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Published in:
Advanced Materials Technology

Link to article, DOI:
10.1002/admt.201900378

Publication date:
2019

Document Version
Peer reviewed version

Link back to DTU Orbit

Citation (APA):
Article type: Full Paper

Sacrificial Polymer Substrates in Photopolymerization-based Micro 3D Printing for Fabrication and Release of Complex Micro Components

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Keywords: micro additive manufacturing, sacrificial release layers, 3D microfabrication, micro stereolithography, micro medical device manufacturing

Abstract Text
3D printing technology is widely employed in various scientific disciplines as well as industrial applications such as hearing aid manufacturing. While technological advances and increasing resolution are making 3D printing accessible for microfabrication purposes, one question remains: how can small and delicate components like micro gears, lattices or micro medical devices be released from the build surface of the 3D printer without manual intervention? Herein, a method for 3D printing on top of water-soluble sacrificial substrates made from polyvinyl alcohol (PVA) is presented. Pre-fabricated sacrificial PVA substrates can be mounted onto a customized holder and serve as build surface during the 3D printing operation. The substrates do not only facilitate a mild release of 3D printed objects after dissolution of the sacrificial material, they also potentially allow for a convenient manipulation and further array-based processing of pre-determined patterns of printed structures subsequent to the 3D printing procedure. This, in turn, may enable a full integration into automated production lines. The fabrication of PVA substrates is thoroughly
characterized and the 3D printing of various exemplary structures on sacrificial substrates is demonstrated. Finally, the release of 3D printed objects from PVA substrates is shown.

1. Introduction

3D printing has attracted interest since the release of the core inventions of stereolithography (SL) in 1986 and fused deposition modeling (FDM) in 1992, and continues to be a hot topic.\(^{[1,2]}\) The use of 3D printing spans a broad range of applications in different areas such as architecture, automotive industry and medicine. It is used to rapidly produce prototypes as well as functional end-products. Especially in the medical field, 3D printing holds great potential due to the possibility to fabricate customized components with high complexity.

Examples of already successful implementations of 3D printing in industrial fabrication of medical products comprise for example dental appliances and hearing aids.\(^{[3,4]}\) Medical and biomedical applications often require miniaturization of products.\(^{[5]}\) Due to advances in resolution and material availability, 3D printing has become a viable alternative to other microfabrication methods in many areas, including biomedical research.\(^{[6]}\) Research efforts in this area cover a broad variety of 3D printed prototypes, ranging from micro medical components such as bioresorbable vascular stents to microscale 3D scaffolds for tissue engineering, oral modified-release dosage forms as well as to propulsion-capable artificial microfish intended for toxin-neutralization applications.\(^{[7–10]}\)

As pointed out by Quinlan et al., 3D printing, as of yet, has a low overall build rate when compared to other manufacturing processes, e.g. injection molding, and is therefore potentially less attractive for serial production. However, the start-up as well as maintenance costs related to conventional manufacturing processes like injection molding and machining can be very high, especially when the design complexity of the product increases.\(^{[11]}\) The low capital costs of 3D printing and its inherent flexibility thus makes it increasingly attractive for small to medium scale serial production in industry, for serial production of products that require mass customization (e.g. hearing aids), as well as for the rapid prototyping required in...
When compared to other lithography- or micromachining-based fabrication techniques, micro 3D printing is advantageous as the other techniques are limited with respect to three-dimensional complexity, ease of operation and production of assemblies with moving parts. One major drawback of 3D printing, however, is the low resolution compared to e.g. photolithography (minimum feature size of 2-3 µm) and electron beam lithography (down to 5 nm). It must be noted though, that 3D printing resolution is a subject of development and progress is shown on a frequent basis (e.g. custom built 3D printer by Gong et al. with a resolution of 7.6 µm). Using common digital light processing (DLP)-based SL as well as conventional SL 3D printing, voxel sizes down to 30 µm can easily be achieved. Current micro 3D printing requires manual removal of the printed objects from the build surface by human intervention. This presents an obstacle towards automation and serial production as any component pattern enabling further computer numerical control (CNC) or other array-based processing is corrupted. Additionally, small prints are easily damaged during the manual print removal process. The release of single structures from a common substrate, such as a silicon wafer, by means of a sacrificial release layer, e.g. a water-soluble release layer, is a common procedure in micromachining and microfabrication. In FDM 3D printing, Polyvinyl alcohol (PVA) and high impact polystyrene (HIPS) are used as built-in sacrificial support structures. Here, we use a rapidly exchangeable solid sacrificial material substrate as the build surface in vat photopolymerization-based 3D printing. The substrate enables further processing steps in an automated production line, and it allows for a mild release by dissolution of the sacrificial material. The working principle of the presented concept is that the solid sacrificial material substrate can be inserted into a vacuum actuated holder (Figure 1a). This assembly can be inserted into a desktop DLP-SL 3D printer, in which the PVA substrate serves as the build surface (Figure 1b). Later, the substrate can be utilized for easy manipulation of the 3D
printed structures as well as for further processing steps. The PVA substrate can finally be
dissolved in water to release the individual 3D printed structures. Advantages of PVA, in this
case, include water solubility, chemical resistance against many solvents and
biodegradability.[17]

