A two-layer skin construct consisting of a collagen hydrogel reinforced by a fibrin-coated polylactide nanofibrous membrane

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Background: Repairs to deep skin wounds continue to be a difficult issue in clinical practice. A promising approach is to fabricate full-thickness skin substitutes with functions closely similar to those of the natural tissue. For many years, a three-dimensional (3D) collagen hydrogel has been considered to provide a physiological 3D environment for co-cultivation of skin fibroblasts and keratinocytes. This collagen hydrogel is frequently used for fabricating tissue-engineered skin analogues with fibroblasts embedded inside the hydrogel and keratinocytes cultivated on its surface. Despite its unique biological properties, the collagen hydrogel has insufficient stiffness, with a tendency to collapse under the traction forces generated by the embedded cells.

Methods: The aim of our study was to develop a two-layer skin construct consisting of a collagen hydrogel reinforced by a nanofibrous polyl-lactide (PLLA) membrane pre-seeded with fibroblasts. The attractiveness of the membrane for dermal fibroblasts was enhanced by coating it with a thin nanofibrous fibrin mesh.

Results: The fibrin mesh promoted the adhesion, proliferation and migration of the fibroblasts upwards into the collagen hydrogel. Moreover, the fibroblasts spontaneously migrating into the collagen hydrogel showed a lower tendency to contract and shrink the hydrogel by their traction forces. The surface of the collagen was seeded with human dermal keratinocytes. The keratinocytes were able to form a basal layer of highly mitotically-active cells, and a suprabasal layer.

Conclusion: The two-layer skin construct based on collagen hydrogel with spontaneously immigrated fibroblasts and reinforced by a fibrin-coated nanofibrous membrane seems to be promising for the construction of full-thickness skin substitute.

Keywords: full-thickness skin substitutes, collagen hydrogel, fibroblast and keratinocyte co-cultivation, fibrin, nanofibrous membrane

Introduction

The skin is the organ of the human body that is most exposed to external agents. It provides a natural barrier between the surrounding environment and the human internal organs. From the tissue engineering point of view, the most important feature of the skin is its ability to self-remodel and regenerate, and to replace the tissue that is continuously being lost. Anatomically, the skin tissue can be divided into three layers: epidermis, with its main cellular type of keratinocytes; dermis, consisting mainly of fibroblasts; and hypodermis, which is rich in adipose cells.

For many years, tissue engineers have been trying to construct under in vitro
conditions an appropriate full-thickness skin equivalent that contains all three layers of the skin. However, there are still many limitations.

Two-dimensional (2D) flat materials are commonly used for cell cultivation, but these materials can induce an apical-basal polarity, which can restrict the adhesion and the migration of the cells in the x-y axis, leading to non-physiological behavior of the skin cells, e.g., fibroblasts. A more complex three-dimensional (3D) microenvironment is therefore necessary for a non-prescribed polarity of the fibroblasts, and also for their spreading and migration in all three dimensions. Novel 3D biomaterials aim to mimic the physiological environment of natural extracellular matrix (ECM), and thus to ensure better conditions for cell adhesion, spreading, proliferation, migration, matrix protein synthesis, and differentiation.

Synthetic materials made of polylactide (PLA) or polycaprolactone (PCL) are commonly processed into fibrous scaffolds by conventional electrospinning, followed by cell seeding. Although the synthetic scaffolds are non-toxic and mimic the mechanical and morphological properties of native tissues, they often do not provide sufficient support for cell adhesion and proliferation. The synthetic materials are therefore often modified with biomolecules in order to improve their attractiveness for cells, and thereby to accelerate the healing process. For example, in our previous studies, nanofibrous PLA membranes were modified with fibrin, a key protein occurring during wound healing, in order to enhance the adhesion and growth of human dermal fibroblasts. Moreover, we found out that two morphologically different structures of fibrin coatings had a significant influence on the cell behavior. In addition, other studies have shown that collagen-containing composites or collagen-coated nanofibers are more attractive for cells than pure synthetic scaffolds, due to their greater biocompatibility and biodegradability. Fu et al. found that collagen-coated PCL nanofibers increased the proliferation of skin fibroblasts. Fibrinogen-coated nanofibers decreased the growth of fibroblasts, but they supported the migration and the expression in these cells of α-smooth muscle actin, which is a typical marker for fibroblast differentiation into myofibroblasts. It was also found that the PCL/collagen nanofibrous membrane alone did not stimulate keratinocyte migration, while further coating of this membrane with an ultrafine fibrous collagen network significantly increased the cell motility. Mahjour et al. used a layer-by-layer method to construct PCL/collagen nanofibers with fibroblasts and keratinocytes, and applied this construct to full-thickness wounds on nude mice. Native collagen in any form therefore increases fibroblast chemotaxis, angiogenesis and keratinocyte migration.

