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Mechanical properties of polycaprolactone (PCL) scaffolds for hybrid 3D-bioprinting with alginate-gelatin hydrogel

Fritz Koch a,b,\*, Ole Thaden a, Stefan Conrad b, Kevin Trondle ¨ a, Günter Finkenzeller c, Roland Zengerle a,d, Sabrina Kartmann a,d, Stefan Zimmermann a, Peter Koltay a,b

a *Laboratory for MEMS Applications, IMTEK - Department of Microsystems Engineering, University of Freiburg, Georges-Koehler-Allee 103, D-79110, Freiburg, Germany* b *Freiburg Center for Interactive Materials and Bioinspired Technologies (FIT), University of Freiburg, Georges-Koehler-Allee 105, D-79110, Freiburg, Germany* c *Department of Plastic and Hand Surgery, Medical Center - University of Freiburg, Faculty of Medicine, University of Freiburg, Hugstetterstraße 55, D-79106, Freiburg, Germany*

d *Hahn-Schickard, Georges-Koehler-Allee 103, D-79110, Freiburg, Germany*

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**1. Introduction**

ABSTRACT

The generation of artificial human tissue by 3D-bioprinting has expanded significantly as a clinically relevant research topic in recent years. However, to produce a complex and viable tissue, in-depth biological under standing and advanced printing techniques are required with a high number of process parameters. Here, we systematically evaluate the process parameters relevant for a hybrid bioprinting process based on fused deposition modeling (FDM) of thermoplastic material and microextrusion of a cell-laden hydrogel. First, we investigated the effect of the printing temperature of polycaprolactone (PCL), on the junction strength between individual fused filaments and on the viability of immortalized mesenchymal stem cells (iMSC) in the sur rounding alginate-gelatin-hydrogel. It was found that a printing temperature of 140 ◦C and bonds with an angle of 90◦ between the filaments provided a good compromise between bonding strength of the filaments and the viability of the surrounding cells. Using these process parameters obtained from individual fused filaments, we then printed cubic test structures with a volume of 10 × 10 × 10 mm3 with different designs of infill patterns. The variations in mechanical strength of these cubes were measured for scaffolds made of PCL-only as well as for hydrogel-filled PCL scaffolds printed by alternating hybrid bioprinting of PCL and hydrogel, layer by layer. The bare scaffolds showed a compressive modulus of up to 6 MPa, close to human hard tissue, that decreased to about 4 MPa when PCL was printed together with hydrogel. The scaffold design suited best for hybrid printing was incubated with cell-laden hydrogel and showed no degradation of its mechanical strength for up to 28 days.

drop-on-demand bioprinting, where low viscous fluids are printed as

droplets (Binder et al., 2011; Troendle et al., 2021), a compromise of

3D-bioprinting has become a significant research area in the last years and shows great potential for future applications in regenerative medicine (Cui et al., 2017; Keriquel et al., 2010; Murphy and Atala, 2014) and in vitro models (Arslan-Yildiz et al., 2016; Mandrycky et al., 2016). For both applications, extrusion-based bioprinting is with 76% the most frequently reported bioprinting technology (Pedroza-Gonz´alez et al., 2021) to print cells and other biomaterials (Choudhury et al., 2018) (Ozbolat and Hospodiuk, 2016). Its low cost and ease of use are particularly advantageous compared to, for example, laser-based methods (Guillotin et al., 2010; Seol et al., 2014). Compared to

mechanically stable biomaterials is found that allows printing into the third dimension while ensuring sufficient cell viability (Heid and Boc caccini, 2020; Malda et al., 2013).

Although the processes and materials for extrusion-based bioprinting are becoming more and more advanced, the mechanical stabilities of hydrogels are still far from those of human hard tissue. One way to improve mechanical stability without reducing cell viability by increased hydrogel stiffness is to decouple the mechanical and biological requirements. This can be achieved by combining several materials in a multi-material bioprinting process referred to also as hybrid bioprinting

\* Corresponding author. Laboratory for MEMS Applications, IMTEK - Department of Microsystems Engineering, University of Freiburg, Georges-Koehler-Allee 103, D-79110, Freiburg, Germany.

*E-mail address:* fritz.koch@imtek.uni-freiburg.de (F. Koch).

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process. For this purpose, a stiff material that supports the mechanical load can be combined with a softer material that serves as an extracel lular matrix (Kang et al., 2016; Ruiz-Cantu et al., 2020).

Polycaprolactone (PCL) is a biodegradable polymer that has been extensively studied in the biomedical field due to its excellent me chanical stability and bio-compatibility (Endres et al., 2003; Schage mann et al., 2010). A broad use of PCL has already been reported for fabrication of scaffolds for a wide range of tissues, such as nerve (Ciar delli and Chiono, 2006; Woodruff and Hutmacher, 2010), cartilage (Alves da Silva, M L et al., 2010; Woodruff and Hutmacher, 2010) and, bone (Hutmacher et al., 2001; Ren et al., 2017; Woodruff and Hut macher, 2010). Hutmacher (Hutmacher et al., 2001) and Zein (Zein et al., 2002) were among the first groups to use fused deposition modeling (FDM) to generate porous PCL scaffolds, which can be incu bated with cells. They investigated the compression modulus and other mechanical properties of the scaffolds and demonstrated the great po tential of such PCL scaffolds for medical applications. Novel PCL-based composite materials loaded with organic or inorganic fillers can help to control mechanical properties and integrate further functionalities (Maietta et al., 2018). Such hybrid PCL-based scaffolds have been shown to be suitable for long-term in vitro cultivation of up to 4 weeks with high stability (Lee et al., 2010).