2. Results and Discussion

2.1. Compatibility Study using Raman Spectroscopy

As a prerequisite for the presented concept to work, it is of importance to ensure that the
contact between the substrate material and the liquid uncured 3D printing photopolymer does
not alter the chemical status of the photopolymer and therefore not interfere with the
photocuring reaction during the 3D printing procedure. Raman spectroscopy was performed
and the spectra (molecular fingerprints) of untreated photopolymer as control and
photopolymer after different durations of exposure to potential PVA contamination (1 h, 3 h,
1 d, 5 d) compared (Figure 2). No changes could be observed and the PVA did not dissolve in
the photopolymer or affect it otherwise. Consequently, we conclude that PVA does not alter
the chemical status of the photopolymer, does not interfere with the 3D printing process and
thus is a suitable substrate material. Supplementary experiments with further 3D printing
photopolymers and PVAs support these findings and their results can be found in the
supporting information (Figure S 1, Supporting Information).

2.2. PVA Substrate Fabrication

The substrates used in this work for vat photopolymerization 3D printing were fabricated by
FDM 3D printing of a precursor substrate and subsequent compression molding (Figure 3a,
b). This fabrication route has been chosen due to accessibility and increased flexibility with
respect to the iterative nature of substrate fabrication optimization. Moreover, it demonstrates
the successful fabrication at lab scale with most simple and affordable equipment.
To demonstrate the possibility for cheap and scalable substrate production we also performed laser cutting of a sheet of PVA (Figure 3c) as well as direct injection molding of PVA-polymer substrates (Figure 3d).

2.3. Geometrical Characterization of PVA Substrates

For 3D printing, the fabricated PVA substrates must be produced with suitable surface characteristics as well as uniform thickness. It is required that the substrates have sufficient surface flatness to ensure good contact between the substrate and the polymerization interface during the printing procedure, especially when the first layers of the objects are created. The peak-to-valley flatness parameter (FLTt; ISO 12781) was locally probed in areas of 1.27 x 0.96 mm using optical profilometry with digital interferometry (DI) and confocal (CF) observation conditions and analyzed after applying a robust gaussian filter (cut-off: 25 µm; ISO 16610) to eliminate noise, outliers and short-wave details (Figure 4a). Different substrates were analyzed: compression molded (CM), hand-roughened (CM-S) and injection molded (IM) PVA substrates. A commercial anodized aluminum 3D printer build platform (BP), plain aluminum substrates (Al) and a silicon wafer (Si) were included as reference substrates. BP and Al substrates served to compare the fabricated PVA substrates to frequently used 3D printing surfaces, Si exclusively served as a quality reference. The analysis of the flatness measurements (Figure 5a) shows that, except for CM and Al samples, which exhibit similar flatness (FLTt ≈ 2.4 µm; DI), samples have significantly different FLTt values with large effect sizes (Table S 1, Supporting Information). During the compression molding, the polymer surface adapts the negative of the molds’ surface texture. Thus, CM and Al samples have similar flatness as CM substrates were molded with the use of flat aluminum sheets. While BP has the lowest flatness (FLTt ≈ 12.11 (DI) and 15.55 µm (CF)), Si has the highest flatness with an FLTt value (0.18 µm; DI) up to two orders of magnitude lower than the ones of the other samples. In comparison with CM samples (FLTt ≈ 3.37 µm; CF), CM-S samples show a reduced flatness (FLTt ≈ 5.46 µm; CF), which can be explained
by the hand-roughening treatment as sanding marks can be observed (Figure 4a). IM samples also show lower flatness (5.78 µm; DI) when compared to CM. In the case of IM samples, the surface texture is determined by the manufacturing of the molding tool.

