Shape-defined solid micro-objects from poly(D,L-lactic acid) as cell-supported counterparts in bottom-up tissue engineering

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ABSTRACT

In bottom-up tissue engineering, small modular units of cells and biomaterials are assembled toward larger and more complex ones. In conjunction with a new implementation of this approach, a novel method to fabricate microscale objects from biopolymers by thermal imprinting on water-soluble sacrificial layers is presented. By this means, geometrically well-defined objects could be obtained without involving toxic agents in the form of photoinitiators. The micro-objects were used as cell-adhesive substrates and cell spacers in engineered tissues created by cell-guided assembly of the objects. Such constructs can be applied both for in vitro studies and clinical treatments. Clinically relevantly sized aggregates comprised of cells and micro-objects retained their viability up to 2 weeks of culture. The aggregation behavior of cells and objects showed to depend on the type and number of cells applied. To demonstrate the micro-objects' potential for engineering vascularized tissues, small aggregates of human bone marrow stromal cells (hMSCs) and micro-objects were coated with a layer of human umbilical vein endothelial cells (HUVECs) and fused into larger tissue constructs, resulting in HUVEC-rich regions at the aggregates' interfaces. This three-dimensional network-type spatial cellular organization could foster the establishment of (premature) vascular structures as a vital prerequisite of, for example, bottom-up-engineered bone-like tissue.

1. Introduction

Tissue engineering (TE) typically makes use of a combination of porous scaffolds from biomaterials of synthetic or biological origin, growth factors and the patient's own cells to ultimately repair or replace diseased, damaged, or lost tissue [1–3]. The scaffolds provide an environment for cells to adhere and proliferate, which at the same time provides mechanical stability, particularly in hard-tissue replacement, and supports the cells to ultimately gain and retain the desired function.

Engineering tissues in vitro can be subdivided into the classical top-down and the more recent bottom-up TE [4–6]. In top-down TE, a tissue construct is engineered with respect to, among others, size and shape of the tissue to be restored. These tissue constructs generally comprise a correspondingly large, one-piece porous biomaterial serving as a scaffold when adding cells. A disadvantage when using the top-down approach lies in the often obtained inhomogeneous cell distribution throughout the scaffold [7]. Furthermore, most top-down approaches do not allow for a high degree of tissue complexity, for example, by applying multiple cell types while controlling their spatial organization, and also not for an early dynamic tissue remodeling before the single-piece, static scaffold is degrading. Moreover, top-down TE typically requires invasive procedures to be able to implant the reconstructed tissue in the defect site in vivo.

In contrast, traditional bottom-up TE overcomes some of these problems by, for example, embedding cells in a hydrogel in which they can be distributed homogeneously when mixing them with the...
corresponding precursor before gelation and which can enable much earlier remodeling (Fig. 1A) [8–12]. Griffin et al. [13] have reported a bottom-up method in which cells are not encapsulated inside hydrogels but reside in between self-assembled hydrogel-based spheres. In that study, the cells show a homogenous distribution between the spheres and remain highly accessible for nutrient and gas supply and metabolic waste exchange in a wound healing application. Yet, hydrogels have a limited mechanical stability and typically a high degradation rate, making them inapplicable for load-bearing applications, such as in bone TE, or as long-term cell support. To overcome this well-known limitation in the mechanical properties of hydrogels, already before the aforementioned study, we introduced a method to create cell-assemblable micro-objects from the relatively stiff, epoxy-based polymer SU-8 [14]. Although SU-8 has a documented biocompatibility [15,16], the fact that the objects were made by photolithography poses the risk of toxic traces [17], in the form of free radicals, which might (unnoticedly) bias readouts in applications made by photolithography poses the risk of toxic traces [17], in the form of free radicals, which might (unnoticedly) bias readouts in applications.