The most recent studies have shown that fibroblast behavior is more physiological if the fibroblasts are entrapped in a 3D structure of hydrogels. The hydrogels are able to assemble from a liquid monomeric phase to a polymeric mesh network under a certain temperature, pH, and enzymatic activity. The hydrogels are covalently or non-covalently cross-linked, which enables the encapsulation of living cells. Various nature-derived hydrogels for wound healing and regeneration applications have been described in earlier studies. These hydrogels were derived mainly from collagen, fibronectin, alginates, gelatin, hyaluronic-chitosan or chitosan-gelatin, which are molecules similar to the macromolecular-based components of the body. However, one of the most discussed tasks in tissue engineering is the correlation between the stiffness of a hydrogel and the kinetics of cell adhesion and migration, which is related to the degradation and the remodeling time of the material. The degradation time and the remodeling time should be in balance, and both are highly dependent on the behavior of the encapsulated cells. When the cells are entrapped in hydrogels, the surrounding environment changes their gene expression pattern and stimulates biosynthesis of their own ECM molecules, which results in degradation and remodeling of the hydrogels. Various matrix metalloproteases (MMP-1, MMP-2, MMP-8, MMP-13 and MMP-14) are involved in the degradation processes and in the migration of the cells in the hydrogels. In addition to the degradation, the cells can also synthesize their own molecules of ECM, and in this way they can remodel the hydrogels. The morphology of fibroblasts in 3D collagen hydrogels varies from bipolar to dendritic in shape, depending on the stiffness and the tension of the hydrogels. Jiang and Grinnell and Grinnell also described the mechanical interactions between the collagen hydrogel and the fibroblasts. Entanglement of the fibroblasts into the surrounding matrix moves their forces to the hydrogel, and this results in contraction or compaction of the hydrogels. In order to develop a large-size clinically applicable transplant, Braziulis et al. used plastic compression for a collagen type I hydrogel to increase the stability and to improve the mechanical properties of the material. Another way to improve the mechanical properties of the hydrogels is to reinforce them with the use of biodegradable synthetic scaffolds, e.g., poly(lactic-co-glycolide) (PLGA) nanofibrous membranes or knitted meshes.
In this study, we prepared a two-layer cell construct consisting of a poly-L-lactide (PLLA) nanofibrous membrane coated with a thin nanofibrous fibrin mesh and seeded with human dermal fibroblasts. After the attachment and the initial growth of the fibroblasts, a collagen hydrogel was formed on the fibroblast-seeded membrane. Finally, human epidermal keratinocytes were seeded on top of the collagen hydrogel. We hypothesized that the 3D-structured collagen hydrogel would enable the fibroblasts to migrate into the hydrogel from the membrane, and the collagen hydrogel with naturally immigrated fibroblasts would therefore simulate the skin dermis. The top of the collagen gel with a layer of adhering and growing keratinocytes would simulate the skin epidermis. The biodegradable PLLA nanofibrous membrane would serve as a substrate for the initial attachment and growth of fibroblasts, and as a mechanical support for a soft collagen hydrogel with skin cells. Coating the membrane with a thin nanostructured fibrin mesh was expected to enhance fibroblast adhesion and proliferation, leading to accelerated fibroblast migration into the collagen hydrogel. We supposed that spontaneous migration of the fibroblasts from the membrane into the collagen hydrogel could lower the tendency of these cells to contract and to shrink the gel by their traction forces, which occurs when fibroblasts are directly embedded into the collagen hydrogel during the gelling process.\(^{17,32,33}\) The aims of this study were to evaluate the ability of human dermal fibroblasts to migrate into the collagen hydrogel from the nanofibrous membrane, to compare the behavior and the migration of the fibroblast from a non-coated membrane and from a fibrin mesh-coated membrane, and to evaluate the behavior of human epidermal keratinocytes adhering on the collagen hydrogel surface.

**Materials and methods**

**Preparation of nanofibrous PLLA membranes**

A nanofibrous PLLA membrane was prepared by the electrospinning process. PLLA with a molecular weight of 45,000–55,000 Da (Polyscitech, Akina Inc., Lafayette, IN, USA) was dissolved in a solvent system composed of a mixture of chloroform and ethanol in a ratio of 9:1 (v/v, Penta s.r.o., Prague, Czech Republic) to a final polymer concentration of 10 wt%. The polymer solution was then used for the electrospinning process. Nanofibrous membranes were prepared using Nanospider NS 1WS500U (Elmarco s.r.o., Liberec, Czech Republic). The resulting fibers 940±340 nm in diameter were collected on a spunbond layer at a distance of 190 mm from the electrospinning string. The collector rolled at a rate of 10 mm/minute. The applied voltage was 40 kV for the string and 10 kV for the collector. The process was carried out at an ambient temperature of 25°C and at relative humidity of 30%.

**Cell harvesting and cultivation**

The two-layer construct consisted of human neonatal dermal fibroblasts and human epidermal keratinocytes. The human neonatal dermal fibroblasts were purchased from Lonza (Basel, Switzerland, Cat. No. CC-2509) and were cultivated in Dulbecco’s Modified Eagle’s Medium (DMEM; Sigma-Aldrich Co., St Louis, MO, USA) with 10% of fetal bovine serum (FBS; Sebak GmbH, Aidenbach, Germany) and 40 µg/mL of gentamicin (LEK, Ljubljana, Slovenia). The human epidermal keratinocytes were harvested from the skin of a young adult donor with approval of the local ethics committee (Ethics committee, Medical Faculty of Charles University, approval number EK-VP/15/0/2017) and after confirmed written informed consent provided by the donor that is in compliance with the Declaration of Helsinki. The skin graft was digested in 0.25% trypsin solution (Sigma-Aldrich Co.) for 1 hour at 37°C. After that, the epidermis was separated from the dermis. The epidermis was inserted in a drop of the cultivation medium. After a while, the cells were spontaneously released from the epidermis graft, and they were collected and seeded on mitomycin-inactivated 3T3 mouse embryonic fibroblasts adhered on the bottom of a cultivation flask. The 3T3 mouse embryonic fibroblasts were purchased from Sigma-Aldrich Co. (Cat. No. 93061524-1VL). The keratinocytes were cultivated in a mixture of the cell cultivation DMEM medium and Ham’s Nutrient Mixture F12 medium (DMEM/F 12, Sigma-Aldrich Co.) in a ratio of 3:1. The mixture was supplemented with 10% of FBS (Sebak GmbH, Germany), 10 ng/mL epidermal growth factor (EGF, Sigma-Aldrich Co.), 10 µg/mL insulin (Sigma-Aldrich Co.), 0.4 µg/mL hydrocortisone (Sigma-Aldrich Co.), 10\(^{10}\) M cholera toxin (Sigma-Aldrich Co.), 1000 U/mL of streptomycin and 0.1 mg/mL of penicillin (Sigma-Aldrich Co.).
The fibroblasts and keratinocytes were cultivated in an incubator at 37°C with a humidified atmosphere saturated with 5% of CO₂ in the air.