For 3D-objects thicker than a few millimeters, it can also be of great advantage to incorporate porosity in the printed scaffolds through intercalated microchannels made of cell-laden hydrogel and PCL. The remaining porosity allows for better diffusion in vitro and *in vivo* to reduce tissue necrosis (Kang et al., 2016). For printing PCL in combi

nation with cell-laden hydrogel, shear-thinning rheological properties of the hydrogel are desirable (Paxton et al., 2017). This allows the printing of high viscous hydrogels with low shear stress. Immediately after printing, the hydrogel should have maximum stability, and not disin

tegrate from its printed shape.

It has already been shown that compression moduli up to 6 MPa can be achieved (Schuurman et al., 2011). This is in the range of natural tissue, such as *cartilage with 4.*1 MPa (Woodfield et al., 2004) or *trachea with 3.*3 MPa (Trabelsi et al., 2010). The molecular weight and filament diameter of the PCL, the fiber orientation and the pore size were important factors to influence the mechanical properties of printed scaffolds. In general, an increase in porosity led to a decrease in elastic properties (Moroni et al., 2006).

In summary, multi-material bioprinting processes have already been used to fabricate tissue-like structures and demonstrated their added value for regenerative medicine. However, multi-material bioprinting requires the use of different printing technologies and biomaterials, leading to increasingly complex and costly printing systems and pro

cesses. To enable such processes, we have equipped in our previous work a low-cost open-source FDM 3D-printer with an additional syringe extruder for the alternate processing of thermoplastics and cell-laden hydrogels in a single process (Koch et al., 2021). However, the pro

cesses presented here can be implemented with any 3D bioprinter that has at least one printhead for printing thermoplastics and one printhead for extruding cell-laden hydrogels. The number of possible process variations can increase exponentially with the number of different ma

terials and printing methods. In order to vary only relevant parameters, a systematic investigation of the printing temperature as first parameter can be helpful, because the FDM extrusion temperature affects both: the bond strength between the filaments and the cell viability, alike. To date, the printing temperature in hybrid-bioprinting has not been sys

tematically investigated regarding its influence on bond strength of filament junctions and cell survival in the surrounding hydrogel. Mostly only successful printing designs have been presented in the literature (Kang et al., 2016; Schuurman et al., 2011) without showing how the printing temperature and other parameters were optimized.

Here, we present a systematic investigation of the printing temper ature of FDM-printed PCL on the mechanical compressibility and the cell viability of the surrounding cells. First, fracture forces in simple PCL test

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structures were investigated in order to determine suitable printing parameter settings. Based on these findings, four different infill patterns were printed and evaluated for their mechanical strength as stability promoting porous PCL scaffolds when printed as bare structures from PCL only and when combined with cell-laden alginate-gelatin hydrogel. The mechanically most promising design was then used to investigate the influence of hydrogel and incubation conditions on the mechanical stability of the printed part after up to 28 day of incubation. The final use case for such hybrid designs with high mechanical stability made from PCL and cell laden hydrogels might be in the field of fabrication of regenerative bone tissues as stability-promoting scaffolds or bone replacement implants.

**2. Materials and methods**

*2.1. Bioprinter and characterization*

A commercial 3D-printer Prusa i3 MK3 from Prusa Research A.S (Prague, Czech Republic) was used as a bioprinter platform. The FDM printer was upgraded with an extrusion printhead to allow alternate printing of hydrogel and thermoplastic material in a single process. The final printer design and the complete instructions for making, assem

bling and operating the printer are documented in an open access publication (Koch et al., 2021). In all experiments polycaprolactone (PCL) Facilian PCL100 from 3D4Makers B.V. (Haarlem, Netherlands) with 50,000 g/mol and 1.75 mm filament diameter was printed using a 0.4 mm nozzle by E3D-Online (Oxfordshire, UK). The fracture force of the bonds between the PCL filaments was tested by pulling the filaments with the bond tester Dage 4000 from Dage (Feldkirch, Germany). Compression modulus was measured with a compression test machine EZ-SX from Shimadzu Europe (Duisburg, Germany). Each sample was compressed with 1 mm/min loading speed until 30% strain to reach the plastic region. Due to plastic deformation, the measurements could only be performed once per sample. Compression modulus was defined as the slope of the linear region of the stress/strain curve. The linear region was in different ranges for the tested scaffolds, but always between 5% and 10% strain. For the hydrogel-only samples, which were significantly less stable due to the absence of PCL, the linear region was defined from 0% to 10%.

