6. K. Gao, P. Kumar, E. Cortez-Toledo, D. Hao, L. Reynaga, M. Rose, C. Wang, D. Farmer, J. Nolta, J. Zhou, P. Zhou and A. Wang, Stem Cell Res. Ther., 2019, 10, 34.
7. S. K. Vishwakarma, A. Bardia, C. Lakkireddy, N. Raju, S. A. B. Paspala, M. A. Habeeb and A. A. Khan, Mater. Sci. Eng., C, 2019, 98, 861–873.
8. T. Kunath, A. Natalwala, C. Chan, Y. Chen, B. Stecher, M. Taylor, S. Khan and M. M. K. Muqit, Eur. J. Neurosci., 2019, 49, 453–462.
9. P. Assinck, G. J. Duncan, B. J. Hilton, J. R. Plemel and W. Tetzlaff, Nat. Neurosci., 2017, 20, 637–647.
10. M. Omami, J. J. McGarrigle, M. Reedy, D. Isa, S. Ghani, E. Marchese, M. A. Bochenek, M. Longi, Y. Xing, I. Joshi, Y. Wang and J. Oberholzer, Curr. Diabetes Rep., 2017, 17, 47.
11. A. Pileggi, R. D. Molano, C. Ricordi, E. Zahr, J. Collins, R. Valdes and L. Inverardi, Transplantation, 2006, 81, 1318–1324.
12. A. R. Pepper, B. Gala-Lopez, R. Pawlick, S. Merani, T. Kin and A. M. J. Shapiro, Nat. Biotechnol., 2015, 33, 518–523.
13. J. Kriz, D. Jirak, T. Koblas, G. Vilk, P. Girman, M. Hajek and F. Saudek, Transplant. Proc., 2011, 43, 3226–3230.
14. N. A. Forster, A. J. Penington, A. A. Hardikar, J. A. Palmer, A. Hussey, J. Tai, W. A. Morrison and S. J. Feeney, Islets, 2011, 3, 271–283.  
15. A. J. Hussey, M. Winardi, X. L. Han, G. P. L. Thomas, A. J. Penington, W. A. Morrison, K. R. Knight and S. J. Feeney, Tissue Eng., Part A, 2009, 15, 3823–3833.  
16. A. M. Smink, S. R. Li, D. T. Hertsig, B. J. de Haan, L. Schwab, A. A. van Apeldoorn, E. de Koning, M. M. Faas, 

Biomater. Sci.