Preparation of the two-layer skin construct

A two-layer cell construct was composed of a PLLA nanofibrous membrane pre-seeded with human dermal fibroblasts, a collagen hydrogel and human dermal keratinocytes (Figure 1). The membrane samples were fixed in Cell Crown inserts (Scaffdex Ltd., Tampere, Finland) fitting into 24-well cell culture plates (TPP, Trasadingen, Switzerland) and seeded with human dermal fibroblasts. The fibroblasts were seeded on the membrane at a density of 30,000 cells, and were cultivated in DMEM supplemented with 10% of FBS (Sebak GmbH, Germany) and 40 µg/mL of gentamicin (LEK, Slovenia). To enhance the adhesion and proliferation of the fibroblasts, the PLLA membrane was coated with a thin nanofibrous fibrin mesh before cell seeding. After 3 days of fibroblast cultivation, the collagen hydrogel was prepared on the fibroblast-seeded membrane, and the fibroblasts started to migrate into the hydrogel. The ability of the fibroblasts to migrate into a collagen hydrogel from a non-coated membrane and from a fibrin-coated PLLA membrane was compared in four time intervals (on days 1, 3, 7 and 14 after collagen preparation, ie on days 4, 6, 10 and 17 after cell seeding). The preparation of the fibrin mesh and the collagen hydrogel is described in detail below in the following sections.

The construct consisting of the nanofibrous fibrin-coated PLLA membrane was then selected for seeding with keratinocytes on top of the collagen hydrogel (Figure 1). The keratinocytes were seeded at a density of 60,000 cells after 4 days of collagen hydrogel preparation (resp. of fibroblast migration). After the keratinocytes had been seeded, the medium was replaced by DMEM/F12 with supplements, as mentioned above. After 2 days of keratinocyte cultivation, the culture medium was further supplemented with 2-phospho-L-ascorbic acid trisodium salt (AA) at a concentration of 50 µg/mL, in order to stimulate the cells to synthesize ECM proteins. The medium was changed every 2 days.

Preparation of the fibrin mesh on the nanofibrous membrane

The fibrin mesh on the membrane was prepared according to our previously published work. The fibrin was formed by activation of water-soluble human fibrinogen (341576; EMD Millipore, Billerica, MA, USA) with human thrombin (T6884; Sigma-Aldrich Co.). Briefly, the fibrinogen in a concentration of 10 µg/mL, diluted in TRIS buffer (consisting of 50 mM TRIS-HCl, 100 mM NaCl and 2.5 mM CaCl₂), was adsorbed on the membrane for 1 hour. After rinsing with TRIS buffer, the adsorbed fibrinogen was activated with thrombin (2.5 U/mL in TRIS buffer) for 15 minutes. The thrombin-activated surface was directly treated without TRIS rinsing with a

**Figure 1** Scheme of the preparation of the constructs for evaluating the migration of the fibroblasts, consisting of a non-coated membrane or a fibrin-coated PLLA nanofibrous membrane seeded with human dermal fibroblasts and covered with a collagen hydrogel. The construct with the fibrin-coated PLLA nanofibrous membrane was then selected for seeding with keratinocytes and for preparing a bilayered construct simulating the dermis and the epidermis of the natural skin.

**Abbreviation:** PLLA, poly-L-lactide.
solution of 200 µg/mL of fibrinogen in a TRIS buffer and 0.5 U/mL of antithrombin III (Chromogenix, Milano, Italy) in deionized water for 1 hour to form a stable thin fibrin mesh. The antithrombin III blocked the unreacted thrombin remaining on the membrane in order to prevent the formation of a 3D fibrin gel.

Preparation of the collagen hydrogel on a nanofibrous membrane pre-seeded with fibroblasts

The type I collagen used for preparing the hydrogel matrix for the migration of the fibroblasts was isolated from rat tails. The tendons were removed from the rat tails, and were digested in 0.1 M acetic acid for 48 hours. The collagen digested in acetic acid was ultracentrifuged at the maximum speed (approx. 46,000 x g) for 1 hour in order to remove the tissue debris. The supernatant containing the collagen was neutralized with 0.1 M sodium hydroxide and was centrifuged at 1,500 x g for 10 minutes. The supernatant was removed and the collagen pellet was lyophilized. The lyophilized collagen I was dissolved in 0.02 M acetic acid to a concentration of 5 mg/mL and was stored as a stock solution.

The collagen hydrogel was prepared on day 3 after seeding of the fibroblasts. For 1 mL of 3 mg/mL collagen suspension 600 µL of collagen stock solution (5 mg/mL), 390 µL of DMEM (supplemented with 10% FBS and 40 µg/mL of gentamicin) and 10 µL of sodium bicarbonate (Sigma-Aldrich Co.) were mixed, and a total volume of 400 µL was applied on the membrane in order to prevent the formation of a 3D fibrin gel. The collagen hydrogel disks were approximately 3 mm in thickness for seeding the cells. The 840 µL of collagen stock solution (5 mg/mL) was diluted in 546 µL of DMEM (supplemented with 10% FBS and 40 µg/mL of gentamicin) and 14 µL of sodium bicarbonate (Sigma-Aldrich Co.). The total volume of 1400 µL of collagen solution (3 mg/mL) was applied on the PLLA membrane without cells. The prepared unconfined samples were 12.6 mm in diameter, and the compression test was performed on six parallel samples. The fresh samples in PBS were placed between two stainless steel cylinders at 21°C. The cylinder was attached to the Kistler 9203 force sensor (1000 Hz, sensitivity 0.001 N, range±10N, Kistler, Switzerland) and moved in quasi-static mode at 2 mm/minute (stepper motor PD28-3-1021, Trinamic, Hamburg, Germany). The initial length l₀ was the length at detecting positive force (height F≠0 =1.82±0.27 mm).

The force and the position data were recalculated into the stress-strain curve reflecting l₀, σ and A₀, respectively. The mechanical properties of the composite material were characterized by the Young’s moduli in the linear parts of the stress-strain curves. The data were acquired by the Dewetron data acquisition system (Dewetron, Grambach, Austria) and were analyzed in Matlab 2017a (Mathworks, USA).

Morphology of the fibrin mesh and the collagen hydrogel on the nanofibrous membrane

The morphology of the fibrin mesh and the collagen hydrogel prepared on a nanofibrous membrane was studied by scanning electron microscopy (SEM). The morphology of the fibrin mesh degraded by cells while they were being cultivated on the fibrin-coated membrane was further observed by fluorescence confocal microscopy.