The roughness of a surface can affect adhesion and has been found to be associated with the bond strength of adhesives. In this regard, Shahid et al. found that the average roughness (Ra) is linearly correlated with average cleavage strength of steel/adhesive/steel cleavage specimens, probably due to an increase in effective surface area and through the formation of “mini scarf joints on adherend surfaces at micro level”.[20] Likewise, the successful use of surface roughening treatments for increased adhesive bonding of titanium to polymer composites has been thoroughly discussed.[21] During the 3D printing procedure, it is fundamental that the first layer of cured photopolymer adheres well to the build surface since the 3D printed objects are subject to tensile stress due to continuous movement of the Z-axis and subsequent separation from the polymerization interface. The local surface roughness, more specifically the arithmetical mean height (Sa), was determined using digital interferometry-based optical profilometry (Figure 4b).[22] The evaluation of conducted Sa measurements (Figure 5b) show significant differences with large effect sizes between the different samples, except for CM-S (Sa ≈ 573 nm) and IM samples (Sa ≈ 623 nm) (Table S 2, Supporting Information). Si has the lowest roughness (Sa ≈ 2nm), which matches the specifications of the manufacturer, while BP appears to have the roughest surface (Sa ≈ 1.79 µm). When compared to BP, CM has a significantly lower roughness (Sa ≈ 134 nm). The hand-roughening treatment is seen to greatly increase the roughness of CM-S substrates when compared with CM substrates, which can also be seen in the example of the very complex surface in Figure 4b. Even though CM-S and IM have similar Sa values, the surfaces exhibit very different surface morphologies. The Sa value does not give any indications of the surface morphology and therefore we calculated the developed interfacial area ratio (Sdr), which is a measure for surface complexity and also a better indication of adhesive properties (Figure
The analysis of Sdr values shows significant differences between all samples (Table S 3, Supporting Information). The results mainly follow the trend that could be observed in Sa measurements, with better differentiated values for CM-S and IM. The Sdr-value for CM-S was two orders of magnitude higher than for CM.

Thickness measurements of different PVA substrates were conducted (Figure 5d), and the measurements on deviation from target thickness show that values obtained for the compression molded PVA substrates (CM) lie in a range of \( \approx 26 \, \mu m \). In the case of hand-roughened compression molded samples (CM-S) and injection molded samples (IM), the measurements lie in a range of \( \approx 43 \, \mu m \) and \( \approx 23 \, \mu m \), respectively. To ensure a successful printing without the need for recurring calibrations, the thickness deviation of the substrates should be smaller than the layer thickness of the individually exposed layers during the 3D printing procedure. As the layer height of the 3D printer in this case was 25 \( \mu m \), a thickness deviation above 25 \( \mu m \) could call for recurring homing calibrations. The lack of precision in thickness repeatability for CM substrates can partially be explained by the deviation in material dispensing during FDM 3D printing of the precursor substrate. Here, an observed weight deviation with a range of 8.66 mg (\( N = 10 \)) can translate into a 16-17 \( \mu m \) thickness deviation when taking the final substrate dimensions into account and assuming a PVA density of 1.19-1.31 g cm\(^{-3}\).[24] Furthermore, the manual handling during the molding procedure leaves room for error. It is to be expected that the thickness deviation is higher for CM-S substrates than for CM substrates, since it is likely that the hand-roughening treatment unevenly affected the final thickness of the substrates. We note that the deviation for CM substrates is not much higher than for injection molded substrates. A further optimization of the CM fabrication processes can lead to a much higher precision in thickness repeatability, allowing for users without access to injection molding to fabricate their own high-quality substrates.
Since standard deviations were smaller than ± 25 µm in all cases, the study was continued based on the same CM fabrication process and without recurring homing calibrations.