Therefore, we developed a novel, radical-free method based on a combination of hot embossing [18], or thermal imprinting, and reactive ion etching (RIE), known as (nano)imprint lithography (NIL) [19,20]. The imprinting was performed on a poly(D,L-lactic acid) (PDLLA) film placed on a water-soluble sacrificial layer, to create free- or isolated standing and releasable engineered micro-objects. Medical-grade PDLLA was chosen as an exemplary or model thermoplastic biopolymer in conjunction with its comparatively low material costs, good thermal processability, and suitable, for example, mechanical properties and clinical relevance in the already mentioned field of bone TE [21,22].

In this potential main area for clinical application, the engineered micro-objects could, for example, be a component of an injectable formulation to treat or fill critical-size bone defects [14]. But also in other TE fields, for example, where conventional scaffolds in the form of porous biomaterials are researched so far, micro-objects from similar materials can, with the cells and the extracellular matrix (ECM) deposited by them acting as a binder for the objects, provide comparable material-based initial mechanical support or stability to the engineered tissues. At the same time, in contrast to the static macroscale scaffolds, the microscale objects – as dynamic scaffolding entities that can be moved by the cells – allow immediate local remodeling of the cell-material construct and might even give it more flexible and adaptive biomechanical properties as a whole. In a previous study, the bottom-up TE of tubular structures scaffolded by micro-objects could be demonstrated [14], as they could similarly represent a potential preliminary stage of a future tissue-engineered trachea or ureter. Another application field could be the use of the objects as advanced cell expansion carriers, particularly when provided with engineered cell-instructive surfaces, for example, by micro- or nanoimprinting, as a result of a previous screen of such surfaces condensed on a miniaturized library created by means of the same imprint technology [23]. Last but not least, as already suggested, the micro-objects could be used in 3D in vitro research models, as similarly demonstrated in the present study. When doing so, the objects could, for example, bring the powerful instructive potential of micro- and nanoengineered surfaces [24,25] to the third dimension when after the objects' highly defined creation, including such engineered surfaces, on wafer-type 2D technical substrates releasing the objects from the substrates and aggregating them with cells in a full 3D fashion.

The PDLLA objects were fabricated in three different shapes as a mean to vary the amount of material per overall volume. The objects were used as cell-adherent substrates and cell spacers in engineered tissues created by cell-guided assembly (Fig. 1B). These constructs can then be
aggregated into (geometric) modular tissues, which in turn can be further applied as building blocks to construct larger and more complex, random or organized assemblies. In such aggregates or assemblies, the micro-objects are expected to avoid (too) high cell or tissue contraction as observed in cells-only bottom-up TE approaches [26–28]. This contraction otherwise may hamper integration with surrounding tissues after implantation.

2. Materials and methods

2.1. Microfabrication process of micro-objects

2.1.1. Dip-coating and film transfer

Fig. 2 shows a simplified representation of the required steps in the microfabrication process. Polymer films from PDLLA (Purasorb 7028; Corbion) were obtained by vertically dipping a 4-inch stainless-steel wafer into a solution of (9% w v⁻¹) dichloromethane (DCM; Sigma-Aldrich) and withdrawing it from the solution with a speed of 40 mm/min (Fig. 2A). After 30 min of solvent evaporation, the dip-coated PDLLA film was transferred from the stainless-steel wafer and laminated onto a precoated 4-inch silicon wafer (Fig. 2B). The coating consisted of a layer of Omnicoat™ (MicroChem), which was spun onto the silicon wafer with a speed of 3000 rpm for 30 s and baked at 180 °C for 1 min to let residual solvent evaporate. This Omnicoat™ layer was applied to improve the adhesion of the second layer consisting of poly(vinyl alcohol) (PVA; Mw: 13,000–23,000; Sigma-Aldrich). The PVA was dissolved in dH₂O (5% w v⁻¹) and filtered (mesh size: 0.2 μm) before application. The PVA layer was spun on top of the Omnicoat™ layer with a speed of 2000 rpm for 30 s and baked for 15 min at 120 °C. The PVA coating acts both as a promoter for the adhesion of the PDLLA film to the wafer and as a water-soluble sacrificial layer in the object release process after patterning the film.