For visualization of the PCL scaffolds, PhotoView 360 from Solid works 2017 from Dassault Syst`emes (V´elizy-Villacoublay, France) was used. The GCode was generated with PrusaSlicer version 2.1.1 based on Slic3r from Prusa Research A.S (Prague, Czech Republic). The porosity of the PCL scaffolds was calculated with the following equation

porosity = (1 − *m*scaffold

*ρ*PCL ⋅ *V*scaffold) ⋅ 100% (1)

where *m*scaffold is the weighted mass of the scaffold, *ρ*PCL the constant density of PCL of 1.1 g/cm3 as specified by the manufacturer, and *V*scaffold the optically measured volume of the scaffold. The number of bonding errors was determined by visually inspecting at least 10 scaf

folds and manually counting the visible number of broken or missing bonds. Images for the optical volume determination were acquired using a digital microscope Dino-Lite Premier from AnMo Electronics (Hsinchu, Taiwan). For each scaffold design, at least three samples were measured at five representatively selected lengths along each dimension.

*2.2. Bioink*

The hydrogel was prepared by dissolving 4 w/v% alginate A1112- 100G and 16 w/v% gelatin 39465-500G both from Sigma-Aldrich (St. Louis, United States) in PBS each. Gelatin solution was heated up to 80 ◦C and alginate stirred for 24 h without heating to dissolve fully. Both stock solutions were mixed 1:1 to result in a hydrogel with 2 w/v% alginate and 8 w/v% gelatin. The bioink was filled in 2 ml medical

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syringes from BBraun (Melsungen, Germany) and printed with conical dispensing nozzles from Globaco GmbH (Rodermark, ¨ Germany) with an inner diameter of 410 μm. The hydrogel was cross-linked by incubating for 30 s in a 10% CaCl2 (223506) solution from Sigma-Aldrich (St. Louis, United States) dissolved in water. A cell-laden hydrogel was prepared by gently mixing cells in a concentration of 500,000 cells/ml into the 37 ◦C warm hydrogel.

*2.3. Cell culture*

Immortalized mesenchymal stem cells (iMSCs) were kindly provided by Prof. Dr. Matthias Schieker from the Laboratory of Experimental Surgery and Regenerative Medicine of the Ludwig-Maximilians University of Munich. They were immortalized by lentiviral gene transfer of human telomerase reverse transcriptase (Bocker ¨ et al., 2008). The cells were cultivated in MEM alpha Medium including 10% FBS both from Gibco (Carlsbad, USA) at 37 ◦C and 5% CO2.

*2.4. Live/dead assay*

Cell viability was determined with a viability/cytotoxicity kit for mammalian cells (Invitrogen L3224) from Fisher Scientific GmbH (Schwerte, Germany). Cells were stained as recommended in the pro tocol for 30 min and imaged with a fluorescence microscope Observer Z1 microscope from Carl Zeiss (Oberkochen, Germany) with 470 nm and 555 nm excitation wavelength. Image analysis was performed with ImageJ (V1.52). Since the cells were due to extrusion printing randomly distributed in the hydrogel rather than in a single focal plane, an in tensity threshold was defined in each image. Values higher than 70% of the maximum intensity were defined as valid live or dead fluorescence signals, as described in detail in a previous publication (Koch et al., 2020). All quantitative data were assessed for statistical significance by

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performing an ANOVA with a Tukey post-hoc test. A statistically sig nificant test result was considered for *p <* 0.05.

**3. Results and discussion**

First, the hybrid bioprinter was characterized for its properties using only the FDM-printhead. For the subsequent hybrid printing of porous structures, which may contain cell-laden material, it was of general in terest to determine how the PCL printing temperature influences the mechanical stability of the bond between single fused filaments and the viability of the surrounding cells. Therefore, a test structure was designed with a single filament printed at an angle of 90◦ to the previous layers. The filament bridged a distance of 2 mm between the two pre vious layers. Detailed images of the test structure are shown in the supplemental Fig. A1. The printing temperature was gradually increased by 10 ◦C between 100 ◦C and 180 ◦C. For each temperature at least 44 samples were investigated by pulling up the filament in the pull tester until failure and measuring the fracture force. Supplemental Fig. A2 il lustrates the pull tester with a test structure and the two failure mech anisms that occur (bond and filament break).

Fig. 1a shows the mean fracture force obtained from these pull tests as a function of the temperature. The fracture force increases from around 2.5 N to around 4.5 N for increasing printing temperature from 100 ◦C to 140 ◦C. For temperatures above 140 ◦C, no significant change in fracture force was measured. This may result from a changed failure mechanism, as shown in Fig. 1b. For increasing temperature, the failure mechanism changed from breakage of the bond between the two fila

ments (bond break) to breakage of the filaments itself (filament break). An increasing number of filament breaks indicates that the fracture force has increased close to the maximum fracture force given by the material properties (see also supplemental Fig. A3 in which the bond break and filament break were considered individually).

