

MRS 50th Anniversary Prospective



\triangleright review of materials used in tomographic volumetric additive

manufacturing

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Abstract

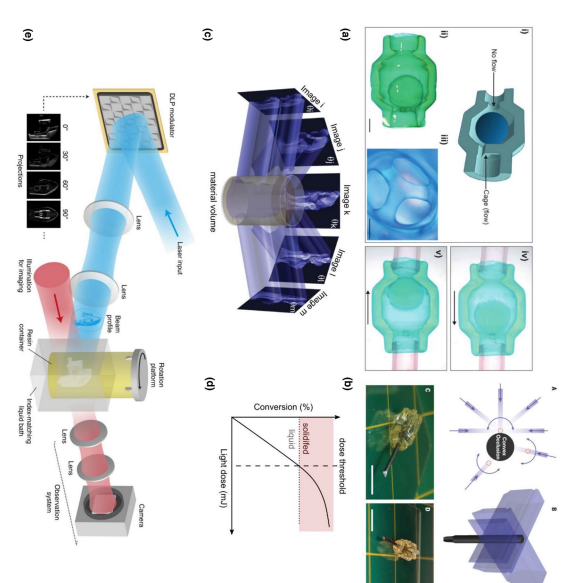
and discuss the perspectives they present. hydrogels for cell-laden biofabrication. Here we review these materials, highlight the challenges to adapt them to volumetric additive manufacturing formulations, tunable stiffness thiol-enes and shape memory foams, polymer derived ceramics, silica-nanocomposite based glass, and gelatin-based for printing. These include viscous acrylates and elastomers, epoxies (and orthogonal epoxy-acrylate formulations with spatially controlled stiffness) methods like stereolithography or digital light printing, volumetric additive manufacturing offers new opportunities for the materials that can be used dose within the photosensitive resin, solidifying the volume of the desired object within seconds. Departing from established sequential fabrication (CT), the method projects dynamic light patterns into a rotating vat of photosensitive resin. These light patterns build up a three-dimensional energy Volumetric additive manufacturing is a novel fabrication method allowing rapid, freeform, layer-less 3D printing. Analogous to computer tomography

Introduction

more recently two photon polymerization (2PP),^[5,6] xologthe object is printed within the resin itself, which supports for complex designs with overhangs or cavities. In others, phy.^[9,10] In many of these methods, support struts are needed metric additive manufacturing or computed axial lithograraphy,^[7] light sheet microprinting,^[8] and tomographic volu-Processing (DLP),^[2,3] selective laser sintering (SLS),^[4] and development of stereolithography (SLA),^[1] Digital Light Different ways of bringing light to the print have led to the crosslinking reactions in liquid resins, thus solidifying them. certain light-sensitive molecules to trigger polymerization or geometries. Light-based 3D printing exploits the ability of try by simplifying the fabrication of designs with complex 3D printing has revolutionized the manufacturing indus-

Instead, light from the subsequent tomographic by solidifying one voxel, one line, or one layer at the time ing is layerless, meaning that it does not fabricate objects methods, tomographic volumetric additive manufactur-[Fig. 1(c)].^[9,10] Unlike most other additive manufacturing able resin from multiple angles with dynamic light patterns ously solidified by irradiating a volume of liquid photocur-In VAM, an entire three-dimensional object is simultanetomographic volumetric additive manufacturing (VAM). overprinted around a metallic shaft, both printed through valve with free-floating elements and a screwdriver handle and (b) illustrates design freedom with a fluidic ball-cage additive manufacturing offer full design freedom. and more recently xolography and tomographic volumetric it. Volumetric methods, such as two photon polymerization, Fig. 1(a) patterns





around its metallic shaft (scale bars = 10 mm, Copyright AAS. (c). In VAM, an entire three-dimensional object is simultaneously solidified by irradiating a liquid photopolymer volume from multiple angles with dynamic light patterns (Copyright AAAS). (d) VAM exploits the thres-holded response of corresponding photosensitive materials to light-induced polymerization. Thanks to this solidification threshold, only VAM (Copyright Springer-Nature). from multiple angles. The liquid unpolymerized resin can be washed away after the print. (Copyright De Gruyter). (e) Exemplary setup for the target object is printed, even if the resin outside the object's target volume inevitably receives some light after having been illuminated Figure 1. (a) Design freedom of tomographic volumetric additive manufacturing (VAM) is demonstrated by the fabrication of a fluidic ball-cage valve with free-floating elements (scale bars=1 mm, Copyright Wiley) and (b) by the overprinting of an acrylic screwdriver handle

builds up an energy dose within the complete volume of the target object. Typical printing times are in the order of tens of seconds^[10] for cm-scale prints with resolutions down to 50 to 80 µm.^[11,12] Additionally, the technique has proven versatile and has been used to fabricate objects in materials such as acrylates, thiol-enes,^[13–15] nanoparticle-loaded composites,^[11] polymer-derived ceramics,^[16] epoxies,^[17] silk bioinks,^[18] and cell-laden hydrogels.^[10,19–23]

In this review, we will cover the underlying functioning mechanism of VAM, briefly discuss the optical and computational strategies developed so far to improve fidelity and

> resolution, cover in detail its applicability to a large variety of materials (Table I), and present our perspective on future applications, challenges and possibilities.

3D printing as tomographic back-projection

A series of 2D light patterns are projected from many different angles onto the vial of photocurable resin to build a cumulative light dose matching the geometry of the printed object [Fig. 1(c)]. Although the entire volume will be irradiated with

Туре	Formulation	Photoinitiator (concentration)	Wavelength	Typical viscosity	Other components	Notes
Acrylate ^[12,34]	Di-pentaerythritol pen- taacrylate	TPO (0.6 mM)	405 nm	>10 Pa·s	TiO2 was added into the resin in one of these works to make it scattering	
Silicone ^[12]	93 wt% vinyl-terminated PDMS+4.7 wt% fumed silica reinforced vinyl-ter- minated PDMS+2.3 wt% (mercaptopropyl) methylsi- loxane-dimethylsiloxane	TPO-L (2.25 mM)	405 nm			Demonstration of 3D printing of full silicones
Acrylate ^[9]	BPAGDA 75 wt%+PEGDA 25 wt%	CQ (5.2 mM)+EDAB 1:1 (weight ratio)		5.2 Pa·s		
Acrylate ^[48]	Urethane Dimethacrylate	2-Benzyl-2-dimethylamino- 1-(4-morpholinophenyl)- butanone-1 (6 mM)	405 nm	11 Pa·s		Printed in microgravity
Acrylate ^[48]	Aliphatic urethane acrylate diluted in isobornyl acrylate (EBECRYL 242 N)	2-Benzyl-2-dimethylamino- 1-(4-morpholinophenyl)- butanone-1 (6 mM)	405	25 Pa·s		Printed in microgravity
Acrylate—epoxy ^[17]	PEGDA / BPAGDA / EEC	CQ (0.02 wt%) EDAB (0.02 wt%) CAT2 (2.9 wt%)	455 / 365 nm			CAT2 first dissolved in propyle ene carbonate
Acrylate ^[17,31]	PEGDA / BPAGDA	CQ (0.1 wt%) EDAB (0.25 wt%)	455 nm	0.32 Pa·s		
Acrylate ^[31]	PEGDA / BPAGDA	CQ (0.1 wt%) EDAB (0.25 wt%)	455 nm	0.32 Pa·s	TEMPO (0.004 or 0.01 wt%)	
Acrylate ^[17,31]	PEGDA / BPAGDA	CQ (0.1 wt%) EDAB (0.25 wt%)	455 nm	1.40 Pa·s		Viscosity adjusted adding BPAGDA
Acrylate ^[30,31]	TEGDMA / BisGMA	CQ (0.2 wt%) EDAB (0.5 wt%)	460/365 nm		o-Cl-HABI (1 or 3 wt%)	HABI pre-dissolved in THF
Acrylate ^[30,31]	TEGDMA / BisGMA	CQ (0.1 wt%) EDAB (0.25 wt%)	460/365 nm		<i>o</i> -Cl-HABI (0.4 wt%)	HABI pre-dissolved in THF
Polymer-derived ceramic ^[16]	Polysiloxane (SPR 684) 85 wt%+1,4-butanediol diacrylate 15 wt%	TPO (2 mM, 0.063 wt %)	405 nm	0.87 Pa·s		
Nanocomposite for glass ^[11]	296 g/mol trimethylol- propane triacrylate (TMPTA)+hydroxyethylm ethacrylate(HEMA)	CQ (0.117 wt%)+EDAB (0.117 wt%)	442 nm	10 Pa·s	TEMPO (0.2 M) in TMPTA (0.5 vol%)	
Shape memory foam ^[14]	TEGDAE: TA-ICN: TME- ICN (0.1: 0.9: 1 molar equivalent of functional groups)	2-Methyl-4'-(methylthio)-2- morpholinopropiophenone (10 mM)	405 nm		23.5 mM ANPHA+0.1 mM TEMPO	

 Table I.
 Formulations used for volumetric additive manufacturing.