2.1.2. Mold fabrication and imprinting

Micromolds based on 4-inch silicon wafers and containing arrays of cavities for replicating three different shapes of micro-objects (cubes, donut-, and LEGO™-like objects) were made by standard UV lithography followed by deep RIE (AMS 100SE; Adixen). To facilitate demolding, an antisticking coating of perfluorooctyltrichlorosilane (FOTS; Sigma-Aldrich) was applied to the molds. The laminated PDLLA film was imprinted between the mold and the PVA layer with a pressure of 30 bar (E) after etching, free-floating micro-objects were obtained by dissolving the sacrificial PVA layer underneath the objects when immersing the wafer in demi water. PDLLA, poly(D,L-lactic acid); DCM, dichloromethane; PVA, poly(vinyl alcohol); FOTS, perfluorooctyltrichlorosilane. While

2.2. Formation of cell and micro-object assemblies

2.2.1. Cell culture and media

Human bone marrow stromal cells (hMSCs) were isolated and proliferated as described previously [29]. Bone marrow aspirates were obtained from patients who had given written informed consent. Briefly, aspirates were cultured in minimal essential medium (α-mem; Gibco, Life Technologies) complemented with 10% v/v heat-inactivated fetal bovine serum (FBS; SA origin; Sigma-Aldrich), 100 U mL⁻¹ of penicillin and 100 μg mL⁻¹ of streptomycin (pen/strep; Gibco), 0.2 mM l-aspartic acid 2-phosphate magnesium salt (ASAP; Sigma-Aldrich), and 1 ng mL⁻¹ of basic fibroblast growth factor (bFGF; Isokine).

Human umbilical vein endothelial cells (HUVECs, ATCC) were cultured in endothelial cell growth medium (EGM-2; Gibco). While

![Fig. 2](image-url) A schematic representation of the five sequential processing steps to fabricate free-floating PDLLA-based micro-objects. (A) PDLLA was dissolved in DCM (9% w v⁻¹). PDLLA films with a thickness of approximately 5 μm were created from this solution on a stainless-steel wafer by dip-coating. (B) Before imprinting, the dip-coated PDLLA film was peeled from the stainless-steel wafer and transferred onto a silicon wafer, which was then spin-coated with a layer of PVA. (C) As a mold, a silicon wafer containing the inverse structures of the micro-objects to be molded and coated with an FOTS antisticking layer was applied. Micro-objects were thermally imprinted under vacuum at a temperature of 130 °C. (D) The imprinted micro-objects were then still connected through a residual layer of PDLLA, which was removed by directional oxygen plasma etching. (E) After etching, free-floating micro-objects were obtained by dissolving the sacrificial PVA layer underneath the objects when immersing the wafer in demi water. PDLLA, poly(D,L-lactic acid); DCM, dichloromethane; PVA, poly(vinyl alcohol); FOTS, perfluorooctyltrichlorosilane.
cocluturing cells, a mixture of α-mem and EGM-2 medium (1:1) was used. In cases in which solely hMSCs or HUVECs were cultured, α-mem and EGM-2 medium were used, respectively.

2.2.2. Agarose-coated well plates and agarose microwell arrays
To obtain hemispherically shaped cell-repellent layers, a solution of agarose (2% w/v; UltraPore; Thermo Scientific) in PBS was added to conventional flat-bottom 96-well plates to form a gel. A similar solution was applied to obtain microwell arrays by means of casting the agarose solution on a patterned polydimethylsiloxane (PDMS) disk. This PDMS disk was prepared by curing PDMS solution in a Plexiglas™ mold, which in turn was patterned by means of an in-house-built microdrilling device.

2.2.3. Live-dead staining
Cell viability was assessed by staining the aggregates with 5 μg mL⁻¹ of calcein AM (Thermo Fisher Scientific) and 40 μg mL⁻¹ of propidium iodide (PI; Sigma-Aldrich) stains, where calcein AM stains viable cells green and PI stains nuclei of necrotic and apoptotic cells red. The stained aggregates were imaged by fluorescence microscopy (EVOS FL; Life Technology).