**Fig. 1.** Influence of nozzle temperature and raster angle on the mechanical stability of fused filaments and cell viability during FDM printing of PCL. Fracture force was measured by pulling on a bridging filament connected to two filaments below. The temperature was varied between 100 ◦C and 180 ◦C (a) Failure was analyzed and sorted into two categories, either the separation of two layers on joints (bond break) or the breakage of the filament itself (filament break) (b). Fracture force was measured for angles of 90◦, 45◦, and 30 ◦between the fused filaments (c). Cell viability was determined for cells embedded into a hydrogel of alginate and gelatin and injected below the printed filament (d).

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In addition to the influence of the printing temperature, the fila ment’s angle on the contact points was also investigated. This raster angle between the filaments in contact to each other was varied from 90◦ to 45◦ and 30◦. The resulting fracture force was measured for 50 fused filaments for each angle in the range from 130 ◦C to 150 ◦C and is shown in Fig. 1c. It can be seen that the fracture force significantly decreases with reduced raster angle. This may be due to a smaller area interacting with shear forces when the raster angle is reduced. Thus, all junctions between fused filaments printed with angles unequal to 90◦ should result in a lower bond strength. This finding is also in line with most published experiments on similar test structures in 3D-bioprinting (Hutmacher et al., 2001; Zein et al., 2002). Based on the measure ments shown in Fig. 1c, only a raster angle of 90◦ was used for the subsequent experiments.

The printing temperature does not only affect the cohesive strength between the filaments but also influences the viability of the sur rounding cells. To achieve a high mechanical stability between the fil aments on the one hand, the printing temperature should be selected as high as possible up to 150 ◦C. At temperatures between 150 ◦C and 180 ◦C, a plateau is reached and the fracture force does not increase any further, as shown in Fig. 1a. On the other hand, the printing temperature should generally be as low as possible for the printing process with living cells, because the printed PCL filaments will heat up the surrounding hydrogel and thus might damage the cells. In the experiments carried out to investigate this effect, PCL filaments were printed in straight lines at a federate of 10 mm/s directly onto an approximately 1 mm thick crosslinked cell-laden hydrogel with different nozzle temperatures. Immortalized mesenchymal stem cells (iMSCs) were encapsulated in a hydrogel of 2% alginate and 8% gelatin in a concentration of 500,000 cells/ml. Cell viability was assessed with a live/dead assay at ten regions of interest and is shown for different nozzle temperatures in Fig. 1d. The viability remains between 60% and 70% for increasing temperature up to 140 ◦C and decreases for higher temperatures to viabilities down to 40%. The viability in control samples without contact to heated PCL filaments had a viability of 72% ± 17%.

From these measurements of the fracture force and the cell viability both being influenced by the printing temperature, it was decided for all further experiments to choose a printing temperature of 140 ◦C and print only connections with a raster angle of 90◦ between the fused filaments to achieve highest stability and viability. Beyond a temperature of 140 ◦C, an increase in printing temperature would not result in a further increase in fracture force and a reduction of printing temperature would not result in a significant increase in cell viability.

In addition to the influence of the individual bonds between fused filaments on the mechanical stability, the print design plays a role in the structure’s overall mechanical stability. Based on the results related to

**Table 1**

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temperature and raster angle, four different scaffold designs were printed at 140 ◦C and 90◦ raster angle between fused filaments. The four scaffold designs, shown in Table 1, had a similar mass and dimensions of 10 × 10 × 10 mm3. Therefore, the theoretical porosity was between 71.5% and 71.6% and a slightly higher porosity between 74.0% and 75.9% was calculated according to equation (1) based on experimental weighing results. The deviation of these values from the theoretical measured porosity may be due to a systematic loss of mass during printing, a discrepancy in the density specification of the PCL from the manufacturer, deviations from the ideal cubic volume or a combination of these. The designs were chosen so that the porosity is as similar as possible but with varied maximum pore sizes. Scaffold 1 was chosen as the most straightforward print design with alternating filaments in x and y-direction, forming 90◦ junctions with the following layer. For each layer, seven lines are printed as a meander. The pore sizes are uniform and with 0.53 mm3, the smallest of the four scaffolds. In scaffolds 2 and 3, gradients in the x- and y-positions of the filaments were created, resulting in bigger maximum pore sizes in the center of the scaffolds of 1.52 mm3 and 2.92 mm3, respectively. This is in reference to natural tissue where, for example in bones, most of the mechanical load is absorbed by the outer walls of the structure and therefore allows the center to be more porous and suitable for vascularization. The gradient increases from scaffold 2 to scaffold 3, so that scaffold 3 has a solid outer wall. Scaffold 4 is identical to scaffold 1 but has twice the maximum pore size of 1.06 mm3 because double layers with the same orientation are printed on top of each other.