2.4. 3D Printing on PVA Substrates

Using a commercial DLP-SL 3D printer and a custom vacuum-actuated holder (see Figure 1a), we were able to 3D print various exemplary structures on CM PVA substrates (Figure 6). The workflow allowed us to 3D print arrays of defined geometrical objects on top of PVA substrates and to remove the entire substrate from the holder after the finished 3D printing operation. 3D printed example structures include those, e.g. helical micro-gear and micro-truss lattice, which are nearly impossible to fabricate by other conventional manufacturing techniques, such as injection molding or micromachining.

Employing a specifically for this purpose designed and 3D printed test object (Figure 6h) and a texture analyzer, we determined the detachment force to study the relationship between surface characteristics of the build surface and bond strength of the 3D print. In this regard, we interpret the detachment force to be proportional to the bond strength as a higher detachment force is caused by a higher bond strength. The footprint of the test object matches the dimensions of areas probed for the flatness characterization. Arrays of the test object were 3D printed on BP, Al, CM and CM-S substrates (Figure 7a, b, c) and using a customized texture analyzer setup (Figure 7d), detachment force as well as work of adhesion (area under curve of detachment graph) were determined (Figure 7e). The evaluation of the detachment force (Figure 7f) shows statistically significant differences with large effect sizes between all samples (Table S 4, Supporting Information). Hand-roughening of PVA substrates significantly affected the bond strength between the test objects and CM-S substrates, thus revealing a much higher detachment force when compared to CM substrates. Despite having a rougher surface, Al and BP have lower detachment forces while Al has the lowest. An explanation for this might be the influence of other adhesion promoting factors, which could lead to the occurrence of increased polymer-polymer (cured photopolymer-PVA) interactions.
An example for those could be electrostatic adhesion between the PVA substrates and the 3D printed objects according to the electrostatic theory of adhesion, hence leading to a higher bond strength.\textsuperscript{[25]} This possible explanation is consistent with the observation that the detached 3D printed objects kept sticking to other objects, e.g. the custom microgripper after they have been detached from the PVA substrates. The evaluation of the work of adhesion (Figure 7g) follows a similar trend, except for the fact that no significant difference between BP and Al can be found (Table S 5, Supporting Information).

The results indicate that the surface of the proposed PVA substrates may be tailored for optimized performance, e.g. through a surface roughening treatment for increased adhesion of the 3D printed structures to the substrate. However, as already noted in the case of potential influence of electrostatic adhesion, there are most probably further parameters, which can positively or negatively influence the adhesive bond (e.g. surface chemistry and interfacial failure). Moreover, the formation of the adhesive bond itself does not represent the only potential influence that is relevant for a successful 3D printing outcome. As mentioned earlier, the adhesion of 3D printed objects to the substrate was considered to be crucial as the freshly printed layers are subject to tensile stresses caused by the vertical motion of the build platform and the repetitive contact to and separation from the polymerization interface. It is to be expected that the same tensile stresses act on the PVA substrate as well and they could potentially lead to deformation or detachment of the entire PVA substrate. Although these issues could not be detected during the experiments, the different forms of mechanical stress and the consequences they might have should be considered with regards to the potential setting and application of the proposed method.

2.5. Release of 3D Printed Objects from PVA Substrates

An array of helical micro-gears (Figure 6e) 3D printed on CM PVA substrates was released from the substrate within 150 min. (Figure 8a; Movie S 1, Supporting Information). Scanning electron microscopy of the harvested individual micro-gears shows that the gears are intact.
and free of substrate material (Figure 8b, c). The dissolution rate of PVA is highly dependent
on the type of PVA (degree of polymerization, degree of hydrolysis) and also on the
temperature.\textsuperscript{[16]} Furthermore, the time needed for the dissolution depends on the amount of
material to be dissolved. In Figure 8a it is visible that the PVA substrate was inserted into the
dissolution medium in a 90-degree orientation. Since, upon contact with water, the PVA
started to dissolve and became rather jelly-like, the substrate lost its mechanical rigidity and
deformed. The latter can be observed as a progressing trend within the first 90 minutes. At \( t = \)
90 min some individual released micro-gears were visible, whereas the remnants of the
substrate transformed into a clot and engulfed all other micro-gears that were still in contact
with the PVA. As a consequence, the PVA needed to be fully dissolved in order to release all
3D printed micro-gears, which accounts for the 150 min release time.