2.2.4. Cell tracker
To distinguish the two cell types (hMSCs and HUVECs) in coculture, they were labeled with Vybrant Di-I (red) and Di-O (green) tracker dyes (Molecular Probes, Invitrogen). To label the cells, 1 x 10⁶ cells mL⁻¹ were resuspended in serum-free medium developed at Roswell Park Memorial Institute (RPMI) and 5 μL of dye solution was added per milliliter of cell suspension. While labeling, the cell suspension was incubated for 20 min at 37 °C. The cell suspension was centrifuged at 390 x g for 5 min, and the supernatant was aspirated. Afterward, the cells were resuspended in RPMI and the centrifugation steps were repeated for another 2 times. After seeding the cells, a recovery time of at least 10 min was taken into account before imaging the cells.

2.2.5. Scanning electron microscopy (SEM)
Cellular attachment and distribution within the hMSC-micro-object aggregates were characterized by scanning electron microscopy (SEM) analysis (XL 30 ESEM-FEG; Philips). After 7 days of culture, the aggregates were fixed for 30 min in 10% formalin. The fixed aggregates were dehydrated in sequential ethanol series and critical point-dried from liquid carbon dioxide (CPD 030 Critical Point Dryer; Balzers). The dried aggregates were gold sputter-coated (108auto; Cressington) before SEM analysis. SEM images were obtained with an acceleration voltage of 30 kV and a working distance of 10 mm.

2.2.6. Histology
After 1 week of culture, unless stated otherwise, aggregates were fixed for 30 min in 10% formalin and stored in PBS at 5 °C until further processing. Samples were dehydrated using a sequential ethanol series (60, 70, 80, 90, 96, and 100% ethanol; 30 min for each step) and subsequently embedded in glycol methacrylate (GMA). The obtained blocks were sectioned at 5-μm intervals and stained with hematoxylin and eosin (Sigma-Aldrich) for visualization of the nuclei and cytoplasm, respectively. The stained sections were imaged using bright-field microscopy (Nanozoomer; Hamamatsu).

2.3. Image analysis
Circularity was quantified using ImageJ [30]. The projection area A and the corresponding perimeter p of the aggregates were analyzed and measured. Circularity was then calculated as a function of A and p, following Eq. (1).

\[ \text{Circularity} \% = \left( \frac{4A}{p^2} \right) \times 100 \]  
(Equation 1)

2.4. Statistical analysis
Results are presented as mean ± standard deviation and compared using one-way analysis of variance (multiple conditions) with a Bonferroni post-test. Statistical significance was set to p-value < 0.05 (*).

3. Results and discussion

3.1. Microfabrication process
A large amount of approximately 2.5 x 10⁶ objects is obtained within one imprinting cycle after which they are still connected by a thin residual layer (Fig. 2B). Along with the design of the mold, the optimal thickness of the dip-coated polymer film was calculated to minimize the thickness of the residual interconnecting layer without compromising the mold filling and consequently the replication fidelity. In this conjunction, the aforementioned uniformity of the polymer film thickness is crucial. In literature, several approaches to obtain free-standing or residual layer-free micro-objects in connection with polymer micromolding or replication are proposed [31]. These are, among others, based on selective mold functionalization and filling followed by microtransfer molding or related printing processes [32-34], through-thickness embossing with a rubber-assisted etching mechanism [35], or (micro)punching [36].

In the process presented here, similar to original NIL, an optimized oxygen plasma etch protocol removes the residual layer between the micro-objects after demolding the imprint (Fig. 2C), while simultaneously etching the top part of the objects. This renders the objects slightly more hydrophilic on all accessible sides because of the introduction of oxygen-containing groups and roughness on the surface. A surface profile plot of the mold and of the imprinted polymer film before oxygen plasma etching was obtained by white light interferometry (Fig. S1). The imprinted film showed the inverted surface profile of the mold, indicating a successful filling of the mold and a successful demolding of the imprinted film. By removal of the residual layer by plasma etching, free-standing PDLLA micro-objects were obtained. After etching, a surface scan was performed by AFM (Fig. S2). The height profile on the top of the object, which appears as a sort of rim, is a result of the mold fabrication process. The fact that this irregularity is conserved after oxygen plasma etching shows the high degree of directionality of the oxygen plasma etching procedure. The underlying sacrificial layer from PVA was dissolved in demic water, releasing the micro-objects from the wafer (Fig. 2D).