These four scaffolds were tested for their mechanical stability with a compression test. The compression modulus was determined as the linear region in the stress/strain diagram for a compression speed of 1 mm/s. It can be seen in Fig. 2a, that the mechanical stability of the scaffolds barely differs despite the different maximum pore sizes. All scaffolds had an average compression modulus of about 6 MPa for ten tested scaffolds each. A significant difference could only be observed between scaffold 3 and scaffold 4. Although the scaffolds were of similar porosity and mechanical stability, the occurrence of errors during the printing process was different. Errors were defined as not fused fila

ments in the z-direction. If this junction was not bonded, it could not support any load and was easily displaced in x- and y-direction. The total number of bonding errors was counted for each scaffold and shown in Fig. 2b. Since scaffold 2 had a significantly higher number of errors than scaffold 1 and scaffold 4, it was not considered for further experiments with cells. The design of scaffold 3 would prevent diffusion of nutrients during in vitro cultivation due to the solid outer walls and as it exhibited no superior compressive modulus than scaffolds 1 and 4 it was also excluded from further experiments. (Exemplary images of the bonding errors are shown in supplemental Fig. A4 for each scaffold).

Overview of various PCL scaffold designs used for mechanical testing. Scaffolds 1–4 were designed to have similar porosity but different maximum pore sizes. The pore sizes are increased in scaffold 2 and 3 by creating a gradient of horizontally aligned layers from the center to the outside of the structures, ending in a closed solid wall surrounding scaffold 3. The pore sizes are increased in scaffold 4 by using the same filament spacing like in scaffold 1, but printing two identical layers on top of each other each time.

Scaffold 1 Scaffold 2 Scaffold 3 Scaffold 4

**Porosity**

Theoretical 71.6% 71.5% 71.6% 71.6% Measured (n = 10) 74.2% ± 0.2% 74.0% ± 0.4% 75.9% ± 0.5% 74.7% ± 0.3%

**Maximum pore size**

Theoretical 0.53 mm3 1.52 mm3 2.92 mm3 1.06 mm3  Measured (n = 10) 0.53 mm3 ± 0.08 mm3 1.65 mm3 ± 0.23 mm3 2.73 mm3 ± 0.28 mm3 1.10 mm3 ± 0.03 mm3

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**Fig. 2.** Compression moduli for different scaffold designs (1–4) measured under constant compression with 1 mm/s (n = 10) determined in the z-direction (a). Scaffold 2 was excluded from the following experiments due to the high number and variation of errors (b), and scaffold 3 due to the design not suitable for in vitro incubation. Scaffolds 1 and 4 were investigated in detail by also measuring the compression modulus in x- and y-direction (c) and the increase in porosity due to the reduction of filaments aligned against compression from 84 down to 36 (n = 3) (d).

Scaffold 1 and scaffold 4 were investigated further by measurements of their compression modulus in the x- and y-direction, as shown in Fig. 2c. The compression modulus in the x- and y-direction was compared to the values in the z-direction, as already shown in Fig. 2a. Both scaffolds showed significantly higher compression moduli in the x

and y-direction than in the z-direction, increasing from about 6 MPa to about 8 MPa. Furthermore, the influence of the total number of fila ments in the x- and y-direction was investigated by experiments. For this purpose, the distance between the filaments in x- and y-direction of scaffolds 1 and 4 was increased, and thus, the total number of filaments was reduced from 84 to 60 and to 36 in both designs. This corresponds to a reduction of filaments perpendicular to the compression direction

from 7 to 5 or 3 for each layer. The resulting compression moduli are shown in Fig. 2d. It can be seen that the compression modulus decreases linearly with a reduced number of filaments. At the same time, the porosity increases due to the lower number of filaments. Such mea

surements allow to forecast the mechanical stability of a specific design and therefore to customize the scaffolds based on porosity and me chanical requirements.

To study the interaction of the PCL printing process with the micro extrusion of hydrogel, the two scaffolds 1 and 4, previously rated with the highest compression moduli, were printed at a nozzle temperature of 140 ◦C and in the same run hydrogel filaments were printed layer by layer alternating to the PCL extrusion. For this alternating, hybrid

**Fig. 3.** a) Compression moduli of scaffolds 1 and 4 printed from PCL only (gray) and the same designs printed as hybrid constructs by combination of PCL and hydrogel (blue) determined directly after printing. For comparison pure hydrogel structures with zero porosity are also shown (red). B) Compression modules of hybrid scaffolds containing a total amount of 250 mg of hydrogel and exhibiting a porosity of about 50%. The hydrogel contained 500,000 cells/ml and the samples were incubated in vitro for up to 28 days. Compression moduli were measured from two samples each on days 0, 7, 14, and 28. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

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printing process the same volume of hydrogel as for the PCL was printed between the PCL filaments in each layer. Due to the additionally added hydrogel, the total porosity of the printed object decreased from about 75% to about 50%. In Fig. 3a, the compression modulus of these hybrid structures (blue) was compared with the pure PCL scaffolds (gray). As further reference, a pure hydrogel cube with the same edge length as the other scaffolds was printed. The results of the pure hydrogel samples are shown for comparison (red), as well. Despite the maximum filling of the hydrogel cube (i.e. zero porosity), the hybrid PCL scaffolds are about 150 times more stable than the pure hydrogel cube.