To illustrate that the release time can be reduced, composite CM PVA substrates with a non-
dissolving polylactic acid (PLA) core were fabricated. The PLA core was fully encapsulated
by the surrounding PVA and reduced the total amount of PVA by 50%. Using this substrate,
the same array of micro-gears could be released within 90 min. (Figure S 2, Movie S 2,
Supporting Information). In this case, the non-dissolving PLA core helped to largely maintain
the mechanical rigidity of the substrate, so that a clot formation was impossible and only the
PVA interfacing the 3D printed micro-gears and the PLA core had to dissolve in order to
release all micro-gears. It has to be emphasized that the two experiments are only examples of
how the release process could look like, as many influential parameters, such as type and
properties of the utilized PVA, have not been investigated. Beyond the dissolution properties
of the PVA, factors like excitation type (ultrasound, stirring, flow etc.) and substrate porosity
might have a positive influence on diffusion and convection of the dissolving PVA and
therefore on the dissolution time. A further optimization of the release procedure can be
expected to drastically reduce the required dissolution time.
3. Conclusion

In summary, we have demonstrated the use of water-soluble PVA sacrificial substrates in vat photopolymerization-based 3D printing. The fabrication of substrates with suitable flatness, roughness and thickness characteristics was accomplished at lab scale, and their specifications are compatible with industrial fabrication. The substrates were chemically compatible with different 3D printing photopolymers and exhibited good bond strengths to the 3D printed objects. Using a custom-made vacuum-actuated holder, PVA substrates could be rapidly exchanged and taken from the 3D printer, thereby enabling further array-based processing and potential integration into automated production lines. We showed that advanced 3D printed objects can be released through dissolution of the substrate, thereby eliminating the need for manual intervention. Consequently, the proposed method might be potent of promoting the application of 3D printing for the serial production of complex micro components.

4. Experimental Section

Materials: All chemicals and reagents were used as received. For the fabrication of PVA substrates different kinds of PVA material were used: RS Pro PVA 3D printing filament (RS Components A/S, Denmark), MOWIFLEX™ C17 and MOWIFLEX™ C600 (Kuraray Nordic Ab Oy, Finland). HTM 140M V2 3D printing photopolymer (EnvisionTEC GmbH, Germany) was used to 3D print onto PVA substrates. Further photopolymers were used for a compatibility study: PIC100 (EnvisionTEC GmbH, Germany) and Form Clear resin (Formlabs GmbH, Germany). 2-propanol (Sigma-Aldrich Denmark A/S, Denmark) was used for the post-treatment of 3D printed structures.

Compatibility study using Raman spectroscopy: A compatibility assay was performed by incubating 200 mg of solid polyvinyl alcohol (PVA) material in 1 ml of liquid 3D printing photopolymer and analyzing a sample of the liquid after successive timepoints (1 h, 3 h, 1 d, 5 d) by Raman spectroscopy. When considering the use of one PVA substrate with a weight of 805 mg in the supplied vat of the 3D printer, which is filled with 150 ml of 3D printing
polymer, the concentration amounts to 5.37 mg ml\(^{-1}\). The ratio of PVA to 3D printing polymer in the compatibility study was chosen to be multiple times higher. 3D printing polymer which was not in contact with PVA served as control. Raman spectroscopy was employed to determine molecular fingerprints of the samples.

Raman spectra were acquired with an in-house-built Raman spectroscopy system with improved sensitivity for Raman scattering registration in case of liquid samples. The system is based on a high power (500 mW) multimode laser with a wavelength of 785 nm. The laser had an intensity of 20 mW µm\(^{-2}\) and was focused on the sample through a liquid container with a CaF\(_2\) bottom plate. Measurements were carried out with a spectral resolution of 1.8 cm\(^{-1}\) in the range from 350 to 2100 cm\(^{-1}\) and collected using a CCD sensor. Wavelength and spectral sensitivity calibration of the instrument was performed according to ASTM 1840 and ASTM E2911 international guidelines.

