

Three-dimensional printing with sacrificial materials for soft matter manufacturing

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Three-dimensional (3D) printing has expanded beyond the mere patterned deposition of melted solids, moving into areas requiring spatially structured soft matter—typically materials composed of polymers, colloids, surfactants, or living cells. The tunable and dynamically variable rheological properties of soft matter enable the high-resolution manufacture of soft structures. These rheological properties are leveraged in 3D printing techniques that employ sacrificial inks and sacrificial support materials, which go through reversible solid–fluid transitions under modest forces or other small perturbations. Thus, a sacrificial material can be used to shape a second material into a complex 3D structure, and then discarded. Here, we review the sacrificial materials and related methods used to print soft structures. We analyze data from the literature to establish manufacturing principles of soft matter printing, and we explore printing performance within the context of instabilities controlled by the rheology of soft matter materials.

Introduction

A major challenge in developing effective biomedical technologies is the difficulty of shaping hydrogels,^{1–4} biopolymer networks,^{5,6} silicones,^{1,7–9} and cells^{10–12} into finely detailed three-dimensional (3D) structures. Generally, these soft materials must be shaped while in a liquid state before solidifying within seconds or minutes for biopolymer gelation,^{13–15} or within several days for cells producing the extracellular matrix.¹⁰ The need to create high-resolution structures from these soft materials has driven 3D printing technology far beyond the traditional practice of liquefying, extruding, and resolidifying solid materials, entering a paradigm of shaping liquids in 3D space. The challenge of shaping liquids has led to a convergence among new printing technologies—they often leverage materials that exhibit large, reversible rheological changes resulting from small physical or chemical perturbations. This behavior is a defining characteristic of soft matter as a class of material.¹⁶ Beyond its typically low elastic modulus (usually below 10 MPa), this sensitivity of soft matter to

small perturbations has been critical for developing new materials for bioprinting and for improving 3D printing of soft materials in general.

Until recently, it was practically impossible to reproducibly form soft materials into complex 3D structures at high spatial resolution; manufacturing processes resembled art more than manufacturing.¹⁷ Manufacturing soft structures has been enabled by new methods and materials that leverage the unique properties of soft matter, which we review here. Significant progress has been made using “sacrificial” materials that are not ultimately part of manufactured structures, but leverage the highly responsive rheological properties of soft matter. We therefore limit the scope of this article to sacrificial inks and support materials. With sacrificial inks, a temporary structure is printed, surrounded by a permanent material, and then removed to create hollow structures (**Figure 1a–c**). With sacrificial support materials, a permanent structure is printed directly into a sacrificial material that acts as a support matrix during the curing or maturation of the manufactured structure (**Figure 1d–g**).

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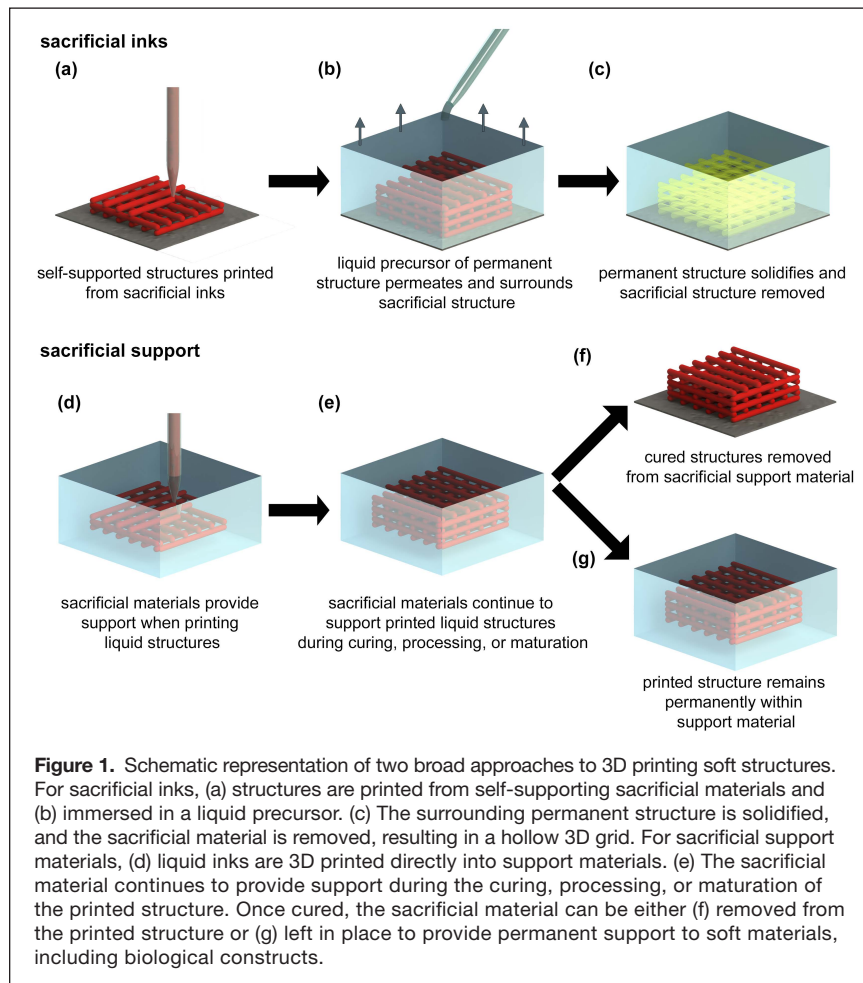