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Туре	Formulation	Photoinitiator (concentration)	Wavelength	Typical viscosity	Other components	Notes
Thiol-ene ^[13]	TEGDA+TEGDAE+TAE- ICN+TA-ICN+TME-ICN (in different mixing molar ratios)	2-Methyl-4'-(methylthio)-2- morpholinopropiophenone (10 mM)	405 nm		TEMPO 0.1 mM	
Poly(e-caprolactone) ^[15]	PCL-ene/ PETA-4SH	TPO-L (0.12-0.25% w/v)	442 nm		TEMPO (0.1 mg mL-1)	
Hydrogel (cell-laden) ^[10]	GelMA 10% w/v	LAP (0.037% wt)	405 nm	Thermally gelated		Printing time ≈12.5 s, viabil- ity>85%
Hydrogel (organoid-laden) ^[20]	GelMA 5% w/v	LAP (0.1% w/v)	405 nm	Thermally gelated	Iodixanol	Iodixanol concentration needs to be optimized for each material and cell type. Laden with up to 5 million cells mL^{-1}
Hydrogel (cell-laden) ^[19]	GelNB/PEG4SH 2.5% w/v	LAP (0.05% w/v)	405 nm	Thermally gelated		Laden with murine C2C12 myoblasts at 1 million cells mL^{-1}
Hydrogel (cell-laden) ^[23]	GelMA 5, 8 and 15% w/v	LAP (0.1% w/v)	405 nm	Thermally gelated		Performed in presence of melt electrowritten poly(ε- caprolactone) meshes
Hydrogel (cell-laden) ^[45]	GelMA 10% w/v in PBS	LAP $(0.16 \text{ mg mL}^{-1})$	405 nm	Thermally gelated		Laden with human fibroblasts at 0.5 million cells mL^{-1}
Hydrogel ^[18]	Silk sericin 2.5–5% w/v	Ru/SPS (1/10) (0.25–1 mM Ru)	525 nm	Not measured	Water	Printing time~55–80 s C2C12 myoblasts at 5 million cells mL ⁻¹
Hydrogel ^[18]	Silk fibroin 1.25–15% w/v	Ru/SPS (1/10) at (0.125–1 mM Ru)	525 nm	Not measured	Water	Printing time~30–170 s
Hydrogel ^[22]	GelNB/PEG4SH	LAP (0.05% w/v)	405 nm	Thermally gelated		Printing time ≈10–11 s, viabil- ity>95%
Hydrogel ^[22]	GelNB/GelSH	LAP (0.05% w/v)	405 nm	Thermally gelated		
Hydrogel ^[22]	PVA-NB/PEG2SH	LAP (1.7–2 mM)	405 nm	Thermally gelated	Sacrificial Gelatin	

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ommunications

projections are blurred.^[26] To correct for this, projected patterns algorithms can be applied to the resulting physically-incorrect patterns to improve print fidelity.^[9,32,33] are set to zero following a non-negativity constraint. Iterative have employed simpler approaches in which negative values instead of photoinitiation.[29-31] So far, however, most works done through interference $\sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_$ ties is physically possible but technically challenging. It may be tions include negative values. Projecting negative light intensiback-projection algorithms.^[27] However; filtered back-projeccan be filtered in the frequency domain, as is the case in filtered cies get more highly sampled than others, and the resulting described by polar or cylindrical coordinates), some frequen-2D images are projected onto a rotating vial (which is better transform or the Fourier slice theorem.^[24,25] Because cartesian the desired object, and they can be calculated using the Radon projected patterns correspond to the tomographic projections of surpass a solidification threshold is solidified [Fig. 1(d)]. The light, only the portion where light dose is sufficiently high to

Optomechanical setup

The optomechanical setup required for tomographic volumetric printing consists of a light source, a light modulator with a projection system, and a rotational stage for the photocurable resin [Fig. 1(e)]. The light source can be either a laser^[10–12,34] or a LED^[9,13,17,18] with a wavelength matching the absorbance spectrum of the used photo initiator. High-power LED are less expensive than laser sources, but their larger *étendue* makes them more divergent and compromises resolution.^[12] 3D ray tracing can be applied to compensate for optical aberrations from divergence and lack of telecentricity.^[35] The light modulator typically consists of a digital micromirror device that displays the calculated patterns synchronously with the rotation of the vial. Some works have also used commercial beamers as projectors.^[36]

Printing occurs within sealed transparent cylindrical vials, unlike open vats common to DLP or SLA. This reduces contamination risks in bioprinting, for example, as well as it contains fumes when volatile resins are used. However, cylindrical vials induce lensing and aberrations, which can be corrected by using a refractive-index matching bath^[12] or through softwarebased corrections of the illumination patterns.^[37] Additionally, imaging systems can be added to the setup to monitor or control the progression of printing. These systems typically use red light to avoid inducing undesired photopolymerization. Boniface et al. included a motorized linear stage to have the resin follow a helical motion in the printer, enabling the fabrication of larger objects still in a layer-less manner.^[38]

Resolution, print fidelity & smoothness: advanced printing strategies

VAM exhibits unprecedent printing speed and enormous versatility across materials. However, the achievable resolution is still limited to above 50 µm. Different strategies to increase print fidelity and resolution have been presented and include

> adaptations to the calculations of the projected patterns,^[32,34,39] optical corrections to reduce aberrations,^[35,37] and feedback from sacrificial prints^[12,40] or live feedback to stop the excitation light [Fig. 2(b) and (c)].^[41] Moreover, refractive-index changes induced by photopolymerization can produce lensing artifacts, including striations via self-writing waveguides.^[42] Such striations degrade print shape accuracy and give VAM-printed parts layer-like effects despite VAM being free from layering. Rackson et al. presented an ingenious strategy to mitigate striations and produce smooth shapes in VAM by flooding the vat with uniform light the end of the printing process.^[43]

The smallest printable feature size is at best limited by the projected image of the DMD micromirrors. Beam divergence further decreases resolution, which is why low-étendue light sources (such as laser diodes) are preferable than high-étendue sources (such as LEDs). This means that resolution can't be any better than the size of the DMD micromirror on the image plane. As an example, Toombs et al. used lower magnification in their micro-CAL setup to demonstrate the fabrication of 3D objects with minimal feature sizes of 20 and 50 µm in polymer and fused silica glass, respectively; albeit at the cost of smaller printable sizes.^[11] The minimal fabricated features were much larger than the projected DMD mirror images in this work. Resolution is further limited due to materials, light deposition, and tomographic calculations, among others (Tables II and III).

Chemical diffusion of free radicals (be it from the activated photoinitiator or growing polymer chains) also reduces resolution. Radical quenchers, such as TEMPO, can be used to limit the detrimental effects of radical diffusion and dark curing.^[44] In addition to this, limited light contrast also hinders resolution.

As light patterns traverse the entire vial's volume, there is light deposited in regions outside of the target volume. Algorithms that optimize light patterns so less light goes outside of the build volume can improved print fidelity.^[39]

Fabricating objects with sub-wavelength features with a purely back-projection approach will be challenging. However, integrating two-photon or two-step absorption into the fabrication process may bridge this resolution gap.

Resin viscosity

Because objects are not fabricated in a layer-by-layer fashion, the resin does not need to flow at each printing step, as is the case in SLA or DLP; more viscous resins can be used in VAM. This has enabled the use of solvent-free formulations, which have higher monomer concentrations and thus polymerize faster and yield stronger objects, and has also enabled the use as solid or gelled materials, like hydrogels^[10,18–22,34,45] and organogels.^[46] Part sedimentation during the printing process depends on print shape and resin viscosity and could compromise print fidelity if not taken into account properly.^[47] Previous works have shown that sedimentation dos not hinder fidelity in viscous resins, as it occurs mostly once printing has finished.[12, p. Supp. Mats.],[16, p. Supp. Mats.] On the contrary, when the polymerization is highly exothermic, solidified objects can float instead of sinking in VAM. Fabricating

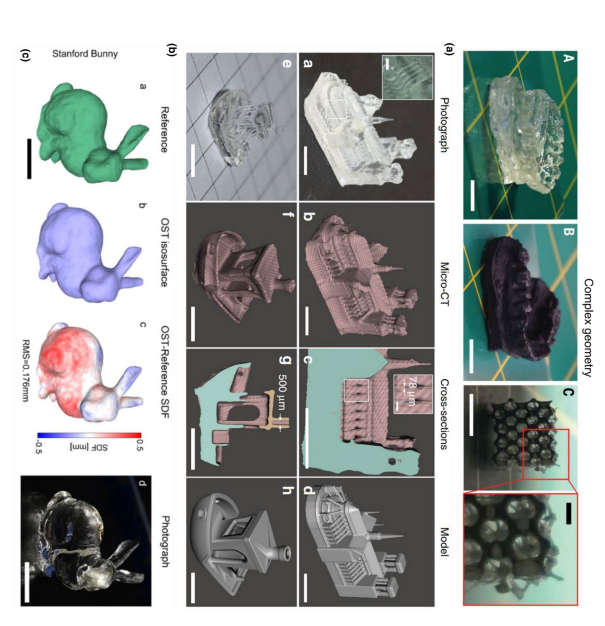


Figure 2. (a) VAM allows for the rapid fabrication of complex geometries into acrylates (scale bars=A.A-B 10 mm, A.C 5 mm, A.D 2 mm; copyright AAAS).^[9] (b) High resolution of positive and negative features is achievable using sacrificial prints (scalebars 5 mm, insets (ba) 1mm, (bc). 0.5 mm; Copyright Springer-Nature).^[12] (c) Optical scattering tomography can be used to determine automatically when to stop the print to maximize fidelity (scale bars=5 mm; Copyright Elsevier).^[41]

objects under microgravity opens the possibility to use less viscous or more exothermic materials.^[48] Viscosity, however, is not a fundamental requirement for VAM, and printing in low-viscosity materials could be done by reducing the amount of time between the beginning of solidification and the end of the print, or by adjusting the displayed patterns to the expected sedimentation.