Fabrication of PVA substrates: Whereas the FDM-3D printing step did not serve to produce the final substrate, but rather as a material dispensing step to fabricate a precursor substrate of a certain size, the compression molding process acted to transform the precursor into a flat substrate of desired shape by using a mold assembly. For the fabrication of the substrate precursor, a commercially available Original Prusa i3 MK2S desktop 3D printer was used (Prusa Research, Czech Republic) to print with likewise commercially acquired RS Pro PVA filament with a 100% infill, a hotend temperature of 210 °C and a print bed temperature of 85 °C (first layer) and 60 °C (following layers). The volume of the substrate precursor was calculated to equal the volume of the mold cavity which is used in the compression molding step. While the FDM 3D printing method can be quite accurate, it is – due to the nature of this technology – not precise enough to exactly dispense the correct volume of material as the layer-by-layer and line-by-line fabrication leads to the creation of small gaps within the print even though the infill ratio is set to 100%. In order to compensate for this phenomenon, the volume of the substrate precursor was increased by 3%, which was found to be an acceptable
value to obtain a good substrate after compression molding. After FDM 3D printing of the precursor substrates, the substrates had an average weight of 804.96 mg (N = 10), ranging from 800.09 to 808.75 mg with a standard deviation of 2.93 mg.

The mold assembly for the compression molding consisted of a 1 mm thick aluminum mold, 90 µm aluminum foil and 1 mm stainless steel sheets. The compression molding procedure was carried out with a pressure of 55 kN and a temperature of 160 °C using a PW-H HKP300 laboratory press (Paul-Otto Weber GmbH, Germany).

For some of the resulting substrates, the surface was modified by sanding one side with 600 grit sanding paper.

Composite compression molded substrates consisting of PVA and polylactic acid (PLA) were as well fabricated using an FDM 3D printing and a compression molding step. PLA inserts were 3D printed with smaller dimensions, constituting 50 % of the final substrate. PVA substrates were designed to have a cavity and the 3D printing procedure was paused as soon as the cavity was completed. Then the PLA insert was inserted into the cavity and the 3D printing procedure was continued. The cavity was closed with the remaining layers of PVA, thereby fully engulfing the PLA in its’ core. The compression molding step transformed the precursor composite substrates into smooth PLA-PVA core-shell substrates of 1 mm thickness using the same conditions as with plain PVA substrates.

In a different approach, a Press 300 SV laboratory platen press (Dr. Collin GmbH, Germany) served to transform 15 g of MOWIFLEX™ C17 PVA polymer pellets into a compressed sheet using a pressure of 50 bar and a temperature of 150 °C for a duration of 1000 s and subsequently cooling it down to 30 °C within 500 s. Substrates of desired shape were cut from the sheet with an Epilog Mini 18 laser cutter (Epilog Laser BV, The Netherlands) which was equipped with a 30 W CO₂ laser. This procedure needed to be performed with the necessary safety precautions as toxic fumes can be release during the procedure.[24]
Injection molding of PVA substrates was performed using an Arburg Allrounder 370A injection molding machine (Arburg GmbH & Co KG, Germany) equipped with an 18mm screw and MOWIFLEX™ C600 PVA polymer. Injection molding parameters were adjusted to 70 bar back pressure, 180 °C melt temperature, 40 °C mold temperature, 50 mm s⁻¹ injection velocity, 500 bar packing pressure, 10 s packing time and 40 s cooling time.

Characterization of PVA and reference substrates: The thickness of the fabricated substrates was measured in the center and in the four corners of each substrate using an RS Pro micrometer screw with an error of 0.001 mm (RS Components, Denmark). A PLu neox optical 3D profiler (Sensofar Metrology, Spain) served to conduct surface topology measurements, using confocal and interferometric microscopy. To analyze the flatness property of the various specimen, 10X interferometry and 20X confocal lenses were used for data acquisition. To compensate the loss in field of view when using the 20X confocal lens, stitching was used to combine four images to one bigger area image. A 50X interferometry lens was used to acquire data for the analysis of the surface roughness. In case of all specimens, a sampling procedure based on a 20 x 20 mm grid was performed to obtain surface measurements in a total of 25 spots in always the same relative positions. A 3” silicon wafer (No. 16013, Ted Pella inc., USA) with a specified roughness and total thickness variation of 2 nm and <20 µm, respectively, as well as the supplied build platform of an EnvisionTec Micro Plus High-Res DLP 3D printer were used as reference surfaces. Treatment and analysis of surface metrology data was done in SPIP 6.7.4 (Image Metrology A/S, Denmark) analytical software.

Computer aided design (CAD): All design tasks were carried out using SolidWorks 2015 (Dassault Systèmes SolidWorks Corporation, USA) and OpenSCAD open source software.