For SEM, the nanofibrous membrane with fibrin was fixed with a 1% solution of osmium tetroxide, and the collagen hydrogel was fixed with 4% paraformaldehyde in PIPES buffer (pH~7) (Sigma-Aldrich Co.). The samples were dehydrated in the standard gradient (20–40 minutes in each of 30%, 50%, 70%, 96%, and 100% ethanol solutions). Samples from absolute alcohol were incubated in the following series of drying solutions: 1:1 ethanol/acetone, 100% acetone, 1:1 acetone/hexamethyldisilazane, 100% hexamethyldisilazane, with each step taking 20–40 minutes. Air drying of the sample for a period of 24 hours was followed by coating with gold-palladium. The surface morphology and the topography of the fibers were observed using a high-resolution
scanning electron microscope with a Quanta FEG 250 field emission cathode (FEI, USA). High-vacuum mode, acceleration voltage 10 kV and secondary electrons mode using an Everhart-Thornley detector (ETD) were used for imaging.

The morphology of the fibrin mesh was evaluated by immunofluorescence staining performed on freshly prepared cell-free samples and on samples cultivated with cells in various time intervals. The samples were treated with a solution of 1% bovine serum albumin and 0.1% Tween 20 in a phosphate-buffered saline (PBS, all Sigma-Aldrich Co.) for 20 minutes at room temperature, in order to block non-specific binding sites. Then the samples were incubated with a primary antibody against human fibrinogen (A0080; polyclonal rabbit antibody, Dako Denmark A/S, Glostrup, Denmark) diluted in the blocking solution (1% albumin and 0.1% Tween 20 in PBS) in a ratio of 1:200 for 1 hour at 37°C. After that, the samples were rinsed with PBS and were incubated with a secondary antibody, ie goat anti-rabbit F(ab')2 fragments of IgG (H + L), conjugated with Alexa Fluor® 488 (A11070; Molecular Probes) diluted in the blocking solution in a ratio of 1:400 for 1 hour at room temperature in the dark. The samples were scanned on a Leica TCS SPE DM2500 upright confocal microscope, objective 40x/1.15 NA oil.

Visualization of the cell morphology and migration in the two-layer construct

The morphology of the dermal fibroblasts adhered on the nanofibrous membrane and migrated into the collagen hydrogel was studied by staining the filamentous actin (F-actin) cytoskeleton and the cell nucleus. The spreading and the morphology of keratinocytes were evaluated by the same staining, and, in addition, by staining the intermediate filaments containing keratin 14.

Before staining, the cells were fixed with 4% paraformaldehyde, and the cell membrane was permeabilized with 0.1% Triton X-100 (Sigma-Aldrich Co.) diluted in PBS for 30 minutes at room temperature. Then the samples were treated in a solution of 1% bovine serum albumin and 0.1% Tween 20 in PBS (all Sigma-Aldrich Co.) to block non-specific binding sites. Subsequently, the F-actin was stained with phalloidin conjugated with tetramethylrhodamine isothiocyanate (TRITC) fluorescent dye (5 µg/mL, diluted in PBS, Sigma-Aldrich Co.), and the cell nucleus was stained with Hoechst #33258 (5 µg/ml diluted in PBS, Sigma-Aldrich Co.) for 1 hour at room temperature in the dark. The keratin 14 of the keratinocytes was stained by incubating the samples with primary antibody to human cytokeratin 14 (ab7800, monoclonal mouse antibody, clone LL002, Abcam, Cambridge, United Kingdom) diluted in a blocking solution (1% albumin and 0.1% Tween 20 in PBS) in a ratio of 1:200 for 1 hour at 37°C. The samples were rinsed with PBS, and were then incubated with a secondary antibody, ie goat anti-mouse F(ab')2 fragments of IgG (H + L), conjugated with Alexa Fluor® 488 (A11017; Molecular Probes) diluted in the blocking solution in a ratio of 1:400 for 1 hour at room temperature in the dark. The samples for evaluating fibroblast migration into the collagen hydrogel were scanned on a Leica TCS SPE DM2500 upright confocal microscope, objective 40x/1.15 NA oil and 20x/20x/0.75 IMM CORR CS2; FWD 0.66, zoom 2x. The maximal depth of fibroblast migration into the collagen hydrogel from the non-coated or fibrin-coated membranes was measured during scanning on three parallel samples. Statistical significance was evaluated using nonparametric Kruskal-Wallis One Way Analysis of Variance on Ranks, Dunn’s Method, statistical significance ($p$≤0.05). Samples of the whole two-layer construct were scanned on a Dragonfly 503 (Andor, Belfast, NI, UK) spinning disk confocal microscope with a Zyla 4.2 PLUS sCMOS camera, objective HC PL APO 20x/0.75 IMM CORR CS2; Free Working Distance =0.66 mm.

Cell mitochondrial activity

The proliferation and the viability of the human dermal fibroblasts that adhered on the membrane and that migrated into the collagen hydrogel were determined by the activity of cell mitochondrial enzymes, using the Cell Titer 96® AQueous One Solution Cell Proliferation Assay (MTS, Promega Corporation, Madison, WI, USA). The metabolic activity was determined on days 1, 3, 7, and 14 after the collagen hydrogel was prepared.

The collagen hydrogel with immigrated fibroblasts was carefully removed from the membrane, and both parts of the sample were moved into fresh cell culture wells to avoid the influence of cells adhered to the bottom of the well. The metabolic activity of the cells was separately determined in the hydrogel and on the membrane beneath the hydrogel. The non-coated
membranes and the fibrin-coated membranes without collagen hydrogel seeded with fibroblasts were used as control samples to observe any difference in the metabolic activity of the cells adhered on the two types of membranes. The assay was performed according to the manufacturer’s protocol. The amount of formazan dye produced by the cells after 2 hours of incubation was quantified by measuring the absorbance. The absorbance was measured with wavelength 490 nm and with reference wavelength 700 nm, using a VersaMax ELISA Microplate Reader spectrophotometer (Molecular Devices Corporation, Sunnyvale, CA, USA).