Optical transparency

As VAM relies on light exciting the full volume at once and not layer by layer, high optical transparency is a requirement for the usable materials. Because of this, photoinitiators must

be used at low concentrations, otherwise light would be rapidly attenuated following the exponential decay described by Beer–Lambert–Bouguer law. Ideal photoinitiators for VAM have low extinction molar coefficients but high polymerization yield. As seen in Table I, BAPO, TPO, and camphorquinone are commonly used initiators in VAM, having molar extinction coefficients in the order of 100–200 L mol⁻¹ cm⁻¹ at the excitation wavelength;^[49] which is orders of magnitude lower than the peak molar extinction coefficients other photoinitiators. Their low concentration is, however, extremely beneficial in some applications such as in bioprinting given the cytotoxicity of most photoinitiators. Light absorbers and dyes, which are

Table II. Fabrication methods combined	Table II. Fabrication methods combined with VAM and their demonstrated applications.	
Combined fabrication method	Materials	Description
Extrusion bioprinting ^[171]	Cell-laden jammed microgels (extrusion)+cell- laden GelMA (VAM)	microgels laden with iß pancreatic cells were extruded into a GelMA hydrogel laden with human mesenchymal stem cells. The surrounding GelMA hydrogel is shaped by VBP
Melt electrowriting ^[23]	Poly-ɛ-caprolactone (MEW)+cell-laden GelMA hydrogels (VBP)	Two-step VBP in human mesenchymal stem cells around a pre-fabricated melt-electrowritten poly- ɛ-caprolactone mesh. The poly-ɛ-caprolactone enhances the mechanical properties of the construct, which can be ultimately seeded with human umbilical vein endothelial cells to pro- duce vein models
Photopatterning ^[22,170]	GelNB (VAM)+growthfactors (photopatterning); PVA hydrogel+fluorescent dyes	GeINB is photocrosslinked into a desired shape by VBP, then growth factors are diffused into the gel and photopatterned onto a desired 3D geometry also through tomographic light projections
Sequential multimaterial VAM ^[9,23,145]	Sequential multimaterial VAM ^[9,23,145] Acrylates around metals; GelNB (different formu- lations); cell-laden GelMA (different cell types)	Overprinting around prefabricated part
Two-photon abalation ^[144]	GeINB	VBP is used to fabricate acellular constructs with hollow cavities (400 µm). Then, microcavities (down to 2 µm) are fabricated into the gel by two-photon ablation

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because they limit the penetration depth of light. common in DLP and SLA, are actually detrimental to VAM

by using longer-wavelength photoinitiators,^[50] upconversion nanoparticles,^[51,52] or multi-photon instead of single-photon excitation.^[5,53,54] putational pipeline.^[34] Possibly, scattering could be reduced by including the scattering profile of the material in the comtive index mismatch within the components of the resin $\ensuremath{^{[11,20]}}$ or effect of scattering has been mitigated by reducing the refraccomputations for the tomographic patterns. The detrimental light from the straight path that it is assumed to follow in the cell-laden hydrogels or composite resins. Scattering deviates Many resin formulations of interest are scattering, such as

cate objects volumetrically in completely opaque materials.^[55] graphically to fabricate 3D objects in opaque media.[56] Acoustic waves could also be used holographically or tomolight, tomographic microwave curing could be used to fabriently subject to lower resolution due to the wave nature of visible spectrum, such as with microwaves. Although inher-Polymerization may be induced with radiation outside of the

Acrylates **Materials used for VAM**

functional domains, polymerize rapidly and in a propagating reaction initiated by a free radical from photoinitiators.^[57] This able mechanical properties. Polyacrylates, having multiple are also highly transparent, even in the near UV, and have tuncoatings, adhesives, and vat 3D printing applications.^[9,12] They their high reactivity, low cost, and availability in commercial VAM was first introduced with acrylated urethanes thanks to

> light dose, which facilitates printing in VAM [Fig. 1(d)].^[59] a non-linear thresholded conversion response of acrylates to scavenges free radicals.^[57,58] Oxygen-mediated inhibition gives chains bind or when molecular oxygen, in dilution in the resin, polymerization reaction terminates spontaneously when two

remain to make the photo-crosslinkable hydrogel edible. for personalized tablet manufacturing, although challenges tablets through VAM within seconds,^[60] opening possibilities hydrogels with paracetamol to simultaneously produce several Pombo et al. loaded polyethylene glycol diacrylate (PEGDA) aids^[12] or dental retainers^[9,38] [Fig. 2(a)]. Also, Rodrígueznumerous industrial applications, like personalized hearing ties for elastomers, challenging materials to 3D print but with retain their shape, but soft enough to open up new possibili-Objects printed in acrylates are typically stiff enough to

Epoxies and orthogonal polymerization

late their mechanical properties spatially enable applications in emerging fields like soft robots and electronics.^[61,62] Two change the glass transition temperature above or below room ing time). For example, illumination can be modulated to dose contrast within the process window (i.e. within the printphotoresponse of the polymer precursor and the established achievable property gradient reflects an overlap between the ing dose and extent of reaction to a property in question. The dose, in combination with a well-defined correlation relatrelies on the spatial accumulation of single-wavelength light polymerization.^[64] In greyscale printing, property modulation a single workpiece: (i) greyscale printing $^{\left[63\right] }$ and (ii) orthogonal strategies are employed to generate property gradients within 3D printers that can not only shape devices but also modu-



	Table III.
	Abbreviations
	used
•	throughout
	this
	text.

lable III. Abbrev	able III. Abbreviations used throughout this text.		
Abbreviation	Definition	Section	Notes
ANPHA	Aluminum N-nitrosophenylhydroxylamine	Thiol-enes	Material (Pot-life stabilizer)
BisGMA	Bisphenol A-glycidyl methacrylate		Material
BPAGDA	Bisphenol A glycerolate (1 glycerol/phenol) diacrylate		Material
CAT2	Triarylsulfonium hexafluoroantimonate salts	Epoxies	Material
CQ	Camphorquinone		Photoinitiator
DIW	Direct ink writing	Sinterable materials	Fabrication method
DLP	Digital light printing		Fabrication method
EDAB	Ethyl 4-(dimethylamino)benzoate	Epoxies	Photoinitiator
EEC	3,4-Epoxycyclohexylmethyl 3,4-epoxycyclohexanecarboxylate		Material
FTIR	Fourier transform infrared	Thiol-enes	Analysis method
GelMA	Gelatin methacryloyl	Hydrogels	Material (hydrogel)
GelNB	Gelatin-norbornene	Hydrogels	Material (hydrogel)
HEMA	Hydroxyethylmethacrylate		Material
LAP	Lithium phenyl-2,4,6-trimethylbenzoylphosphinate	Hydrogels	Photoinitiator
o-CL HABI	2.2'-Bis (2-chlorophenyl)-4,4',5,5'-tetraphenyl1,1'-biimidazole; 1H-Imidazole, 2-(2-chlorophenyl)-1-[2-(2-chlorophenyl)-4,5-di-	Epoxies	Photoinitiator
PCL	Poly(ε-caprolactone)	Thiol-enes	Material
PEG	Polyethylene Glycol	Hydrogels	Material
PEGDA	Polyethylene Glycol Diacrylate		Material
PVA	Polyvinyl alcohol	Hydrogels	Material
Ru/SPS	Ruthenium/sodium persulfate	Hydrogels	Photoinitiator
SF	Silk fibroin	Hydrogels	Material (protein)
SLA	Stereolithography		Fabrication method
SMP	Shape Memory Polymers		Family of materials
SS	Silk sericin	Hydrogels	Material (protein)
TA-ICN	Tri-allyl isocyanurate	Shape memory foams	Material
TAE-ICN	Tris[2-(acryloyloxy)ethyl] isocyanurate	Thiol-enes	Material
TEGDA	Triethylene glycol diacrylate	Thiol-enes	Material
TEGDAE	Triethylene glycol diallyl ether	Shape memory foams	Material
TEGDMA	Triethylene glycol dimethacrylate		Material
TEMPO	2,2,6,6-tetramethyl-1-piperidinyloxy		Material (nitroxyl radical)
THF	Tetrahydrofuran	Epoxies	Material
TMPTA	Trimethylolpropane triacrylate		Material
TME-ICN	(Tris[2-(3-mercaptopropionyloxy)ethyl] isocyanurate	Shape memory foams	Material
TPO	Diphenyl(2,4,6-trimethylbenzoyl)phosphine oxide		Photoinitiator
TPO-L	2,4,6-Trimethylbenzoyldi-Phenylphosphinate (liquid)	Thiol-enes	Photoinitiator
VAM	Volumetric additive manufacturing	Introduction	Fabrication method
VBP	Volumetric BioPrinting. Equivalent to VAM, when applied to	Hydrogels	Fabrication method
	engineering		

temperature, leaving the end product in a rubbery or glassy state, to further increase the achievable contrast.^[63] Dose modulation in TVAM is challenging: the highest dose is capped by the need to minimize over-exposed out-of-part voxels, while the lowest dose is defined by the gelation threshold. The short printing time of VAM limits the achievable hardening, whereas undesired dose accumulation due to full-body illumination limits retainable softness. Nonetheless, stiffness modulation via grayscale printing is achievable using smart algorithm

design, $^{[17,39]}$ or by reversing dose buildup through pseudonegative illumination. $^{[30,31]}$