As indicated, PDLLA was chosen as a model thermoplastic biopolymer and can be substituted by every other biocompatible thermoplastic. PDLLA is known to form acidic products during its usually biodegradation. When about 1 x 10⁷ cube-shaped objects, corresponding to approximately 0.32 cm³ total volume, were stored immersed in 5 mL of deionized (DI) water at room temperature, after 6 months, no change in the pH of the DI water could be observed. Although the degradation behavior of the material has to be assumed to be different under physiological conditions in vivo, it is expected that waste products can be cleared at a sufficient rate to prevent tissue damage and that cells will have ample time to express their own mechanically supporting extracellular matrix scaffold. Regarding the latter, we anticipate that material degradation and taking over of the material’s mechanical stabilization role by formed tissue will be more uniform for many microscaffolds than for a single macroscopic scaffold. This should be particularly the case for polymers rather undergoing bulk than surface degradation, such as the chosen PDLLA, where the sudden structural collapse of a bigger volumetric construct might be critical in comparison with the gradual resorption of many well-distributed small centers of degradation.

In case other materials are preferred for a specific biomedical application, in principle, as indicated, every other amorphous thermoplastic biopolymer can be used and processed, both biodegradable and bioinert ones. For example, the bioresorbable polymers poly-(lactic-co-glycolic
acid) (PLGA) and poly-(1,3-trimethylene carbonate) (PTMC) have already shown a good compatibility with hot embossing and NIL-based procedures [37]. For the suggested applications of the micro-objects in the fields of in vitro models or cell expansion, biostable materials could be preferable. The probably most trusted material in this concern, (tissue culture) polystyrene, was shown by us to be also processable by micro-imprinting [38], the process also used for PDLLA in this study (even the same equipment was used). For each polymer, the method has of course to be optimized with respect to dip-coating, film transfer, spin-coating (of a sacrificial layer material and the affinity to it to ensure proper demolding after the imprint), imprinting, and etching parameters. The decoupling of making the film to be imprinted on a first substrate and the sacrificial layer on a second one, in each case from dissolved polymers, circumvents problems, such as local thickness variations of the two materials, related to a (partial) dissolution of the sacrificial material during coating of the material to be imprinted. This opens up many possibilities concerning material choice. Crystallinity of the polymer should be avoided because it can lead to an inhomogeneous progress of the plasma etching for residual layer removal, also affecting the surface roughness of the micro-objects.

3.2. Shapes and sizes of the micro-objects

Previously, we could show that by varying the shape and size of objects and the ratio of cells to objects, the compaction of the aggregates could be altered [14]. Consequential changes of the aggregates' permeability may in turn change the cells' survivability within the aggregates. Many other studies have revealed the influence of substrate geometry, or shape, and topography, both at a micro- and nanoscale, on the morphology of cells and thus their function [24,39,40]. To demonstrate the degree of freedom in terms of object shapes that can be realized using the fabrication method described here, besides cubes, also donut- and LEGO™-shaped micro-objects were fabricated (Figs. 3A–C). In case of the latter two shapes, open free volume is introduced in the objects that cannot be occupied by other objects upon assembly and therefore has an impact on aggregate compaction. Instead, the volume(s) can be entered by cells, or (temporarily) perfused by culture medium or interstitial fluid. The many microlumens do not only introduce a large surface-to-volume ratio that can be beneficial for cell adherence but could also aid (nutrient) diffusion. The corresponding mold designs can be found in Fig. S3. Many more shapes and sizes of micro-objects can be realized by this method with freedom of design in so-called ‘2.5D’, namely, in 2D with the third dimension only being variable over an entire object but not within the same. The dimensionality of the 2.5D micro-objects is increased by them being dynamically (re-)arranged by the cells and thereby also being rotated (Figs. 3F–J). Micro-objects that are more three-dimensional, with surfaces stepped, sloped, or curved out of the mold(ing) plane, can be replicated in mold cavities being created, for example, by various 3D silicon micromachining technologies [41], or two-photon polymerization laser lithography [42] on a suitable substrate for the creation of a master mold in combination with copying the master into a working mold.