Four parallel samples were used for each experimental group and time interval. The experiment was performed twice. The data was presented as the mean ± standard deviation (from eight measurements). Statistical significance was evaluated using parametric analysis of variance (ANOVA), with the Tukey post hoc test for pairwise comparison. Values of \( p \leq 0.05 \) were considered as significant.

**Results**

**Morphology of the fibrin mesh and the collagen hydrogel on the nanofibrous membrane**

The PLLA membrane consisted of randomly-oriented fibers. The diameter of the fibers varied in a large range, from tens of nanometers to micrometers. The average diameter of the fiber was 940 nm ±340 nm (Figure 2A and B).

The fibrin covered the fibers of the membrane, and it also formed a thin homogeneous nanofibrous mesh on the surface of the membrane and among its fibers. The diameter of the fibers of the fibrin mesh ranged from tens of nanometers to approx. 100 nm (Figure 2C and D).

The nanofibers were formed during the collagen gelling process. They were randomly-oriented in the bulk with a diameter in tens of nanometers. The average diameter of the nanofibers was 47±7.6 nm (Figure 2E and F).

**Collagen hydrogels unconfined compression test**

The stress strain curves representing the collagen hydrogel disks displayed a linear response up to 21.29±4.36%
deformation at unconfined compression. The Young’s modulus was 3.0±1.4 kPa in this region. The next linear region was detected from approximately 56.69±9.91% deformation, where the Young’s modulus was 89.4±13.2 kPa at unconfined compression; however, structural changes occurred in the sample disks in this region.

**Migration of dermal fibroblasts into the collagen hydrogel from fibrin-coated vs non-coated membranes**

We prepared a construct consisting of a pristine, ie non-coated, PLLA nanofibrous membrane or consisting of a membrane coated with a thin nanofibrous fibrin mesh, human dermal fibroblasts and the collagen I hydrogel. The collagen hydrogel was prepared on the fibroblast-seeded membrane after 3 days of cell cultivation. We compared the ability of fibroblasts to migrate into the collagen hydrogel from a non-coated membrane and from a fibrin-coated PLLA membrane in four time intervals (on days 1, 3, 7, and 14 after collagen preparation, ie on days 4, 6, 10, and 17 after cell seeding).

The cell proliferation and viability were determined by measuring the activity of cell mitochondrial enzymes (ie cell metabolic activity). We separately determined the metabolic activity of the fibroblasts migrating in the collagen hydrogel and the fibroblasts adhered on the membrane beneath the hydrogel. Moreover, the metabolic activity of the fibroblasts adhered on the control membranes (non-coated and fibrin-coated membranes) without the collagen hydrogel was evaluated in order to see the effect of the fibrin mesh on fibroblast proliferation. There was greater metabolic activity of the cells proliferating on the control fibrin-coated membrane (without collagen) than on the control non-coated membrane (without collagen) (Figure 3, the first pair of columns). We observed these differences in all cultivation time intervals, with statistical significance on day 1. Similarly, the fibrin mesh also increased the fibroblast proliferation on the membrane beneath the collagen (Figure 3, the second pair of columns), with statistical significance on days 1, 3 and 7 after the collagen preparation.

![Figure 3](image_url)

**Figure 3** Mitochondrial activity of human dermal fibroblasts migrated into the collagen hydrogel from the non-coated PLLA membrane (Pristine) or from the fibrin-coated (Fibrin) PLLA membrane on day 1 (A), day 3 (B), day 7 (C) and day 14 (D) after preparation of the collagen. Cell mitochondrial activity on the control membrane without the hydrogel (PLLA, first pair of columns), separately on the membrane beneath the hydrogel (PLLA beneath gel, second pair of columns) and in the hydrogel (Gel, third pair of columns). Arithmetic mean ± SD from 8 measurements, ANOVA, Student-Newman-Keuls method, statistical significance (p≤0.05): * compared with a non-coated membrane (Pristine).

**Abbreviation:** PLLA, poly-L-lactide.
hydrogel was prepared (Figure 3A–C). The metabolic activity of the cells migrated into the collagen hydrogel from the fibrin-coated membrane was significantly higher than the activity of the cells migrated into the hydrogel from the non-coated membrane in all time intervals (Figure 3, third pair of columns).

The results for cell mitochondrial activity were confirmed by visualization of the cells migrating from the membrane into the collagen hydrogel using immunofluorescent confocal microscopy. The cells migrating into the collagen hydrogel and the cells adhering on the membrane beneath the hydrogel were separately visualized in various time intervals of cell migration (Figure 4). In addition, the fibroblasts adhered on the control membranes (non-coated and fibrin-coated membranes) without the collagen hydrogel were observed (Figure 5). Before the preparation of the collagen hydrogel on the membrane after the 3rd day of cell cultivation, the cells were almost in a confluent layer on the fibrin-coated membrane, while on the non-coated membrane, the cells had reached a lower population density (Figure 5, Day 0). On day 1 of cell migration, the cells started to migrate into the collagen hydrogel only from the fibrin-coated membrane. On the non-coated membrane, the cells remained attached to the membrane or to the bottom of the hydrogel (Figure 4, Day 1). In other words, the cells adhering and proliferating on the fibrin-coated membrane started to migrate 1 day after the hydrogel had been prepared, while the cells on the non-coated membrane started migrating later. The supporting influence of the fibrin mesh on the attachment of cells to the membrane and on their proliferation led to a higher number of cells migrating into the collagen hydrogel in all time intervals (Figure 4, the 1st and 2nd rows). In addition, the cells migrated deeper into the hydrogel from the fibrin-coated membrane than from the non-coated membrane, and this difference reached statistical significance on day 3. The depth of the cell migration increased significantly from the 3rd day to the 7th day (Figure 6). On day 14, the cells migrated throughout the collagen hydrogel, and they formed a confluent layer on top of the hydrogel, mainly in the case of the fibrin-coated membrane (Figure 4, 1st and 2nd row).