Introducing orthogonal mechanisms of photo-polymerization adds another dimension to property modulation. Freeradical mediated chain growth is advantageous in VAM because the curing threshold set by pre-dissolved radical quenchers is essential for establishing a desired dose contrast that guarantees the geometric fidelity of reconstruction, and the termination of propagation is amenable to actinic control. In contrast, cationic

true fluidic devices and the peripheral.^[68] adopted the strategy to enhance the interfacing between microusing a DLP-based, dual-wavelength setup.^[67] Schmidleithner cations of orthogonal polymerization in 3D- and 4D-printing rials. Schwartz and Boydston demonstrated promising appliranging from stiff thermoset to much softer hydrogel-like matecan generate a continuous variation in mechanical property the extent of both reactions separately. The cited formulation covalently bonded, allowing for greyscale printing to control initiate acrylate. As a result, two polymers are interlaced but not Brønsted acid that initiates epoxy, and free radical species that without initiating the epoxy group, whereas the opposite is not cursors, showed that blue light could excite the free radical PI Larsen et al., ^[64] using a mixture of hydrogel and epoxide preonic PI when exposed to light of wavelength below 385 nm. The coexisting epoxy monomer was only initiated with a catithe acrylate with a radical photoinitiator (PI) using visible light. acrylate group, and the authors selectively polymerized only ter work, a monomer is functionalized with an oxetane and an explored by Decker,^[65] and later by Ruiter et al.^[66] In the latexposure to different light sources in a one-vat setup was first idea of selectively curing a resin into different materials with lenges for arresting conversion and for post-processing. The sustained chain growth, even in the dark, poses additional chalthat separates in-part voxels from out-of-part voxels, and the polymerization does not offer a threshold-setting mechanism -upon excitation by UV, a cationic PI produces both a

greyscale printing proved challenging for a balanced print fidelity and property contrast via dual-color in their intricate geometries [Fig. 3(g)]. Generating the patterns without support struts [Fig. 3(f)], and with stiffness modulation to fabricate objects with suspending and enclosed structures achieved. The geometric design freedom of VAM was exploited (e)] revealed that an average gradient of 5 MPa μ m⁻¹ could be surface analysis in selected regions of interest [Fig. the printing resolution of the setup used [Fig. 3(c)]. A detailed stiffness changes were created within 300 µm, comparable to in the lateral plane [Fig. 3(b)]. In the radial direction, stepwise greyscale printing to generate a continuous property gradient end product [Fig. 3(a)]. This correlation may be combined with between the UV-to-visible dose ratio and the stiffness of the ents. It was first established that a strong correlation exists $\lambda_2 = 365$ nm) to generate internal mechanical property gradiphotoresins and used a two-color VAM setup ($\lambda_1 = 455$ nm, Larsen et al.^[64] and Kelly et al.^[9] into a wavelength-sensitive dent wavelength. Wang et al.^[17] combined the formulation of free radical quenchers, remains valid regardless of the incishaping, which relies on contrast-building via pre-existing erization, meaning that the principle underpinning geometric exposure of out-of-part voxels. UV triggers free radical polymacrylate. Slow superacid propagation avoids immediate overcals and polymerization of epoxide being slower than that of thanks to photolysis of cationic PI also generating free radi-Stiffness modulation by multiwavelength TVAM is viable 3(d) and

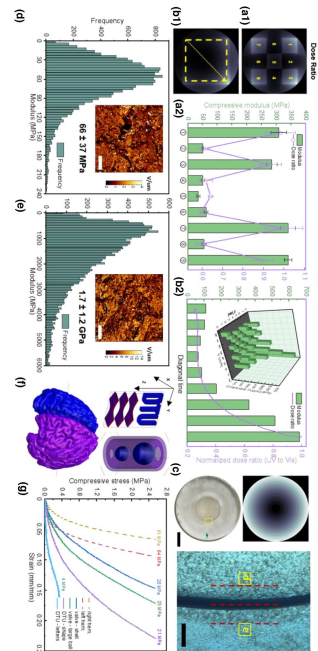


Figure 3. Property modulation via orthogonal photo-polymerization. (a) Stiffness of end-product correlates strongly with UV-to-visible dose ratio. (b) A continuous property gradient can be generated laterally by using greyscale printing to modulate the relative extent of conversion for free radical- and cationic polymerization. (c) The radial resolution of stiffness control is 300 µm or better, comparable to the printan achievable modulus gradient of 5 MPa/µm. (f) and (g) Dual color property modulation can be realized in structures challenging to print using conventional AM methods. (Copyright pringer-Nature).^[17] ing resolution of the same TVAM setup (scale bars=left 3 mm, right 150 µm). (d) and (e) Analysis using atomic force microscope reveals



Sintered materials

Ceramics and glasses are materials with remarkable properties like hardness, thermal resistance, chemical resistance and inertness, and, in the case of glass, optical transparency and refractivity. Thanks to these properties, ceramics and glasses have numerous industrial and technical applications, from everyday kitchenware to implantable prosthetics and insulators for outterspace satellites. It is also because of their high mechanical resistance and brittleness, that these materials are difficult to shape into complex geometries, particulary by means of traditional subtractive manufacturing.^[69,70] Recently, developments in additive manufacturing have opened new possibilities for the fabrication of objects and devices with greater design freedom from glass and ceramics.

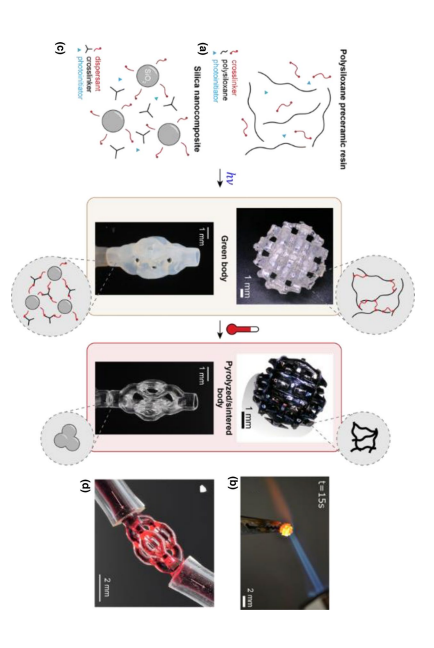
The fabrication of ceramics and glasses requires high temperatures or pressures to give the desired mechanical, chemical, or thermal properties to the material. Plastics and organics are, on the other hand, easy to shape; although they lack mechanical, chemical, and thermal resistance.

In the additive manufacturing of ceramics and glasses, a well-established strategy is to combine the shapeability of

plastics and the resistance of ceramics by fabricating a softer version of the object, also know as the green body. The green body is rich in organics and has low stiffness, but has already been molded to the desired shape. The green body can be formed from polymer derived ceramics or nanocomposite suspensions, as we will cover in this section. Then, it can be pyrolyzed or sintered at high temperatures in a furnace, usually above 800°C, in a process that burns all the organic components and leaves the inorganic ceramic or glass behind, as shown in Fig. 4. Pyrolisis or sintering comes with shrinkage and mass loss; thus reducing cracks from burning the organic part is an active field of study.^[71]

Polymer derived ceramics

The fabrication of polymer-derived silicon-based objects by the pyrolsis of organosilicon polymers was developped in the 1960s.^[69] Since then, numerous ways to pyrolyze these polymers into ceramic parts have been presented and the library of materials has grown from binary systems as SiC and SiN to more complex systems like SiCN, SiBCO, SiBCN, SiBCO, and SiAlCO. Pyrolyzing polymeric precursors requires typically



crosslinker and a photoinitiator is shaped with light in the volumetric printer. After sintering the green body at 1300°C, (d) transparent glass devices, like this microfluidic structure, are obtained.^[11] burnt and evaporated away from the green body, leaving an amorphous inorganic SiOC ceramic part. (b) The transformation makes the object highly temperature resistant.⁽¹⁶⁾ (c) A nanocomposite resin composed of silica nanoparticles coated with a dispersant, an acrylic Figure 4. Volumetric additive manufacturing of ceramics and glasses. (a) To fabricate ceramics, a photo-curable polysiloxane preceramic resin is shaped by exposing it to light in a volumetric printer. The resulting the printed green body, has a cross-linked organic mesh at its microscale. Then, the green body is pyrolsyed at 1000°C. At these high temperatures, the organic moleties of the preceramic resin are

lower temperatures (900–1100°C) than sintering ceramics from powders (1700–2000°C).

During pyrolysis, organic moieties are eliminated by breaking the lower-energy C-H bonds; releasing gases and leading to mass loss and shrinkage.^[69] However, when gases cannot escape rapidly enough from the bulk of the greenbody, pressure increases rapidly thanks to high temperature, which results in the formation of cracks. Cracks are generally undesirable and are responsible for low yield rates of sintering processes. To minimize the formation of cracks, sintered objects are designed to be thin (<1 mm) or porous,^[72] and they are sintered following long, slowly-increasing heating profiles with holding steps at the temperatures of solvent evaporation and polymer decomposition.

Although polymeric precursors can be formed by molding or thermally-induced gelling; the advent of photocurable materials has enabled the shaping of ceramics with higher precision and flexibility. Introduced by Liew et al., the UV photopolymerization of a polysilazane allowed the fabrication of SiCN microelectromechanical systems using lithographic masks.^[73] More recently, Zanchetta et al. demonstrated the stereolithographic fabrication of crack-free SiOC microcomponents from polysiloxanes;^[74] which are insensitive to air and moisture, unlike polysilazanes. The method has been expanded to fabricate multi-cm heat shields using high-area rapid printing (HARP)^[3] or crack-free micrometric injectors using twophoton polymerization.^[75]

a mass loss of 54%. Additional Raman, FTIR, and X-Ray Pho-14) baths. and being immersed in corrosive acidic (pH 2) and basic (pH resistance after being subject to heating cycles up to 1400°C and spherical woodpiles, exhibited high thermal and chemical intrusions.^[76,77] The fabricated components, including screws mediate mixed silicon oxycarbide species and free-carbon organic green body to an inorganic SiOC ceramic with intertoelectron Spectroscopy demonstrated the conversion from an smooth surfaces and an isotropic linear shrinkage of 31% and pyrolysed at 1000°C, yielding dense ceramic componets, with bodies exhibited resolutions down to 80 µm. They were then concentration could be kept low. The resulting printed green The crosslinker reduced printing times while photoinitiator trimethylbenzoyl)phosphine oxide (TPO) as photoinitiator. 1,4-butanediol diacrylate as a crosslinker with diphenyl(2,4,6cally clear resin composed of a commercial polysiloxane and ble to polymer-derived ceramics.^[16] They developed an opti-Kollep et al. demonstrated that VAM could be also applica-

Glass

Traditionally, glass has been shaped by forming technologies like blowing and casting or subtractive methods like grinding and chemical etching. Freeform 3D shape control of glass originated with molten glass filament deposition^[78,79] and powderbased laser sintering.^[80] These, being direct structuring processes, require localized high temperatures to sinter or melt the feedstock at the time of printing. Alternatively, multistep