Machining of customized 3D printer build platform: A customized 3D printing build platform featuring a four-point spring leveling mechanism and a vacuum-actuated holding cavity for a print substrate was made to retrofit a Micro Plus High-Res digital light processing (DLP) 3D
printer (EnvisionTec GmbH, Germany). The platform was machined by an external machining shop using a combination of CNC milling and electrical discharge machining.

3D printing on PVA substrates: The 3D printing on PVA substrates was conducted with an EnvisionTec Micro Plus High-Res DLP 3D printer with a XY resolution of 30 µm pixel size and a Z resolution of 25 µm. The 3D printer was retrofitted with a customized build platform to enable a flush leveling of the platform to the polymerization interface of the printer.

Perfactory RP software (EnvisionTEC GmbH, Germany) served to create print files from the prepared CAD models. After the printing procedure, the PVA substrate with printed structures on top was first cleaned from excess printing material in a beaker with 2-propanol placed in an ultrasound bath for 5 min and subsequently post-cured in an UV oven for 10 min (EnvisionTEC GmbH, Germany).

Scanning electron microscopy: All scanning electron microscopy was performed using a TM3030Plus tabletop scanning electron microscope (Hitachi High Technologies Europe GmbH, Germany). A 208HR high resolution sputter coater (Cressington Scientific Instruments, UK) equipped with a gold target was used to coat the specimens with a thin layer of gold (≈ 20 nm) prior to observation.

Determination of detachment force: A TA.XT plus Texture Analyzer (Stable Micro Systems, UK) equipped with a 10 kg load cell and a customized probe was used to measure detachment forces needed to separate a printed sample from different 3D printing substrates. Detachment forces and work of adhesion were computed with a customized python program.

Release of micro 3D prints from PVA substrates: 3D printed structures were released from the PVA substrate by retaining the substrate in a small box with a bottom of fine stainless-steel mesh and placing it in a de-ionized water-filled beaker, which in turn was placed into an ultrasound bath at a temperature of 55 °C. The samples were kept in the ultrasound bath until all PVA was dissolved. A waterproof USB endoscopic camera and Video Velocity Free software (Candy Labs Media, Canada) were used to record time-lapse photos during the
release procedure. The samples were ultimately taken out of the water and left to dry in an oven at 37 °C.

Statistics: All presented statistics were computed using R programming language and RStudio software (RStudio Inc., USA) as well as Microsoft Excel (Microsoft Corporation, USA). As in case of reference samples Si and BP only one specimen was available each, t-test results comparing those with Alu, CM, CM-S and IM samples are based on the assumption that the measured reference samples constitute ideal and representative samples of their kind. The results obtained in these cases can serve as an indication only, because resulting p-values might be distorted. Consequently, the reported effect sizes (Hedges’ g) are more reliable.
Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

Acknowledgements

The authors would like to acknowledge the Center for Intelligent Drug Delivery and Sensing Using Microcontainers and Nanomechanics (IDUN) whose research is funded by the Danish National Research Foundation (DNRF122) and Villum Fonden (Grant No. 9301).

Dr. Guanghong Zeng acknowledges funds from the Danish Agency for Institutions and Educational Grants, metrology for additively manufactured medical implants (short name: MetAMMI, project number: 15HLT09) from the EMPIR program. The EMPIR program is co-financed by the participating states and the European Union’s Horizon 2020 research and innovation program.

The authors would also like to thank Dr. Kristian E. Jensen and Dr. Tommy Sonne Alstrøm for help with the manuscript preparation, Francesco Regi for performing injection molding and Jesper Scheel for photography. Furthermore, the authors would like to thank Ann-Britt Aspholm-van der Brugge and Kuraray Nordic Ab Oy for straightforward communication, great service and provision of free polymer samples.