**Figure 4** Human dermal fibroblasts in the collagen hydrogel migrated from a non-coated membrane (Gel PLLA) and from a fibrin-coated membrane (Gel PLLA F) for 1, 3, 7, and 14 days, or human dermal fibroblasts proliferated on the non-coated membrane (PLLA beneath gel) or on the fibrin-coated membrane (PLLA F beneath gel) beneath the collagen hydrogel for 4, 7, 10, and 17 days. The fibrin was stained by immunofluorescence (Alexa 488, green). The cells were stained with phalloidin-TRITC (red; F-actin cytoskeleton) and with Hoechst #33,258 (blue; cell nuclei). Leica TCS SPE DM2500 confocal microscope, 20x/0.75 IMM CORR CS2 zoom 2x or obj. 40x/1.15 NA oil, maximal intensity projection images.

Abbreviation: PLLA, poly-L-lactide.
The cells that remained adhering on the membrane beneath the collagen hydrogel were able to divide and proliferate, and their population increased in time (Figure 4, the 3rd and 4th rows). If we compare the cell behavior on the membranes beneath the collagen hydrogel (Figure 4, the 3rd and 4th rows) and on the control membranes (without collagen, Figure 5), the number of adherent cells was significantly higher on the control membranes in all time intervals. On the control fibrin-coated membrane, the cells gradually degraded and reorganized the fibrin mesh. On day 14, the fibrin mesh was almost degraded, and only residues of it remained on the membrane (Figure 5, 2nd row), while the fibrin coating on the membrane beneath the hydrogel was only slightly altered by the cells (Figure 4, 4th row).

The collagen hydrogel was not considerably contracted and shrunk by the fibroblasts migrating in the collagen over a period of 14 days (Figure 7). The fresh collagen hydrogel prepared on the fibroblast-seeded

![Figure 5](image1)

Figure 5 Human dermal fibroblasts on the control non-coated membrane (PLLA) and on the fibrin-coated membranes (PLLA F) without the collagen gel. The cells on the membranes on the 3rd day after they were seeded and before collagen hydrogel preparation (Day 0), and cultivated for 1, 3, 7, and 14 additional days (ie 4, 7, 10, and 17 days in total). The fibrin was stained by immunofluorescence (Alexa 488, green). The cells were stained with phallloidin-TRITC (red; F-actin cytoskeleton) and with Hoechst #33258 (blue; cell nuclei). Leica TCS SPE DMi8500 confocal microscope, objective 40x/1.15 NA oil, maximal intensity projection images.

**Abbreviation:** PLLA, poly-L-lactide.

![Figure 6](image2)

Figure 6 Depth of migration of human dermal fibroblasts into the collagen hydrogel from the non-coated PLLA membrane (PLLA) or from the fibrin-coated PLLA membrane (PLLA + F) on days 3, 7, and 14 of cell migration. Nonparametric Kruskal-Wallis One Way Analysis of Variance on Ranks, Dunn’s Method, statistical significance (p≤0.05): * in comparison with the non-coated membrane (PLLA), # in comparison with day 3.
membrane was approximately 8.9 mm in diameter. After 14 days of cell migration, the diameter of the hydrogel reached 8.4±0.3 mm (presented as a mean ± standard deviation). In addition, there was no obvious difference between the remodeling of the collagen hydrogel by the cells migrating from the non-coated membrane (hydrogel diameter was 8.3±0.4 mm) and from the fibrin-coated membrane (hydrogel diameter was 8.5±0.1 mm).

Creating the two-layer construct consisting of dermal fibroblasts and keratinocytes

A two-layer construct was prepared using a PLLA nanofibrous membrane coated with a thin nanofibrous fibrin mesh and seeded with human dermal fibroblasts. The collagen hydrogel prepared after 3 days of fibroblast seeding allowed the fibroblasts to migrate inside the gel. The human dermal keratinocytes were seeded on top of the collagen hydrogel after 4 days of fibroblast migration.

Before the keratinocytes were seeded, the fibroblasts were allowed to immigrate into the collagen hydrogel (Figure 8A and B). Inside the hydrogel, the cells were spread, and showed a spindle-like morphology with well-developed F-actin microfilaments (Figures 8 and 9), as we also showed in the previous section in Figure 4. After the collagen hydrogel had been seeded with keratinocytes, the fibroblasts continued proliferating inside the hydrogel and migrating through the hydrogel from the bottom to the top (Figure 9). The keratinocytes were able to attach to the collagen hydrogel, and to divide and proliferate throughout the experiment, ie for 14 days. The number of proliferating keratinocytes increased with cultivation time. The cells had already reached a confluent layer on the whole surface of the collagen hydrogel 7 days after keratinocyte seeding (Figure 9C and D). The keratinocytes had well-developed filaments containing cytokeratin 14 (Figure 9). Video S1 shows the keratinocytes and fibroblasts in the collagen hydrogel in individual horizontal sections from the top to the bottom of the collagen hydrogel. On the hydrogel surface, the keratinocytes formed a suprabasal layer of large cells frequently without cell nuclei and a confluent basal layer of small cells that were still dividing and proliferating. The layer of keratinocytes does not allow the fibroblasts to migrate throughout the whole volume of the collagen hydrogel, thus the fibroblasts remained inside the hydrogel without migrating to the surface (Figure 9). If the keratinocytes were not seeded on the collagen hydrogel, the fibroblasts migrated through the whole thickness of the gel and formed a confluent layer on top of the hydrogel (Figure 8E–H).

Figure 10 and Video S2 show the entire two-layer construct after 14 days of keratinocyte cultivation. The bottom of the skin construct is composed of a nanofibrous membrane with a confluent layer of proliferating fibroblasts (red) and with...
residues of the fibrin mesh (green) degraded by the fibroblasts. The fibroblasts are homogeneously immigrated inside the collagen hydrogel, and the keratinocytes are adhered in the confluent layer on top of the collagen hydrogel (green).