> processes transform a body of pre-glass polymer^[81-84] or hybrid glass-polymer nanocomposite^[83-87]-shaped by low-temperature 3D printing methods-into high-purity glass with a thermal treatment. These materials are often designed to be compatible with existing 3D printing technologies like direct ink writing (DIW), stereolithography, and digital light printing (DLP). Fidelity can be improved over high-temperature direct structuring; however, for layer-by-layer processing the rheological properties of the liquid precursor material must be optimized. For instance, desirable properties include shear-thinning for recoating in DLP and fast viscoelastic recovery to retain shape after extrusion in DIW. Moreover, complex geometrics may require support structures which can limit geometric freedom and lead to surface artifacts and roughness which may, at best, require further polishing and, at worst, create anisotropic mechanical properties.

including periodic lattices and 3D branched microfluidic pathand smaller defects were present in beams produced by VAM. tion peak-to-valley roughness, both suggesting more uniform modulus of rupture, and smaller average and standard deviashowed enhanced Weibull modulus, i.e. decreased variation in of solid beams printed volumetrically to DLP-printed beams cal performance. Preliminary comparison of flexural strength ness of 6 nm without additional polishing steps, and fair optisystems and microlens arrays exhibited minimum R_a roughsizes of 50 μm and 150 $\mu m,$ respectively. Multielement lens ways demonstrating minimum positive and negative feature for microscale printing. Complex structures were fabricated section Thiol-enes) to improve polymer conversion contrast to volumetric printing by including a radical quencher (see ing;^[11] a nanocomposite formulated for DLP^[85] was adapted demonstration of transparent glass tomographic VAM printrefractive index-matched. This strategy was used in the first nanoparticles and liquid monomer binder-should be nearly the two phases of a particle-loaded precursorwavelength is a necessary condition for tomographic VAM, ing. As sufficient material transparency at the digital projection irradiation times than bulk polymerization without solid loada sol to solid object thus requiring less dose and thus lower nents of medium molecular weight to sharply transition from formation. Furthermore, high particle loading allows compohigh-viscosity precursor liquids, and layer-less volume-at-once printing. Lack of fluid motion permits high solids loading, Volumetric printing has important advantages for glass -solid silica

Early results show the potential of volumetric printing for glass components, but sub-millimeter maximum cross-sectional thickness and single-material silica composition could impede adoption for certain important applications. Crucial to formulations are proper dispersion of silica into resin, index matching of silica to resin, tuning of solvents that enable drying without cracking, and development of a suitable polymer binder system. Prior resin formulations used in VAM and DLP achieve this through the use of monofunctional monomers, such as hydroxyethyl methacrylate (HEMA), that serve as a polymer

an alternative strategy when mitigation of light scattering is tive indices up to 1.58.[83,84,90] High-index sulfur-containing for phase-modulated volumetric printing. in scattering biological tissues,^[92] could be transferred to VAM ing, which has proved indispensable for deep photostimulation fidelity for low concentrations of $\mathrm{TiO}_2^{[34]}$ and wavefront shapof amplitude projections has resulted in improved VAM print not possible. Software-based light scattering compensation propagation models which account for disordered media offer may be required for still higher refractive index glass. Light length into infrared until a match can be found or even into microwave where scattering will be significantly reduced^[55] ing glasses up to $n_D \approx 1.7{-}1.8.$ Red-shifting the actinic wavemonomers^[91] could be a candidate for nanocomposites containand Ag nanoparticles can enable tuning of glasses with refracwill be challenging. Alternatively, doping with TiO₂, GeO₂, glasses, like metal oxide and chalcogenide glasses $(n_{\rm D} > 2)$, ily available monomers, index-matching high refractive index the refractive index of SiO₂ is easily accomplished with read-Al₂O₃, B₂O₃, AlNaO₆Si₂, TiO₂, and PbO.^[89] Although matching compositions or even gradients of multiple oxides including refractive index lenses, and optical fibers require different oxide applications like scratch or impact resistant glass, gradient Beyond single-material silica composition, advanced glass be optimized to avoid shrinkage-induced stress and warping. parts with more uniform drying, though monomer choice must mers and oligomers mitigates this effect, thus enabling larger the drying process. Polymerization of multifunctional monoproducts. Low molecular weight products can greatly disrupt scale parts results in a wide distribution of the molecular weight polymerization of monofunctional monomers in centimeterbinder while also enabling proper mixing of silica through the formation of a solvation layer.^[85,88] However, the non-uniform

the sol-gel process^[94] represent a second class of transformmetric printing of high index glasses as the refractive index until after printing. This approach could be beneficial for voluafter the printing process, formation of solids can be delayed inducing the condensation reaction via elevated temperature other 3D printing processes like DIW.^[83] More generally, by composition in tomographic VAM that comes naturally with properties will be important for advanced spatial control of controllable pore size [96] This type of greyscale control over photopolymerization-induced phase separation and intensityboron alkoxide precursors and organic monomers that achieved for DLP 3D printing composed of silicon, phosphorous, and cessing is rich for exploration, especially for volumetric print-ing of multicomponent glass.^[95] Moore et al. introduced a resin ymer derived ceramics);^[16] however, the field of sol-gel prooxycarbide ceramics has already been reported (see section Poltomographic VAM, using preceramic precursors to print silicon challenges of the nanocomposite approach. In the context of printing which may circumvent the refractive index matching able materials compatible with photopolymerization-based 3D control of composition and creation of glass and ceramic by Inorganic alkoxide precursors^[93] which allow deliberate

Thiol-enes

tered radical to a thiol.^[107] Thiol-ene networks are significantly more uniform than those formed by acrylates,^[106,108] exhibiting them particularly suited to optical applications. generally have high optical clarity and refractive index, making homogeneous gels, elastomers and glasses.^[109] These materials much narrower glass transitions temperature ranges, forming radical propagation and the chain transfer from a carbon-cencontrolled by the ratio of kinetic constants between the thiyl common to acrylates.[104-106] Polymerization rates are strongly stress and shrinkage in comparison to chain growth processes growth process, delaying gelation to dramatically reduce curing enes. The regioselective propagation reaction follows a step paired with a similarly large family of electron-rich or strained sible polymers is vast because a large library of thiols can be toclick reactions, $[^{98,99]}$ the step-growth thiol-ene mechanism represents the most widely used. $[^{100,101-103]}$ The family of poswith non-toxic end products and mild running conditions.[97] istry defining modular, orthogonal highly efficient reactions In 2001, Sharpless et al. introduced the concept of click chem-When it comes to photochemical click reactions, or simply pho-

Thiol-ene polymers also readily support additional processes after photopolymerization that can modify their properties. Since the step-growth reaction proceeds by alternation to nearly full conversion with no homopolymerization, addition of a stoichiometric excess of thiols or enes results in remaining reactive groups for later reaction stages.^[110] The thiol-thioester exchange reaction or the disulfide addition–fragmentation chain transfer reaction can locally transform the elastic thermoset network to exhibit viscoelastic plastic flow.^[111] Finally, thiol-ene polymers can be engineered as high performance shape memory materials with excellent shape fixity and rapid actuation.^[112]

systems to suppress gelation outside of the desired print challenging, since this inhibitory threshold is used in acrylate ity of thiol-ene reactions to oxygen makes VAM printing more molecular weight thiol monomers. Conversely, the insensitivin a closed volume, alleviating the odors that arise from low hydrogels).^[15,19,22] VAM resins are also completely contained volumetric bioprinting (see section Hydrogels & cell-laden viability and biodegradability in constructs fabricated through patibility compared to acrylate-based systems, improving cell photoclick chemistries may potentially have higher biocoming. Additionally resins and hydrogels made through these fast as fast as 1-10 s, matching the potential rate of VAM printticularly when using strained enes such as norbornene, can be Advantageously, the rapid reaction times of thiol-enes, parin volumetric additive manufacturing and one disadvantage. Thiol-enes have several specific advantages when used

volume. Additionally, similarly to the large refractive index changes seen in (meth)acrylate chain-growth polymerization reactions, refractive index changes during thiol-enes polymerization can result in self-focusing waveguide formation during VAM,^[43] impacting print quality.

of 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) as a radical scavenger were found to be sufficient.^[13] TEMPO, however, material and mechanical properties.^[15] choice of thiol and ene functionality directly impact the bulk material toolkit for VAM, as simple changes to monomer and thiol-ene polymer networks opens up a massive modifiable behavior, providing enticing avenues to target soft robotic and biological applications.^[13–15] Ultimately, the uniformity of the als have improved biocompatibility and unique shape memory growth acrylate counterparts.[13] Additionally, thiol-ene materitunable and robust mechanical properties compared to chainprinting of these thiol-ene resins confirmed the expected highly ranging from 44 to 450% depending on the formulation. VAM orders of magnitude (421 to 0.12 MPa), and ultimate strains flexible ene monomers, with elastic moduli spanning three systems are highly tunable in relation to the ratio of rigid and acrylate compositions, the mechanical responses of thiol-ene generation of the tomographic projections. Unlike comparable ric energy dose and gelation threshold for printed resins for photo-FTIR analysis was used to identify the critical volumet-(a triethylene glycol chain (TEG)) ene monomers for tunable mechanical response.^[13,14] Photorheological and real-time ing rigid (a trifunctional isocyanurate ring (ICN)) and flexible were formulated with trifunctional thiol monomers and vary-With these considerations in mind, thiol-ene resins for VAM and room temperature and improve handling for printing.^[14] found to increase stability of the resin at elevated temperatures pot-life. Additions of pot-life stabilizers such as ANPHA were sity to thermally cure under ambient conditions, reducing vat in adapting thiol-ene polymerizations for VAM is their propenphoto-dosing relative to reaction kinetics. Another challenge O_2 in acrylate systems [Fig. 5(a)], requiring careful tuning of threshold provided by TEMPO is "softer" than that seen with competing effects must be carefully balanced. In addition, the also retards the overall cure speed and delays gelation, so these ene polymerizations for VAM, small concentrations ($\sim 0.1 \text{ mM}$) In order to induce the necessary thresholding within thiol-

Among the unique material properties identified in thiol-ene VAM materials, shape memory behavior is appealing for developing responsive or active materials. Shape memory polymers (SMP) can be physically deformed to a "temporary" shape, then recover a more "permanent" state in response to external stimuli such as temperature, pH, light and electromagnetic fields. The most easily programmed and well-studied of these is temperature, as shape memory behavior is easily controlled by heating the shape memory polymers above its glass transition temperature (T_g) .