Conflict of Interest

The authors declare no conflict of interest.
References


Figure 1. a) Design of customized vacuum-actuated substrate holder for the use in a desktop DLP-SL 3D printer. b) Schematic illustration of working principle of using pre-fabricated PVA substrates in a DLP-SL 3D printer. The PVA substrate is used as the build surface and held in place by the vacuum-actuated holder (build platform) which moves in Z direction. In an industrial production line setting, the holder could be operated by a robotic arm which also carries out further processing steps.
Figure 2. Molecular fingerprint of HTM 140 M 3D printing photopolymer after contact with RS Pro PVA filament for different time durations (1 h, 3 h, 1 d and 5 d) and untreated (Control), as determined by Raman spectroscopy.
Figure 3. PVA substrate fabrication. a) Schematic illustration of two-step PVA substrate fabrication sequence using fused deposition modelling (FDM) 3D printing and subsequent compression molding. b-d) Photographs of differently fabricated PVA substrates. b) FDM 3D printed precursor substrates (substrates placed on mold assembly) and compression molded substrates (front). c) Laser-cut substrates from compressed PVA sheet. d) Injection molded PVA substrates in standard object slide format. Scale bars are equal to 25 mm.
Figure 4. Flatness and roughness measurements obtained by optical profilometry. a) Representative surface renderings of substrates used for flatness analysis. Computed from data acquired with a 20X confocal lens in stitching mode (BP, CM and CM-S) and a 10X interferometry lens (Si, Al and IM). b) Representative surface renderings of data used for roughness analysis (Sa and Sdr). Computed from acquisitions with a 50X interferometry lens.
Figure 5. Geometrical characterization of different 3D printing substrates: Plain aluminum (Al), compression molded (CM), hand-roughened CM (CM-S) and injection molded (IM) PVA and reference substrates: Silicon wafer (Si) and commercial anodized 3D printer build platform (BP). Error bars represent 95% confidence interval in a), b) and c) and standard deviation in d). a) Peak-to-valley flatness deviation (FLTt) measurements from optical profilometry surface data obtained with digital interferometry (DI) and confocal (CF) observation conditions. For statistical comparison see Table S 1. b) Arithmetical mean height (Sa) measurements from optical profilometry surface data. For statistical comparison see Table S 2. c) Developed interfacial area ratio (Sdr) computed from optical profilometry surface data. Statistical comparison available in Table S 3. For a), b) and c) counts: N=5 with 5 different samples in case of Al, CM, CM-S and IM and N = 1 with 25 repeated measurements on the same sample in case of Si and BP. d) Micrometer thickness measurements of PVA substrates adjusted to target values with Y = 0 = target thickness value. N = 10.
Figure 6. Photographs and SEM micrographs of 3D printed structures on compression molded PVA substrates (CM). a) Array of printed structures on PVA substrate inserted in vacuum-actuated holder (see schematics in Figure 1a and b). b) 3D printed crosshairs, facilitating evaluation of alignment of PVA substrate and printed structures. c) Circular array of micro-cones. d) DTU logo assembly from separate 3D printed parts. e) Helical micro-gear with a twist of 25°. f) Surgical staple. g) Complex lattice made from micro-sized trusses. h) Small structure used for evaluation of bond strength of 3D print to PVA substrate.
Figure 7 Determination of detachment forces/bond strengths of 3D printed objects on PVA substrates. a-c) Photographs of manufactured samples. Scale bars are equal to 10 mm. a) Test structures 3D printed on plain aluminum substrates (Al). b) Test structures 3D printed on compression molded PVA substrates (CM) and c) hand-roughened CM PVA substrates (CM-S). d) Schematic illustration of texture analyzer test-setup used for the experiments. e) Schematic illustration of obtained displacement curves. f) Evaluation of detachment forces. g) Determined work of adhesion (WOA), which is equal to the area under the curve (AUC) of the displacement graph. Additional to the manufactured samples, a commercial 3D printer build platform (BP) also served as reference substrate. N = 3-6. Error bars represent 95% confidence interval. Statistical evaluation available in Table S 4 and Table S 5.
Figure 8. Release of 3D printed objects from PVA substrates. a) Time-lapse photos taken with a water-resistant endoscopic camera during the release procedure of 3D printed micro-gears from compression molded PVA substrates (CM). Release procedure was carried out in a customized release-chamber/substrate-holder combination at 55 °C in an ultrasound bath. b) and c) SEM micrographs of 3D printed helical micro-gears (see Figure 6e) on stainless steel filtering mesh after dissolution of compression molded PVA substrates (CM) and subsequent release. b) front side. c) backside.
3D printing on top of sacrificial substrates is demonstrated. The used 3D printing workflow enables the 3D printing on quickly exchangeable substrates, further array-based processing of 3D printed products and easy manipulation, as well as integration into industrial production lines. 3D printed products can be mildly released from the substrates upon dissolution of sacrificial material and harvested.

3D printing

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