Adding keratinocytes into the construct did not significantly promote shrinkage of the collagen hydrogel. There was a similar slight alteration in the morphology and in the contraction of the collagen hydrogel during cell cultivation when only fibroblasts were seeded into the construct, see our comments in the previous section, and Figure 7.

Discussion
Many previous studies have shown that the 3D microenvironment of hydrogels provides better physiological conditions than 2D flat substrates for cell spreading, proliferation, migration and differentiation. The cells embedded in the hydrogels, especially fibroblasts and other mesenchymal cells, tend to be more spread, with a typical spindle-like morphology, and they form a network with cell contacts in all three dimensions. On
the other hand, keratinocytes with their apical-basal polarity prefer a 2D structured surface of the hydrogels. Fujisaki et al.\(^{36}\) and many other researchers have observed that collagen hydrogels, mainly collagen IV and collagen I, support the adhesion, proliferation and stratification of keratinocytes.\(^{17,36,37}\) In addition, the biosynthesis of ECM molecules by fibroblasts and keratinocytes is enhanced by the physiological conditions of the hydrogels, which is important for hydrogel remodeling and for separating the two cell types by forming the basement membrane.\(^{28,38,39}\) Although the encapsulated fibroblasts are separated from the keratinocytes growing on the surface, their communication continues to be mediated by the cytokines and growth factors released from the cells in a paracrine manner, or by cell-hydrogel mechanosensation.\(^{40-43}\) The paracrine-based relationship of the cells can be regulated by changing the permeability and the structural properties of the cell-encapsulating substances. For example, Chiu et al.\(^{44}\) modulated the permeability of fibrin constructs by varying the concentration of fibrinogen and thrombin.\(^{44}\) Bader et al.\(^{45}\) changed the keratinocyte-fibroblast paracrine communication by gelatin-based semi-interpenetrating networks.\(^{45}\) Moreno-Arotzena et al.\(^{46}\) characterized the collagen hydrogel as a biomaterial with a higher void ratio and higher permeability than the fibrin hydrogel.\(^{46}\) One of the current approaches to the construction of full-thickness skin substitutes therefore involves embedding dermal fibroblasts into 3D hydrogels and cultivating epidermal keratinocytes on the surface of hydrogels. Although these 3D co-culture systems have been used for many years for in vitro studies of epithelial-mesenchymal interactions,\(^{47}\) for keratinocyte differentiation,\(^{36}\) for the dynamics of the basement membrane\(^{38}\) and for many other applications, there are many limitations on clinical applications that still need to be overcome.

Although hydrogels have excellent biocompatibility and biodegradability properties for wound healing applications, they are not mechanically stable and their structure tends to be contracted under the traction forces of the embedded cells.\(^{27,48}\) The Young’s modulus of collagen hydrogels oscillates around units of kPa, depending on temperature, pH, duration of neutralization and many other conditions during polymerization.\(^{49,50}\) In order to keep the cells alive during polymerization of the collagen hydrogels, the conditions have to be physiological for the cells, eg 37 °C, pH~7.4. However, it has previously been observed that the collagen hydrogels formed under physiological conditions are less stiff than those polymerized at basic pH.\(^{49,51}\) However, the stiffness of hydrogel fibers plays an important role in cell behavior. Greater stiffness of the fibers suppressed cell adhesion, spreading, proliferation and migration into the hydrogel, due to a lower ability of the cells to transfer their traction forces to the hydrogel fibers.\(^{25,52}\)

Many approaches leading to improved hydrogel stability and improved mechanical properties have therefore recently been studied. Braziulis et al.\(^{21}\) used plastic
compression for a collagen type I hydrogel. Similarly, Kim et al\textsuperscript{53} induced structural and mechanical changes to the fibrin-collagen matrix by compressing it. Lotz et al\textsuperscript{48} reduced the fibroblast-mediated contraction of the collagen hydrogel by cross-linking it with succinimidyl glutarate polyethylene glycol. Another way to improve the mechanical properties of hydrogels is to reinforce them by means of biodegradable synthetic scaffolds.\textsuperscript{22} For example, Franco et al\textsuperscript{23} combined PCL/PLGA membranes with chitosan-gelatin hydrogels in an optimal ratio to achieve appropriate biocompatibility and mechanical properties for creating a two-layer construct of fibroblasts and keratinocytes. Hartmann-Fritsch et al\textsuperscript{31} incorporated nanofibrous PLGA membranes or knitted meshes into a bovine collagen I hydrogel with embedded fibroblasts and keratinocytes seeded on the surface. These authors obtained optimal stability of tissue-engineered full-thickness skin analogues by using a knitted mesh, where the epidermal part was well-stratified and the dermal part was revascularized in in vivo experiments. In the present study, we stabilized the collagen hydrogel by underlaying it with a nanofibrous fibrin-coated synthetic biodegradable PLLA membrane, pre-seeded with skin fibroblasts. Our results are in accordance with the recent studies which have shown that the Young’s modulus of collagen hydrogels formed in physiological conditions varied around units of kPa.\textsuperscript{49,51,52} The collagen hydrogel disks returned to their original state when compressed at low strain (up to 21%). As the scaffolds had a composite character, the mechanical response depended not only on the mechanical properties of the collagen hydrogel but also on the nanofibrous PLGA membrane. The results indicated that collagen hydrogel and PLLA membrane form a serial mechanical connection at compression in which the PLLA membrane start dominating at higher strains (over 56%). In addition, at higher strains, the collagen hydrogel undergoes structural changes. Pre-seeding the membrane with fibroblasts allowed the fibroblasts later to migrate into the collagen hydrogel without significant contraction of the hydrogel. The stability of the collagen hydrogel can be explained by the optimal ratio between the degradation processes and the remodeling processes mediated by the fibroblasts. During collagen polymerization, the fibroblasts adhering on the membrane were already in their proliferating phase, and they might not generate the strong traction forces that they generate while they are spreading. Moreover, after collagen polymerization, the cells could start migrating into the hydrogel by degrading it with matrix metalloproteases, and by synthesizing their own ECM proteins.\textsuperscript{16,17} In other words, we suppose that the traction forces generated by fibroblasts migrating from the PLLA membrane might be weaker than those generated by fibroblasts directly embedded into the collagen hydrogel. Direct embedding of the fibroblasts into the hydrogel during polymerization can be followed by cell spreading and by generating the intensive cell traction forces that are transmitted into the collagen fibrils. However, fibroblasts migrating from the membrane gradually degrade and remodel the collagen, which enables them to migrate spontaneously into the hydrogel, and leaves the collagen hydrogel relatively unchanged.