Schwartz et al. adapted thiol-ene formulations to fabricate devices with facile, controlled thermal programming of shape memory polymers behavior.^[14] The trifunctional thiol

> isotropic, unlike traditional layer-based AM methods where print orientation during printing can imbue anisotropies. [113]. structures without layering, the shape memory behavior was their permanent printed configuration. As VAM produces 3D the $T_{\rm g}$ again, the gripper and tripod structures recovered to deforming the structure before cooling. Upon heating above temporary shape through heating above the T_g to 80°C and (Fig. 5). After printing, structures were programmed to their three-arm gripper and self-standing tripod structures by VAM these characterized thiol-ene materials, the authors printed a etitions at strains up to 18.4%, similar to skeletal muscle. Using components exhibited nearly full shape recovery over four repstrain and $T_{\rm g}$. Dynamic mechanical analysis showed that the functional ene monomer-subunits to modulate ultimate tensile varied the amount of flexible and rigid bifunctional and tricomponent was kept constant in their photoresins; yet they

Photoclick thiol-ene chemistry has also been applied to volumetrically print poly(*e*-caprolactone) (PCL) constructs with improved mechanical properties and lower brittleness than most acrylate cross-linked counterparts.^[15] PCL is degradable and biocompatible, making it the material of choice for many resorbable synthetic implants in the past decades. Thijssen et al. have shown that volumetrically printed thiol-ene PCL constructs, including lattices and branching tubular structures (Fig. 5), were non-toxic and biocompatible in vitro and in vivo. Volumetrically printed PCL could be a promising candidate for the generation of complex cell-free 3D implants.

Hydrogels and cell-laden hydrogels

only after printing, the term biomaterial inks is instead com-monly used.^[116] The unique ability of biofabrication techniques as biotechnological products.[121] biological research,[120] as well as engineered living materials alternatives to animal experimentation for pharmaceutical and regenerative medicine,[119] the generation of tissue models as Key applications include the production of living grafts for salient functions of native organs and biological systems.^[118] to engineer tissues that mimic the anatomical composition and and biomaterials, is being leveraged by biomedical scientists to control the spatial positioning of cells, bioactive molecules bioresins (when used for extrusion or light based printing tech-niques, respectively).^[116,117] If cells are added to the hydrogel of biofabrication, cell-laden hydrogels are termed bioinks or cellular matrix (ECM) present in living tissues. In the context tionality, acting also as temporary analogues of the native extrathe embedding of cells and preserve their viability and funcgels, aqueous networks of hydrophilic polymers that allow for materials. Typically, these materials are in the form of hydrohave been engineered to process also living cells and cell-laden The past two decades saw the emergence of the field of bio-fabrication,^[114,115] in which additive manufacturing techniques

In this context, the bioprinting process needs to be carefully designed not to harm cell integrity, viability, and health during and after printing. Extended printing times (from tens of minutes to hours), necessary to fabricate centimeter-scale,



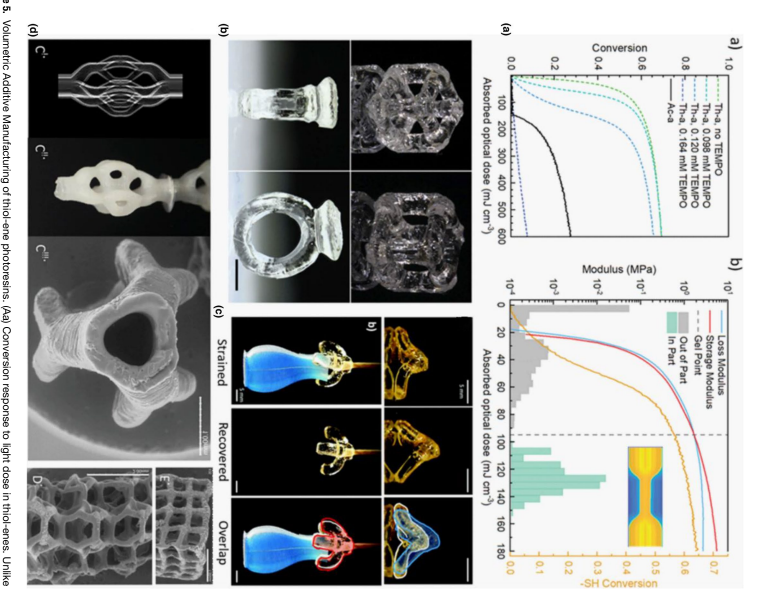


Figure 5. Volumetric Additive Manufacturing of thiol-ene photoresins. (Aa) Conversion response to light dose in thiol-enes. Unlike acrylates, thiol-enes are not subject to polymerization inhibition from molecular oxygen; instead, by adding TEMPO, a radical scavenger, conversion can be thresholded. (Ab) Mechanical properties (storage and loss moduli) of thiol-enes are tunable along a large dynamic range. (b) Resulting prints, exhibiting overhangs and cavities (scale bar = 5 mm, Copyright Wiley).^[14] Thiol-ene printed parts can be thermally treated to exploit shape memory effects, as in this tripod and gripper (scale bars = 5 mm, Copyright Wiley).^[14] light patterns used to generate (cii) poly(ɛ-caprolactone) microfluidic devices and (d) and (e) geometric lattices (scale bars = 1 mm, 2 mm & 1 mm, respectively; Copyright Wiley).^[15]

clinically-relevant structures are characteristic of conventional layer-by-layer bioprinting methods; which may impair cell functionality if cells are kept outside of their optimal culture environment for too long.^[122] Addressing this drawback of classic additive manufacturing approaches, Bernal et al. first introduced the concept of Volumetric Bioprinting (VBP), generating cell-laden, functional tissues in few seconds, by means of tomographic printing.^[10] VBP is simply VAM applied to fabricate cell-laden constructs for tissue engineering and tissue modelling. As photo-responsive bioresin component, gelatin methacryloyl (GelMA) was used for this first study, and since then, the library of materials available for volumetric bioprinting has rapidly expanded. The following section will review the main materials and chemistries applied in and developed for this printing technology.

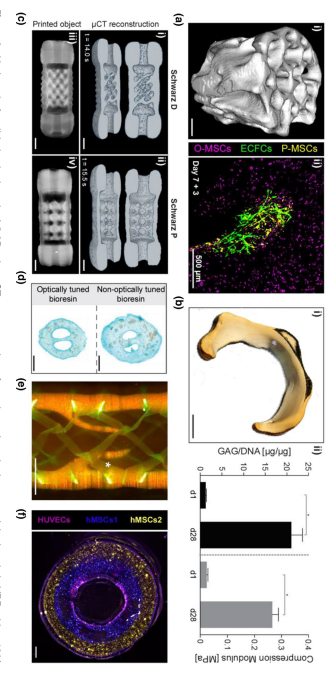
GelMA

Gelatin, a biopolymer obtained from collagen denaturation, has been widely used as biomaterial for tissue engineering and bioprinting. This is due to the material's biocompatibility, controllable degradation profile, and its ability to promote cell adhesion and several cellular functions.^[10] While gelatin undergoes physical gelation when cooled below body temperature, at 37°C the polymer is fully soluble in aqueous solutions. Therefore, thermostable hydrogels with several crosslinking

> modalities can be obtained, for instance through the introduction of photo-responsive moieties.

The first report on the synthesis and utilization of (meth) acryloyl modified gelatin, named GelMA, appeared in the year 2000.^[123] In the presence of a photoinitator, the (meth)acryloyl moieties undergo free-radical polymerization, rapidly forming a covalently crosslinked hydrogel, whose mechanical profile can be tuned by varying the GelMA pre-polymer content and the degree of methacryloyl substitution.^[124] Most notably, GelMA is readily accessible to many research labs through commercially available products or in-house synthesis, following well-described protocols.^[124,125] Moreover, medical-grade, endotoxin-free material processing routes have been established, therefore facilitating the potential translation of GelMA-based constructs towards pharmaceutical and medical products.^[126] As such, GelMA rapidly became one of the most widely used bioinks for extrusion printing.^[117,130]

In 2019, Bernal et al., developed a GelMA-based bioresin for tomographic volumetric bioprinting, using lithium phenyl-2,4,6-trimethylbenzoyl-phosphinate (LAP) as visible-light type I initiator (0.037% wt in PBS).^[10] The system facilitated the generation of anatomical, centimeter-sized trabecular bone models embedding mesenchymal stromal cells (MSC) in 12.5 s, with high cell viability (>85%), comparable to non-printed controls [Fig. 6(Ai)] The bioprinted cells remained functional



valve structure (red) throughout an opaque melt electrowritten tubular mesh (green) (scale bar=2 mm). (f) Multi-cellular, multi tubular construct reinforced with tubular melt electrowritten meshes (dotted line) (Scale bar=500 μ m). Adapted from Ref. 23. micro-CT reconstructions and (ii) digital images (scale bars = 2 mm). (d) Stereomicroscopy images of non-optically tuned 5% GelMA and iodixanol-supplemented bioresins for organoid VBP (scale bars = 2 mm). Adapted from Ref. 21. (e) VBP of GelMA bioresins into a venous GelMA bioresin) cultured for 28 days and (ii) exhibiting significant increases in neocartilage matrix production and compressive proper-ties. Adapted from Ref. 10. (c) Organoid-laden VBP of convoluted, mathematically derived Schwarz D and P structures as shown in (i) a early endothelial cell infiltration into the printed hydrogel (scale bar=500 µm). (b) Digital model of a VBP-printed meniscus model (10% GelMA bioresin (scale bar=2 mm). (aii) MSC/ECFC co-culture seeded into the pores of the MSC-laden trabecular bone model showing Figure 6. Hydrogels & cell-laden hydrogels: (ai) 3D micro-CT reconstruction of a native trabecular bone model printed via VBP with a 10% 2 mm). (f) Multi-cellular, multi-layered in (i) 3D