It can be seen for both scaffold designs that mechanical stability decreases with the addition of hydrogel, despite the fact that the total number of PCL filaments is not reduced. The most likely explanation for the smaller compression moduli of the hybrid scaffolds might be the reduced bonding strength between fused PCL filaments caused by wet

ting with hydrogel that leaks out from the hydrogel filaments. For scaffold 4, this reduction in compression modulus is lower than for scaffold 1. Since both hydrogel and PCL were printed with two identical layers on top of each other, the printing process appears to be less sensitive a reduction of the stability due to hybrid printing. Therefore, scaffold 4 was used in the following for the printing experiment with living cells suspended in the hydrogel.

To investigate potential long-term degradation of PCL under real cell culture conditions, hybrid structures of scaffold 4 were printed with alginate-gelatin hydrogel laden with 500,000 iMSCs/ml. Structures were incubated in cell culture medium, which was changed every three days. Since cell viability is highly dependent on many parameters beyond the used print design, which was the focus of this investigation, cell viability was not systematically quantified within the experiment. However, cell proliferation observed on the bottom of the plates indi

cated unrestricted cell viability in the presence of the PCL scaffolds throughout the whole experiment. Fig. 3b shows the compression modulus of two hybrid scaffolds after 0, 7, 14, and 28 days of incubation. Over the period of 28 days, only slight differences in the range of the previously determined standard deviation could be seen. Therefore, it can be concluded that the compression moduli of PCL scaffolds deter

mined previously are valid for up to 28 days of in vitro cultivation. Thus, hybrid printing of bioinks and porous PCL scaffolds can significantly increase the mechanical stability during an incubation period of at least up to four weeks. The increase of the compression modulus from the kPa to the MPa range is a significant improvement compared to the pure hydrogel-based bioink, which can be beneficial for *in vivo* applications of hard human tissues.

**4. Conclusions**

In this work, process parameters and printing patterns for hybrid 3D bioprinting of PCL and a micro extruded alginate-gelatin-hydrogel were systematically investigated. It was shown that design parameters and printing parameters can be adjusted in view of a defined mechanical stability of the resulting 3D-bioprinted object. First, a systematic study showed the influence of PCL printing temperature on bond strength between PCL filaments and the viability of surrounding cells subjected to thermal stress by the PCL deposition. Second, four different printing designs were investigated by varying infill patterns. The designs were printed with a similar porosity, evaluated for their mechanical stability, and tested for hybrid printing. With the knowledge gained from the study, the final mechanical stability of the printed scaffolds can be predicted and adjusted by the chosen designs as a function of the number of printed PCL filaments. The mechanically most stable design was tested for its long-term stability as a hybrid print construct with a cell-laden hydrogel by incubation for up to 28 days. During this time no significant change in the mechanical stability was detected.

The hybrid bioprinting process described in this work allows to achieve high mechanical stability, similar to human hard tissues, simultaneously with high cell viability in a single 3D-bioprinted object.

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Compared to the use of a one single bioink only, these properties are achieved by combining porous PCL structures with a hydrogel-based bioink, leading to a decoupling of the mechanical and cellular re quirements. But, also for the hybrid printing approach, a compromise between mechanical stability and cell viability must be made, which may differ for each material and application. The printing temperature of 140 ◦C and angles of 90◦ between filaments described here as most stable configuration for printing PCL, while still maintaining a high cell viability, may help for process optimization of other 3D-bioprinting processes, designs, cells and materials.

**CRediT authorship contribution statement**

**Fritz Koch:** Visualization, Validation, Methodology, Investigation, Formal analysis. **Ole Thaden:** Formal analysis, Investigation, Visuali zation. **Stefan Conrad:** Software, Resources, Investigation, Conceptu alization. **Kevin Trondle:** ¨ Resources, Validation, Writing – review & editing. **Günter Finkenzeller:** Validation, Resources, Funding acquisi tion. **Roland Zengerle:** Funding acquisition, Supervision, Writing – review & editing. **Sabrina Kartmann:** Writing – review & editing, Su pervision, Funding acquisition, Conceptualization. **Stefan Zimmer mann:** Project administration, Resources, Writing – review & editing. **Peter Koltay:** Writing – review & editing, Supervision, Formal analysis, Data curation.

**Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Appendix A. Supplementary data**

Supplementary data to this article can be found online at https://doi. org/10.1016/j.jmbbm.2022.105219.

**References**

Alves da Silva, M.L., Martins, A., Costa-Pinto, A.R., Costa, P., Faria, S., Gomes, M., Reis, R.L., Neves, N.M., 2010. Cartilage tissue engineering using electrospun PCL nanofiber meshes and MSCs. Biomacromolecules 11 (12), 3228–3236. https://doi. org/10.1021/bm100476r.

Arslan-Yildiz, A., El Assal, R., Chen, P., Guven, S., Inci, F., Demirci, U., 2016. Towards artificial tissue models: past, present, and future of 3D bioprinting. Biofabrication 8 (1), 14103. https://doi.org/10.1088/1758-5090/8/1/014103.

Binder, K.W., Allen, A.J., Yoo, J.J., Atala, A., 2011. Drop-on-demand inkjet bioprinting: a primer. Gene Ther. Regul. 6 (1), 33–49. https://doi.org/10.1142/

S1568558611000258.