![Fig. 3. Micro-objects of different defined shapes such as (A) cubes, (B) donut-, and (C) LEGO™-shaped (porous) objects can be obtained and are shown here in suspension in culture medium to incubate them before cell seeding. A live-dead assay on the full aggregate of 5 × 10⁴ hMSCs with (D) approximately 2000 cubes and (E) approximately 700 donuts shows minimal cell death after 14 days of culture. SEM images show a compact assembly of (F, G) cubes or (H) donuts with hMSCs after 1 week of culture. (F) One aggregate was disrupted on purpose to be able to study the core of the aggregate. It can be seen that the core of the aggregate shows a diameter of approximately 600 μm. The scale bars represent (A–C, F, J) 100 μm, (G) 20 μm, (H) 50 μm, and (I) 500 μm. hMSCs, human bone marrow stromal cells; SEM, scanning electron microscopy; ECM, extracellular matrix.](image-url)
3.3. Cell viability during culture with micro-objects

The absence of photochemistry-based toxic traces increases the possibilities for clinical translation of the objects in multiple applications in the field of TE and regenerative medicine. To investigate cytotoxicity, approximately 2000 cube- and 700 donut-shaped objects were combined with 5 × 10^4 hMSCs on a cell-repellent agarose layer, which was added to conventional 96-well plates. The number of micro-objects was corrected for the volume of each type of micro-object to make sure that the total volume of objects per well is kept constant in each well. Figs. 3D (cubes) and E (donuts) show fluorescent microscopy images with the focal plane at the widest part of the aggregate, which should correspond to approximately the half height of the aggregate. Because it is not a confocal image, the signal observed is retrieved from the aggregate as a whole. The aggregates with a diameter of around 600 μm after 14 days of culture show viable cells in green and dead cells in red. Although the donuts theoretically introduce more pore volume in a tissue construct than the cubes, which may optionally lead to a higher cell viability, no differences in viability as a result of this pore volume were observed. It can be observed that the majority of cells combined with micro-objects are viable. Higher magnification images and split channels (green = live, red = dead) confirm this finding for 3, 7, and 14 days of culture (Figs. S4, S5, and S6, respectively). Furthermore, the results of controls in the form of aggregates from only 5 × 10^4 cells and from 5 × 10^6 cells combined with approximately 2800 PDLLA microspheres, fabricated by an emulsion-solvent-based extraction method (supplementary information) without oxygen plasma treatment of the spheres, show similar degrees of viability (Figs. S4–6). Yet, the aggregates formed from cells only without the presence of micro-objects or microspheres show smaller diameters after 7 and 14 days of culture, indicating that the addition of objects and spheres indeed allows for larger-sized tissue constructs without the requirement of adding more cells.

Other studies on cells-only aggregates show aggregate diameters up to approximately 600 μm while maintaining viability [43,44]. Rivron et al. [27] introduced an approach to prevascularize microtissues through cell assembly in agarose microwells to potentially overcome cell death related to nutrient deficiency. Yet, their method required multiple sequential assembly steps. Although the resulting tissue constructs showed early indications of vascularization, no evidence for an interconnected vascular network was found. Recently, Vrij et al. [26] have shown to challenge the critical size of in vitro cultured tissue constructs via a cell-only bottom-up approach. In both studies by Rivron et al. [27] and Vrij et al. [26], the assemblies were exceeding diffusion limits in the lateral direction only, while the thickness of the constructs remained beyond a critical size where nutrient supply to the inner cells by diffusion was still possible. Furthermore, in the mentioned studies and in other studies on scaffold-free bottom-up TE, high numbers of cells were required to be able to form tissues with clinically relevant volumes [28, 45]. On the contrary, in the method presented here, submillimeter-sized tissue constructs can already be obtained in a one- or two-step procedure within 3–4 days of culture with a relatively low cell number. Supported by the cell-assembled micro-objects scaffold, the resulting aggregates show sufficient mechanical stability to be transferred by means of pipetting or tweezer.