In order to increase the attractiveness of the PLLA membrane for the adhesion, spreading and proliferation of the fibroblasts pre-seeded on the membrane, we coated the membrane with a homogeneous nanofibrous fibrin mesh. In our previous study,\textsuperscript{10} we showed that a nanostructured fibrin mesh apparently enhanced fibroblast adhesion, proliferation and biosynthesis of ECM proteins, mainly collagen I and fibronectin. In this study, we have additionally found that the fibrin mesh significantly increased the migration of fibroblasts into the collagen hydrogel in comparison with the non-modified PLLA membrane. Similar results were obtained by Fu et al\textsuperscript{13} in experiments with fibrinogen-coated PCL nanofibers. They reported that coating PCL membranes with fibrinogen accelerated the migration of fibroblasts and stimulated their differentiation to myofibroblasts in a transforming growth factor (TGF)-\textbeta{} 1 rich microenvironment.\textsuperscript{13} In addition, due to the greater adhesion and proliferation of the cells caused by the fibrin mesh, a higher number of cells might migrate into collagen hydrogel. In contrast to our previous work,\textsuperscript{8} we observed that the fibrin mesh under the collagen gel was not degraded, even on day 14 after fibroblast seeding. This may have been caused by focusing the fibroblasts on migrating and on remodeling the 3D collagen hydrogel, rather than focusing on the degradation and remodeling of the fibrin mesh, which was described in detail in our previous studies.\textsuperscript{8–10}

In order to create a two-layer cellular skin construct, the collagen hydrogel enriched by spontaneously immigrated fibroblasts was seeded with primary dermal keratinocytes. A monolayer of adhered keratinocytes, with well-developed F-actin and basal cytokeratin 14 filaments, was formed on the whole surface of the collagen hydrogel after 7 days. The cells were able to divide for 14 days of cultivation, and they formed the basal layer with
highly mitotically active cells and a suprabasal layer with large cells frequently without cell nuclei. Our results correlated with other similar studies focused on the cultivation of keratinocytes on a collagen gel \cite{21,28,31,54} or on other forms of collagen substrates. \cite{9,36,55} The previous studies also reported that a PCL/collagen nanofibrous membrane alone did not stimulate keratinocyte migration, while subsequently coating a membrane composed of an ultrafine fibrous collagen network significantly increased the cell motility. \cite{14} Our results and other studies have proved that native collagen in the form of a hydrogel provides a physiological 3D microenvironment for the optimal co-cultivation of fibroblasts and keratinocytes. Furthermore, the high void ratio and the high permeability of collagen hydrogels enable paracrine communication between cells. \cite{46}

Conclusion
We have prepared a two-layer skin construct of fibroblasts and keratinocytes composed of a nanofibrous fibrin-coated PLLA membrane pre-seeded with human dermal fibroblasts, a collagen hydrogel and keratinocytes. The results have shown that the fibroblasts were able to migrate from the membrane upwards into the collagen hydrogel. The PLLA membrane underlying the collagen hydrogel reinforced the whole skin construct, and served as a substrate for the initial attachment, growth and subsequent spontaneous migration of fibroblasts. Moreover, coating the membrane with a nanofibrous fibrin mesh further considerably enhanced the migration of the fibroblasts into the hydrogel. The fibroblasts did not mediate a significant contraction of the collagen hydrogel during their migration, such as has been repeatedly observed when fibroblasts are directly embedded into a collagen hydrogel during the gelling process. The keratinocytes seeded on top of the collagen formed a homogeneous basal layer of proliferating cells. This two-layer skin construct based on a collagen hydrogel with spontaneously immigrated fibroblasts and reinforced by a fibrin-coated nanofibrous membrane seems to be promising for the treatment of extensive full-thickness skin wounds.

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**Supplementary materials**

**Video S1** The horizontal sections through the collagen hydrogel with keratinocytes adhered on the hydrogel surface and fibroblasts homogeneously immigrated inside the collagen hydrogel. The fibroblasts migrated from the membrane into the collagen hydrogel for a period of 11 days. Keratinocytes proliferated on the hydrogel for 7 days. Both cell types were stained with phalloidin–TRITC for the cell F-actin cytoskeleton (red), and with Hoechst #33258 for the cell nuclei (blue). The cytokeratin 14 in the keratinocytes was stained by immunofluorescence (Alexa 488, green). Dragonfly 503 spinning disk confocal microscope with a Zyla 4.2 PLUS sCMOS camera, objective HC PL APO 20x/0.75 IMM CORR CS2.

**Video S2** This video shows the two-layer construct of skin cells, depicted in Figure 10, in dynamic mode. The construct was composed of the fibrin-coated PLLA membrane seeded with human dermal fibroblasts, the collagen hydrogel with fibroblasts migrating inside the gel, and the keratinocyte layer on top of hydrogel. The fibroblasts migrated from the membrane into the collagen hydrogel for a period of 18 days. Keratinocytes proliferated on the hydrogel for 14 days. The PLLA membrane with fibroblasts and the collagen hydrogel with immigrated fibroblasts and keratinocytes on the top of hydrogel were stained separately. Both cell types were stained with phalloidin – TRITC for the cell F-actin cytoskeleton (red), and with Hoechst #33258 for the cell nuclei (blue). The cytokeratin 14 in the keratinocytes and the fibrin mesh on the nanofibrous membrane (at the bottom) were stained by immunofluorescence (Alexa 488, green). Dragonfly 503 spinning disk confocal microscope with a Zyla 4.2 PLUS sCMOS camera, objective HC PL APO 20x/0.75 IMM CORR CS2.