Bocker, ¨ W., Yin, Z., Drosse, I., Haasters, F., Rossmann, O., Wierer, M., Popov, C., Locher, M., Mutschler, W., Docheva, D., Schieker, M., 2008. Introducing a single cell-derived human mesenchymal stem cell line expressing hTERT after lentiviral gene transfer. J. Cell Mol. Med. 12 (4), 1347–1359. https://doi.org/10.1111/j.1582- 4934.2008.00299.x.

Choudhury, D., Anand, S., Naing, M.W., 2018. The arrival of commercial bioprinters - towards 3D bioprinting revolution. Int. J. Bioprint. 4 (2), 139. https://doi.org/ 10.18063/ijb.v4i2.139.

Ciardelli, G., Chiono, V., 2006. Materials for peripheral nerve regeneration. Macromol. Biosci. 6 (1), 13–26. https://doi.org/10.1002/mabi.200500151.

Cui, H., Nowicki, M., Fisher, J.P., Zhang, L.G., 2017. 3D bioprinting for organ regeneration. Adv. Healthc. Mater. 6 (1) https://doi.org/10.1002/adhm.201601118.

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Endres, M., Hutmacher, D.W., Salgado, A.J., Kaps, C., Ringe, J., Reis, R.L., Sittinger, M., Brandwood, A., Schantz, J.T., 2003. Osteogenic induction of human bone marrow derived mesenchymal progenitor cells in novel synthetic polymer-hydrogel matrices. Tissue Eng. 9 (4), 689–702. https://doi.org/10.1089/107632703768247386.

Guillotin, B., Souquet, A., Catros, S., Duocastella, M., Pippenger, B., Bellance, S., Bareille, R., R´emy, M., Bordenave, L., Am´ed´ee, J., Guillemot, F., 2010. Laser assisted bioprinting of engineered tissue with high cell density and microscale organization. Biomaterials 31 (28), 7250–7256. https://doi.org/10.1016/j.

biomaterials.2010.05.055.

Heid, S., Boccaccini, A.R., 2020. Advancing bioinks for 3D bioprinting using reactive fillers: a review. Acta Biomater. 113, 1–22. https://doi.org/10.1016/j. actbio.2020.06.040.

Hutmacher, D.W., Schantz, T., Zein, I., Ng, K.W., Teoh, S.H., Tan, K.C., 2001. Mechanical properties and cell cultural response of polycaprolactone scaffolds designed and fabricated via fused deposition modeling. J. Biomed. Mater. Res. 55 (2), 203–216. https://doi.org/10.1002/1097-4636(200105)55, 2*<*203:AID-JBM1007*>*3.0.CO;2- 7.

Kang, H.W., Lee, S.J., Ko, I.K., Kengla, C., Yoo, J.J., Atala, A., 2016. A 3D bioprinting system to produce human-scale tissue constructs with structural integrity. Nat. Biotechnol. 34 (3), 312–319. https://doi.org/10.1038/nbt.3413.

Keriquel, V., Guillemot, F., Arnault, I., Guillotin, B., Miraux, S., Am´ed´ee, J., Fricain, J.-C., Catros, S., 2010. In vivo bioprinting for computer- and robotic-assisted medical intervention: preliminary study in mice. Biofabrication 2 (1), 14101. https://doi. org/10.1088/1758-5082/2/1/014101.

Koch, F., Thaden, O., Trondle, ¨ K., Zengerle, R., Zimmermann, S., Koltay, P., 2021. Open source hybrid 3D-bioprinter for simultaneous printing of thermoplastics and hydrogels. HardwareX, e00230. https://doi.org/10.1016/j.ohx.2021.e00230.

Koch, F., Trondle, ¨ K., Finkenzeller, G., Zengerle, R., Zimmermann, S., Koltay, P., 2020. Generic method of printing window adjustment for extrusion-based 3D-bioprinting to maintain high viability of mesenchymal stem cells in an alginate-gelatin hydrogel. Bioprinting 20, e00094. https://doi.org/10.1016/j.bprint.2020.e00094.

Lee, J., Guarino, V., Gloria, A., Ambrosio, L., Tae, G., Kim, Y.H., Jung, Y., Kim, S.-H., Kim, S.H., 2010. Regeneration of Achilles’ tendon: the role of dynamic stimulation for enhanced cell proliferation and mechanical properties. Journal of biomaterials science. Polym. Ed. 21 (8–9), 1173–1190. https://doi.org/10.1163/ 092050609X12471222313524.

Maietta, S., Russo, T., Santis, R. de, Ronca, D., Riccardi, F., Catauro, M., Martorelli, M., Gloria, A., 2018. Further theoretical insight into the mechanical properties of polycaprolactone loaded with organic-inorganic hybrid fillers. Materials 11 (2). https://doi.org/10.3390/ma11020312.

Malda, J., Visser, J., Melchels, F.P., Jüngst, T., Hennink, W.E., Dhert, W.J.A., Groll, J., Hutmacher, D.W., 2013. 25th Anniversary Article: Engineering Hydrogels for Biofabrication. Advanced Materials (Deerfield Beach, Fla.), vol. 25, pp. 5011–5028. https://doi.org/10.1002/adma.201302042, 36.