laden GelMA over 42 days.^[21] sion of osteoblastic and osteocytic markers in bioprinted MSCcontext of bone tissue engineering, demonstrating the expres-Long-term culture was also later shown by Gehlen et al., in the ble to that of human knee menisci [~ 0.3 MPa; Fig. 6(Bii)].^[10] endowed the constructs with compressive stiffnesses comparacolonized the hydrogel with fibrocartilage-like matrix, which Fig. 6(Bi)]. Exposed to chondrogenic media, ACPCs readily printing 10 million cells mL⁻¹ into a meniscus-shaped construct articular cartilage progenitor cells (ACPCs) was shown, by biosibility and potential of VBP, long-term (28 days) culture of the original STL files. Moreover, further underlining the feawere experienced, with $\sim 95\%$ volumetric fidelity compared to pended in the bioresin, and no artefacts due to sedimentation gel form prior to printing, cells remained homogenously sus-[Fig. 6(Aii)]. By converting GelMA into a thermoreversible lial cells and pericytes, was observed in the printed hydrogel and ingrowth of capillary-like structures from seeded endothe-

shown for constructs laden with up to $1.5 \cdot 10^7$ cells mL⁻¹ ^[20] organize to capture organ-like behavior. The nozzle-free, shear tumor microenvironment.^[45] duct and studied the inflammatory response of these cells in a where they seeded human pancreatic epithelial cells inside the et al. fabricated fibroblast-laden duct-like structures in GelMA, ins [Fig. 6(d)]. In another soft tissue application, Sgarminato a simple and versatile tool applicable for all types of bioresadditional corrective strategies, iodixanol supplementation is While further increasing the cell content will likely require resolution achieved in the presence of cells to date, were thus densities. Defect-free prints with 40 µm-features, the highest scattering can cause on printing quality when using high cell fore minimizing the negative effects that cell-mediated light could be tuned to match that of intracellular organelles, thereible compound iodixanol,^[132] the refractive index of GelMA supplementing the bioresin formulation with the biocompat-Notably, in the same study, Bernal et al. demonstrated that leveraged to tune the functionality of engineered tissues.^[20] underlining how bioprinting and its freedom of design can be could be varied to tune the rate of ammonia conversion to urea, flow experiment [Fig. 6(c)]. The geometry imposed by VBP the ability to detoxify ammonia over the course of a dynamic ble to express key functions of the native liver, as shown by printing,[131] into complex perfusable hepatic constructs capawhich otherwise tend to fragment during extrusion-based biostress-free nature of VBP allowed to process the organoids, are miniaturized, multicellular 3D structures in which cells selffor VBP have also been optimized to culture organoids, which low stiffness (compressive modulus < 2 kPa) GelMA bioresins Besides offering a platform to engineer connective tissues, The thermoreversible behavior of GelMA prior to covalent

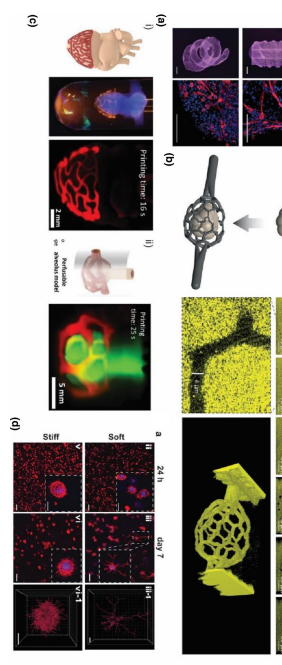
The thermoreversible behavior of GelMA prior to covalent (photo)crosslinking is also beneficial to print composite (bio) materials, in which pre-formed objects are embedded within the bioresin, and secured in place once the thermal gel is formed. Groessbacher et al., utilized this concept to demonstrate the possibility of sculpting (cell-laden) GelMA in presence of

> and vascular formation, [133] and the further functionalization even unmodified proteins and peptides. used for the volumetric printing of tyraminated polymers, or tyramine-like groups onto the polymer, [136] it could also be GelMA, exploiting either the native tyrosine residues, or adding tion of covalent bonds between aromatic side chains in gelatin. trigger di-tyrosine oxidation, therefore catalyzing the formatiator.^[134,135] This ruthenium complex, a type II initiator, can bipyridyl)ruthenium (II)/sodium persulfate (Ru/SPS) as iniof pristine, unmodified gelatin were printed utilizing tris(2,2'ric printing of GelMA perfusable gyroids, within which plugs Soliman et al., recently reported the multi-material volumetwith other (light-responsive) chemical groups. In this line, degrees of methacrylation to facilitate in-gel cell migration material (mammals, fish, or recombinant), exploring different constructs can also follow the selection of the source gelatin ies.^[23] Further optimization of volumetrically-printed GelMA mating the mechanical performance of porcine coronary artercomposite construct, as shown with vessel-like prints approxiing strategy for modulating the mechanical properties of the stiffer thermoplastic polymermeshes also constitutes a promisbination of GelMA as a bioprintable matrix for cell culture with were obtained via volumetric printing [Fig. 6(e), (f)]. The com-(venous valves, branched structures, partly occluded vessels) and structures with varying degree of architectural complexity were produced mimicking the architecture of blood vessels, opaque, poly(*e*-caprolactone) fiber meshes produced by melt electrowriting.^[23] Multi-material and multi-cellular prints While this photochemistry could be used in combination with

GeINB and photoclick biopolymers

process leads to the formation of heterogeneous networks and hydrophobic kinetic chains that are non-biodegradable. This time and the potentially cytotoxic radical initiating species. [139] networks are formed faster, thus reducing the light exposure Due to higher reactivity and insensitivity to oxygen, thiol-ene acryloyl derivatives such a GelMA and PEGDA).[104,109,138] tages when compared to chain-growth systems (i.e., (meth) mechanism comes with a series of biologically relevant advanall, the use of the photoclick thiol-ene step-growth crosslinking makes it highly reactive upon formation of a thiyl radical. Overand methacryloyl moieties); while its ring-strain conformation to Michael-type addition (unlike the more traditional acryloyl the NB functionality has the advantage of being insusceptible groups, the use of norbornene (NB) offers significant advan-tages for tissue engineering and bioprinting.^[99,137] In particular, the most widely used in this field. Among the possible -ene click reactions, the step-growth thiol-ene mechanism represents As introduced in Sect. "Thiol-enes", in the context of photovolume shrinkage, which are undesirable phenomena. In contrast, chain-growth polymerization occurs by creating

In the context of VBP, thiol-NB chemistry has been first introduced by Rizzo et al. with a photoresin composed of gelatin norbornene (GelNB) and 4-arm-PEG-thiol (PEG4SH).^[19] As previously discussed, gelatin possesses both mechanical and **Figure 7.** VAM of GeINB hydrogels. (a) VBP of complex geometries in the presence of myoblasts showing spreading and differentiation into myotubes (Myosin Heavy Chain: red, Nuclei: blue) Scale bars: left 2 mm, close-ups right 200 µm^[19] (b) Two-photon ablated micro-channels connecting cavities printed by VAM (Copyright Rizzo et al.).^[144] (c) Algorithmically designed organ-specific auxetic meshes and perfusable networks volumetrically printed around pre-existing constructs. (Copyright Chansoria et al.).^[145] (d) Cell spreading in VBP printed PVA-NB hydrogels of varying stiffness (scale bars: 100 µm and close-ups 50 µm copyright Wiley).^[22]



print around a preexisting shape,^[9,23,145] GelNB/GelSH photoclick bioresin was also employed by Chansoria et al. to produce organ-specific auxetic meshes wrapped around a heart as well as perfusable geometries enveloping an alveolus model [Fig. 7(c)].^[145]

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ties of tissue VBP to fabricate constructs matching the mechanical properthis method has only been applied to DLP, it may be used with molecular cleavage was introduced by Wang et al. [146] Although softness upon removal, another strategy based on post-printing the use of sacrificial material to ensure good printability and biofabricating soft tissues with structural complexity. Besides is foreseen to play a central role in tackling the challenge of PVA-based network upon thermal removal. In this regard, VBP as sacrificial material to leave behind a soft, cell permissive nating sedimentation-induced artifacts, and at the same time fied gelatin to guarantee reversible thermal gelation, thus elimihol (PVA) derivatives [22] The authors used chemically unmodithe library of VBP bioresins by introducing NB polyvinyl alcorial choice for VBP, a recent work from Qin et al. has expanded Although gelatin has so far prevailed as the preferred mate-

and cell migration capabilities.^[141-143]

Although thiolated PEG is a commonly used bioinert thiol-

biological desirable properties for VBP and thus represents an ideal material platform for photoclick crosslinking. Thanks to the faster crosslinking of photoclick chemistry, printing times for centimeter-size objects in this GelNB/PEG4SH bioresin were reduced to ~ 10 s [Fig. 7(a)]. Lower light exposures and generation of radical species resulted in excellent cell viability of embedded cells (>95%). In addition, thiol-NB crosslinking made it possible to combine such printing performances with a lower gelatin degree of substitution and concentration, allowing to retain the native polymer properties, ^[123,140] and to generate softer matrices known to result in enhanced nutrient diffusion

Volumetric printing has only started to take advantage of the potential of photoclick reactions. Besides thiol-ene, several other photoclick reaction mechanisms, comprehensively reviewed elsewhere,^[98,99,137] remain currently unexplored. It is also worth mentioning that the library of possible photoclick reaction mechanisms goes beyond light-triggered freeradical mechanisms (reactions requiring a photoinitiator generating radical species), with photo-triggered uncaging,^[147]

first work, VBP printing of perfusable constructs was reported with an overall polymer concentration of only 2.5% w/v.^[144] Interestingly, the use of a gelatin-only photoresin made the resulting hydrogel well susceptible to two-photon ablation (2PA); converging VBP and 2PA for the first time. Using this hybrid method, the authors overcame the resolution limit of VBP and created multiscale complex perfusable geometries with features spanning from hundreds (VBP) to few microns (2PA) [Fig. 7(b)]. Leveraging VBP's unique capability to 3D

to obtain a purely gelatin-based VBP photoresin $^{\left[144,145\right] }$ In a

ene crosslinker, the literature provides methods for synthesizing and functionalizing thiol- polymers of various types, including synthetic and naturally-derived polymers. This presents an opportunity to customize the thiol-ene photoresin properties further, biophysically and biochemically. For example, GelNB has been recently combined with thiolated gelatin (GelSH)