3.4. Cell-guided assembly of the micro-objects

In Figs. 3F–H, scanning electron microscopy (SEM) images show the high degree of integration of the cells and the micro-objects. From the inner part of the aggregate (Fig. 3F and Fig. S7), it can be observed that the cells are homogeneously distributed between the micro-objects throughout the aggregate. Only on the outside of the aggregate, a dense skin layer of (only) cells and ECM is observed, which is confirmed by histological analysis (Fig. 3I). This dense layer corresponds to results found in other studies on the self-assembly of cells in cell-only bottom-up TE approaches and is suspected to be related to geometry-induced processes in the actin-myosin complex [27]. In some aggregates, this layer reached thicknesses of more than 30 μm (Figs. 3I and J). Although this relatively thick and dense layer may reduce the tissue diffusivity, histological analysis shows no evidence for cell death in the form of a necrotic core in aggregates that reach up to millimeter-sized diameters.

As indicated, permeability of the obtained tissue constructs is of paramount importance in sustaining viable and functional cells within the constructs. The circularity of the aggregates can be determined as an indirect measure for tissue compaction and therewith tissue diffusivity assuming a homogeneous tissue density throughout the aggregate. Furthermore, changes in circularity over time and fusion of aggregates after compaction are indications of active tissue remodeling, which is essential, for example, in the musculoskeletal area. Fig. 4A addresses the circularity of aggregates in various conditions with cube-shaped micro-objects after image analysis of a series of 2D phase-contrast images (Figs. S8–11). It can be observed that aggregates’ circularity depends on the cell type used, the ratio of cells to objects, and the culture method – coculture by mixing or by a 2-day later addition of a second cell type (referred to as ‘mixed’ and ‘added’, respectively, see next paragraph). Furthermore, it was observed that the circularity of the formed aggregates increased over time for almost all conditions. These results correspond to previous results on SU-8–based micro-objects in which the circularity of the aggregates showed to be dependent on the ratio of cells to objects, the object size and shape, and the cell type used [14].

3.5. Coculture of hMSCs and endothelial cells on micro-objects

Previously, it has been shown that mechanical forces, among others compressive forces, can result in apoptosis in various cell types [46–48]. Here, cell death induction as a result of such forces during cellular assembly is expected to be limited because of the objects’ geometry-based compaction hindrance. Yet, cell death resulting from diffusion limitations in large-sized tissue constructs when cultured for multiple days may occur. Therefore, vascularization remains a process of vital importance in upscaling to the millimeter or even centimeter range [49]. McGuigan and Sefton [49] have reported the application of an endothelial cell layer around hepatocytes-laden gelatin-based modular tissue constructs. By exposing the endothelial cells to the desired shear stress, they aimed at obtaining a non-thrombogenic phenotype of the cells. Unfortunately, the authors were confronted with low cell viabilities probably because of the cross-linking method introduced to obtain the gelatin gels. Nevertheless, approaches to obtain sufficient blood perfusability in tissue-engineered constructs by introducing endothelial cells have shown promising results [50]. Here, to explore the possibilities of introducing vascular endothelial cells as a first approach toward vascularization, HUVECs were applied in combination with hMSCs and micro-objects.

The order of adding the two cell types to the micro-objects determined the rate and compaction of the cell-guided assembly, the resulting cellular organization, and the cell survival (Fig. 4A). When 2.5 × 10^6 HUVECs were added to 10,000, 5000, or 2500 cube-shaped micro-objects, hardly any cell attachment, assembly, and compaction was observed (Fig. S9). In Figs. S12–14, it can be observed that when HUVECs (stained green) and hMSCs (stained red) were mixed in a 1:1 ratio, within 3 days of culture, the number of HUVECs significantly dropped compared with the number of hMSCs. This effect is independent of the ratio of cells to objects. From these data, it becomes clear that hMSCs are the driving force in tissue compaction and remodeling. When a coculture of hMSCs and HUVECs shall be established, the order of adding cells to the objects and the cell numbers have to be optimized.