Mandrycky, C., Wang, Z., Kim, K., Kim, D.-H., 2016. 3D bioprinting for engineering complex tissues. Biotechnol. Adv. 34 (4), 422–434. https://doi.org/10.1016/j. biotechadv.2015.12.011.

Moroni, L., Wijn, J.R. de, van Blitterswijk, C.A., 2006. 3D fiber-deposited scaffolds for tissue engineering: influence of pores geometry and architecture on dynamic mechanical properties. Biomaterials 27 (7), 974–985. https://doi.org/10.1016/j. biomaterials.2005.07.023.

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Murphy, S.V., Atala, A., 2014. 3D bioprinting of tissues and organs. Nat. Biotechnol. 32 (8), 773–785. https://doi.org/10.1038/nbt.2958.

Ozbolat, I.T., Hospodiuk, M., 2016. Current advances and future perspectives in extrusion-based bioprinting. Biomaterials 76, 321–343. https://doi.org/10.1016/j. biomaterials.2015.10.076.

Paxton, N., Smolan, W., Bock, ¨ T., Melchels, F., Groll, J., Jungst, T., 2017. Proposal to assess printability of bioinks for extrusion-based bioprinting and evaluation of rheological properties governing bioprintability. Biofabrication 9 (4), 44107. https://doi.org/10.1088/1758-5090/aa8dd8.

Pedroza-Gonzalez, ´ S.C., Rodriguez-Salvador, M., P´erez-Benítez, B.E., Alvarez, M.M., Santiago, G.T., 2021. Bioinks for 3D bioprinting: a scientometric analysis of two decades of progress. Int. J. Bioprint. 7 (2), 333. https://doi.org/10.18063/ijb. v7i2.337.

Ren, K., Wang, Y., Sun, T., Yue, W., Zhang, H., 2017. Electrospun PCL/gelatin composite nanofiber structures for effective guided bone regeneration membranes. Mater. Sci. & Eng. C, Mater. Biol. Appl. 78, 324–332. https://doi.org/10.1016/j. msec.2017.04.084.

Ruiz-Cantu, L., Gleadall, A., Faris, C., Segal, J., Shakesheff, K., Yang, J., 2020. Multi material 3D bioprinting of porous constructs for cartilage regeneration. Mater. Sci. & Eng. C, Mater. Biol. Appl. 109, 110578 https://doi.org/10.1016/j. msec.2019.110578.

Schagemann, J.C., Chung, H.W., Mrosek, E.H., Stone, J.J., Fitzsimmons, J.S., O’Driscoll, S.W., Reinholz, G.G., 2010. Poly-epsilon-caprolactone/gel hybrid scaffolds for cartilage tissue engineering. J. Biomed. Mater. Res. Part A 93 (2), 454–463. https://doi.org/10.1002/jbm.a.32521.

Schuurman, W., Khristov, V., Pot, M.W., van Weeren, P.R., Dhert, W.J.A., Malda, J., 2011. Bioprinting of hybrid tissue constructs with tailorable mechanical properties. Biofabrication 3 (2), 21001. https://doi.org/10.1088/1758-5082/3/2/021001.

Seol, Y.-J., Kang, H.-W., Lee, S.J., Atala, A., Yoo, J.J., 2014. Bioprinting technology and its applications. Eur. J. Cardio. Thorac. Surg. : Off. J. Eur. Assoc. Cardio-Thorac. Surg. 46 (3), 342–348. https://doi.org/10.1093/ejcts/ezu148.

Trabelsi, O., del Palomar, A.P., Lopez-Villalobos, ´ J.L., Ginel, A., Doblar´e, M., 2010. Experimental characterization and constitutive modeling of the mechanical behavior of the human trachea. Med. Eng. Phys. 32 (1), 76–82. https://doi.org/10.1016/j. medengphy.2009.10.010.

Troendle, K., Rizzo, L., Pichler, R., Koch, F., Itani, A., Zengerle, R., Lienkamp, S.S., Koltay, P., Zimmermann, S., 2021. Scalable fabrication of renal spheroids and nephron-like tubules by bioprinting and controlled self-assembly of epithelial cells. Biofabrication. https://doi.org/10.1088/1758-5090/abe185.

Woodfield, T.B.F., Malda, J., Wijn, J. de, P´eters, F., Riesle, J., van Blitterswijk, C.A., 2004. Design of porous scaffolds for cartilage tissue engineering using a three dimensional fiber-deposition technique. Biomaterials 25 (18), 4149–4161. https:// doi.org/10.1016/j.biomaterials.2003.10.056.

Woodruff, M.A., Hutmacher, D.W., 2010. The return of a forgotten polymer—polycaprolactone in the 21st century. Prog. Polym. Sci. 35 (10), 1217–1256. https://doi.org/10.1016/j.progpolymsci.2010.04.002.

Zein, I., Hutmacher, D.W., Tan, K.C., Teoh, S.H., 2002. Fused deposition modeling of novel scaffold architectures for tissue engineering applications. Biomaterials 23 (4), 1169–1185. https://doi.org/10.1016/s0142-9612(01)00232-0.

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