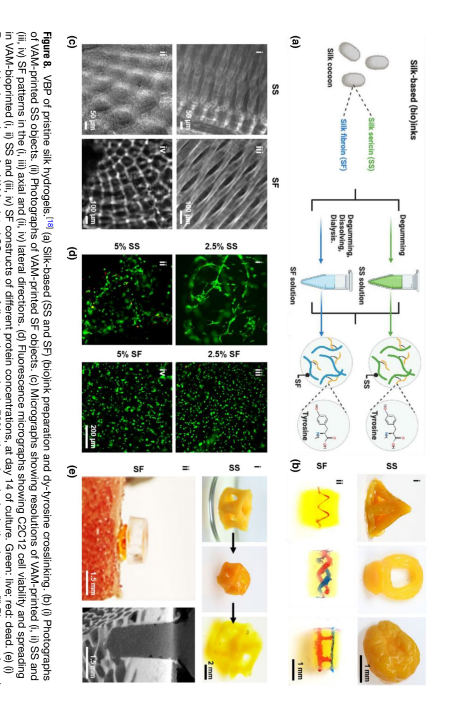
photodimerization,^[148–150] and photoregulated hydrazone/imine formation,^[151,152] presenting radical-free alternatives.

Silk-based hydrogels

cessing bioprinting to exploit on-demand photocrosslinking to pattern cell-laden silk constructs in 3D.^[157] Photocrosslinking is usually achieved by functionalizing SF^[158–160] or SS^[160,161] ties.^[153–155] Reports of applying silk proteins, in particular printing or stereolithography/digital light processing-enabled the formation of dityrosine bonds.^[162] For either extrusion biolight photoinitiator Ru/SPS that facilitates crosslinking through bioink in pristine form due to the development of the visibleit has been suggested that SF can also be used directly as a ture-building in the presence of photoinitiators. More recently, with methacryloyl groups, followed by photoactivated strucbioinks to support cellular functions^[156], or digital light prorial is used alone or in combination with other biomaterials as reports range from extrusion bioprinting where the biomate-SF, towards additive biomanufacturing, are abundant. These different amino acids but of very different ratios and proper-(SF) and silk sericin (SS), both of which are composed of 18 cated Bombyx mori; its main components include silk fibroin Silkworm silk is a natural protein produced by the domesti-

> light-based bioprinting, the rapid formation of 3D silk constructs with sophisticated architectures usually requires protein concentrations > 5%.

a few minutes. The printability range for SS was 2.5-5% SS axial feature size of ~ 57 μm was attainable; in comparison, of printed SS reached 45.9 µm at minimum, while the smallest additive manufacturing of silk proteins. The axial resolution cated the high resolution obtained with tomographic volumetric blocks for SF [Fig. 8(B-ii)]. Resolution assays further indithe brain-like structures for SS [Fig. 8(B-i)], and vascularized nal architectures could be printed, such as a pyramid, ring, constructs containing sophisticated external shapes and inter-0.125-1 mM of Ru. Under optimized conditions, volumetric while that for SF was slightly broader at 1.25-15% SF and paired with 0.25-1 mM of Ru (Ru/SPS ratio was kept at 1/10), were volumetrically printed within a few tens of seconds to both SS and SF [Fig. 8(a)].^[18] Both unmodified SS and SF method was expanded to pristine silk biomaterials, including SPS photoinitiator system, the bioink pool for this bioprinting as modification-free photocrosslinking enabled by the Ru/ requirements with the structural complexity attainable, as well ric additive manufacturing to decouple mechanical property Inspired by the unique ability of tomographic volumet-



remur.

Reversable shape-change for VAM-printed SS structure following immersion in 70% ethanol and rehydration in water; (ii) Photograph and micro-CT image showing ex vivo implantation of VAM-printed SF screw double-crosslinked by immersion in 70% ethanol in a porcine

the lateral resolutions were generally at a poorer resolution, 108 µm for SS and 124 µm for SF [Fig. 8(c)].

Moreover, when cells were embedded in the inks, bioprinting of ultrasoft tissue constructs otherwise not easily obtained with other bioprinting strategies were achieved. When C2C12 myoblasts were encapsulated within the SS and SF bioinks for tomographic volumetric bioprinting, the cells showed high viability throughout 14 days of culture [Fig. 8(d)]. Interestingly, the cells within the bioprinted SS constructs at 2.5% exhibited significant spreading [Fig. 8(D-i)], indicating that the ultralowconcentration SS protein concentrations frig. 8(D-ii)], as well as SF at the same or higher concentrations [Fig. 8(D-ii)]. A similar result was obtained with NIH/3T3 fibroblasts.

be used for fixing hard bone tissues [Fig. 8(E-ii)]. screws in the dry state were sufficiently robust that they could nol (β -sheets only). Thus, printed and double-crosslinked SF higher than those for constructs crosslinked only with ethatreatment (dityrosine bonds+β-sheets), also significantly post-printing (dityrosine bonds only) to > 200 Mpa posttreatment; compressive moduli were elevated from <1 Kpa was more extensive in the printed SF constructs upon ethanol immersed in water [Fig. 8(E-i)]. In contrast, the crosslinking back to its original size and then swell to a larger size when the shrinkage was reversible, where the structure could expand from ~ 1 to ~ 40 kPa, associated with shrinkage in volume, while compressive modulus of the printed 5% SS constructs increased of these hydrogels. For example, upon ethanol treatment, the tent was induced, thus enhancing the mechanical properties post-printing, such as by immersing in ethanol, β-sheet condityrosine bonds to form hydrogels. With secondary treatment constructs, whether SS or SF, only contained photocrosslinked biomaterials towards biomedicine. The as-printed silk protein constructs further expanded the potential applications of these Finally, the tunable mechanical properties of the printed silk

General perspectives for hydrogels and bioprinting

In recent years, the research field of 3D bioprinting and tissue engineering has seen a trend towards the use of high celldensity bioresins (tens, hundreds of million cells mL^{-1}),^[163] which diverges from VBP current capabilities (≤ 15 million cells mL^{-1}).^[20] Although the use of organoids,^[20] refractive index matching compounds,^[20] and optimized algorithms to limit light scattering^[34] have improved VBP performances, there is a major gap with more established methods such as extrusion printing or DLP.^[164] Near-infrared light could be used to increase penetration depth and alleviate the detrimental effects of scattering from high cell densities.

The precise 3D positioning (or patterning) of bioactive cues (i.e., small molecules, peptides and growth factors) within hydrogels has been traditionally obtained with lengthy twophoton triggered processes,^[165–169] VBP technology offers the possibility to rapidly distribute light doses in a 3D manner,

> thus triggering the patterning process in few seconds and for much larger volumes. The concept has been recently demonstrated by thiol-NB reactions,^[22] even to photograft complex patterns of growth factors and chemoattractants (i.e. vascular endothelial growth factor) while preserving their biological functionality,^[170] and is foreseen to become a major biological application of VBP in the near future.

As with every printing method, VBP comes with pros and cons, and while the technology is rapidly improving, a few groups have started to explore possible convergence with other printing methods, from melt electrowriting,^[23] to two photon ablation,^[144] and patterning.^[22,170] Table II lists examples of works which have integrated multiple fabrication methods with VAM/VBP and summarizes the demonstrated applications. Integrating fabrication methods leverages their strengths and mitigates their shortcomings. Exemplifying this, the work of Ribezzi et al.^[171] combines extrusion and volumetric bioprintlow structures, respectively. The team used the two methods to fabricate densely islets laden with pancreatic β -cells that were extruded into a hydrogel laden with human mesenchymal stem cell and volumetrically bioprinted into a geometry mimicking pancreatic ducts.

In addition, VBP has so far mainly been used with single bioresins, therefore limiting the ability to resemble the complexity of human tissues that are made of different cell types and material compositions with regional distribution. Chansoria et al., Groessbacher et al., and Ribezzi et al. have demonstrated approaches for multimaterial and multicellular VBP, ^[23,145,171] opening new opportunities for VBP. Additionally, heterocellular structures have been fabricated through sequential VBP and cell injection in cartilage, bone and pancreatic models.^[10,21,45] Moreover, although not yet investigated in the presence of cells, the 3D stiffness gradients that can be obtained with localized consecutive exposures or dual wavelength exposures^[17] may add a pivotal level of biophysical complexity to the printed tissues.

Finally, from a material perspective, we believe that VBP will align with the most recent trends in the field and broaden its bioresins choice beyond gelatin, PEG and PVA with more biologically relevant systems such as collagen, decellularized ECM and viscoelastic networks featuring dynamic bonds.

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Data availability

Data sharing is not applicable to this article as no new data were created or analyzed in this study.

Declarations

Competing interest

FK is shareholder of Glassomer GmbH which is commerciallizing 3D printed glass. YSZ consults for Allevi by 3D Systems, and sits on the scientific advisory board and holds options of Xellar, neither of which however, participated in or biased the work. RM is a shareholder of Vitro3D which is commercializing volumetric additive manufacturing. RM has filed related provisional patents 63/483, 147 and 63/181,645. RL is scientific advisor for Readily3D SA, which commercializes volumetric printers. CM is a shareholder of Readily3D SA. CM and JMW have filed provisional patent No. WO2022EP63245 20220517 regarding three-dimensional printing in complex media. All other authors declare no competing interests.

Supplementary Information

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