When 3 × 10^4 HUVECs were added to an approximately 1-mm-large 1-day-cultured pregelled tissue construct of 3 × 10^5 hMSCS and approximately 3000 cube-shaped micro-objects, they showed to adhere to the construct and form a coating around the aggregate (Fig. 4B). This cell layer of HUVECs disintegrated a few days later, and the detached HUVECs seem to undergo cell death. Yet, when multiple smaller 4-
day-cultured precompacted aggregates from $1 \times 10^4$ hMSCs and approximately 500 donut-shaped objects were combined each with $2.5 \times 10^3$ HUVECs and were allowed to assemble after 1 day, the HUVECs did not disintegrate but stay aligned as HUVEC-rich regions at the aggregates’ interfaces (Figs. 4C and D and Figs. S15 and S16). This 3D network-type spatial cellular organization could foster the establishment of (premature) vascular structures as a vital prerequisite of, for example, bottom-up-engineered bone-like tissue. Our approach is supported by studies on vascularization in conjunction with bottom-up TE by Zhong et al. [9] and McGuigan et al. [50]. Yet, those studies are still based on classical hydrogel building blocks in which the non-endothelial cells are fully embedded in the gel.

To further control the spatial organization of multiple cell types during cell-guided assembly, a cell-repellent microwell array from agarose gel was fabricated to therein combine micro-objects, hMSCs, and HUVECs (Fig. S17). Still based on conventional cells-only aggregation, Tekin et al. [51] showed a partially similar approach, yet their system is more complicated because it requires microwells having temperature-dependent dynamic properties. Here, we show that a static microwell system is a promising tool in obtaining spatial control in cellular assemblies upon sequential seeding of two cell types. It can be observed that HUVECs form a layer around the micro-aggregates of hMSCs and micro-objects. Optimization of cell numbers, culture conditions, and methods of fusing multiple micro-aggregates may aid in the formation of prevascularized complex tissue assemblies.

3.6. Surface functionalization of the micro-objects for bone TE

As a proof of concept for the functionalization of PDLLA micro-objects for bone TE, we coated the objects with octacalcium phosphate (OCP) (Fig. S18). OCP is known as an osteoconductive material allowing for the integration of surrounding bone with the implanted or injected substitute material. Other methods of functionalization of the micro-objects’ surfaces can involve, for example, instructive nanotopographies, chemical modifications by plasma treatments or in the form of peptide-functionalized coatings, or drug delivery coatings. Furthermore, the cell and micro-object assemblies can potentially be complemented with hydrogels as another functional (e.g. binding, viscosity-determining or drug-loaded) phase [52]. In the future, by combining the micro-objects with other instructive or smart materials, novel advanced TE approaches and a broad range of potential applications can be explored.

4. Conclusions

In summary, we present a bottom-up TE approach utilizing engineered micro-objects with tailorable shapes and sizes on which cells can adhere and that can aid in the formation of viable tissue constructs of clinically relevant size. The micro-objects were successfully created by a novel, photoinitiator-free, thermal imprinting-based process conducted on a water-soluble sacrificial layer for the later release of the objects, onto which before the material for the creation of the objects has been
transferred in the form of a film. The process allows the material to be chosen from a wide variety of thermoplastic biomaterials, such as exemplarily PDLLA in case of this study, depending on the demands imposed by the specific research question or clinical application. The latter may range from advanced bone fillers to cell expansion carriers, and beyond. Here, we show the applicability of the objects in *in vitro* culture settings, including hMSCs and HUVECs as representative cell sources in TE, and as spacer material to reduce the required number of patient-derived cells in potential future cell-based TE therapies.

**Conflict of 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**

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