Figure 1. Schematic representation of two broad approaches to 3D printing soft structures. For sacrificial inks, (a) structures are printed from self-supporting sacrificial materials and (b) immersed in a liquid precursor. (c) The surrounding permanent structure is solidified, and the sacrificial material is removed, resulting in a hollow 3D grid. For sacrificial support materials, (d) liquid inks are 3D printed directly into support materials. (e) The sacrificial material continues to provide support during the curing, processing, or maturation of the printed structure. Once cured, the sacrificial material can be either (f) removed from the printed structure or (g) left in place to provide permanent support to soft materials, including biological constructs.

Whenever possible, we identify the dominating rheological properties of these materials and discuss their consequences on manufacturing capabilities. We also describe the rheological concepts that allow differentiating and categorizing soft matter manufacturing materials. These basic concepts help predict the manufacturing capabilities of different soft materials by comparing stabilizing rheological properties and destabilizing external forces such as gravity, interfacial tension, and inertia. Thus, we identify scaling laws that relate the rheology of soft matter 3D printing materials to the potential instabilities associated with manufacturing processes and their resulting limitations.

Soft matter 3D printing principles Rheological foundations

Recently developed sacrificial soft materials for 3D printing include jammed granular particles,^{1,7,9,10,18} entangled polymer solutions,¹⁹ micelles packed into solid-like phases,^{20,21} and polymer networks with reversible bonds.^{22,23} These materials are designed or formulated to undergo large rheological changes that facilitate the printing process; some transition between solid and fluid states while others are better described as shear thinning fluids. Fundamental rheological concepts and simple

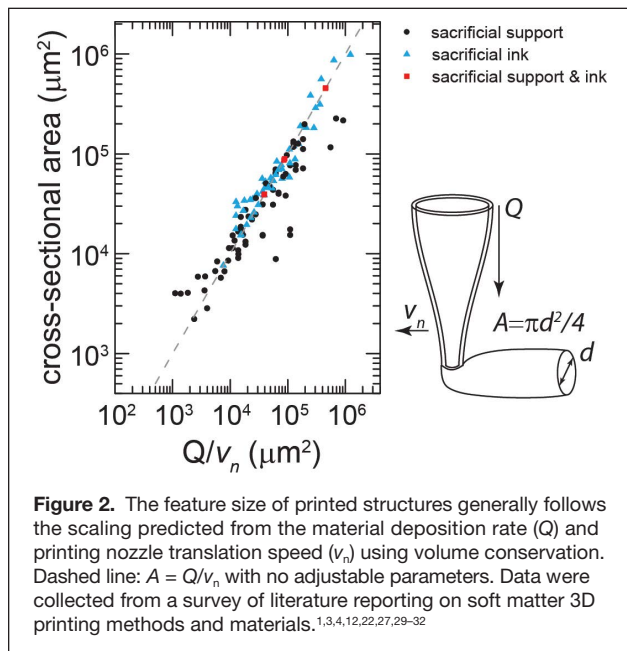
models are useful tools for describing and categorizing these diverse materials, although such an in-depth discussion is outside the scope of this article. Thus, we have provided a “Supplementary material” section that describes basic rheological concepts and models and applies them to current soft matter 3D printing materials. There, we explain how concentrated micelles^{24,25} and polymer networks with weak reversible bonds^{22,23} often behave like liquids at long time scales, while jammed granular microgels behave dominantly like elastic solids.^{1,9,10,26,27} These diverging material properties control how different soft matter 3D printing materials yield during printing and hold their shape after printing, which we elaborate on next.

Feature size and material deposition rate

There are many standardized metrics of quality in additive manufacturing—accuracy is how closely a printed part matches its design; precision is the repeatability of a printing process; a feature is the discrete unit of positive or negative volume created in a printing process, such as a filament or a drop.²⁸ In surveying the soft matter printing literature, we use the cross-sectional area of single features as the standard for comparing different methods and materials. A key question of any soft matter printing method or material combination is whether the cross-sectional

feature area, A , can be predicted using simple volume conservation from the material deposition rate, Q (dimensions of volume per time), and printing nozzle translation speed, v_n , given by $A = Q/v_n$. To answer this question, we identified publications in which A , Q , and v_n were reported or could be inferred. This simple scaling law holds for more than three orders of magnitude in A and Q/v_n , representing a general guideline for soft matter printing in practice (Figure 2).^{1,3,4,12,22,27,29–32} However, the spread in the data indicates that printing quality also depends on the details of each printing approach. For example, measurements of A can deviate from the volume conservation prediction if printed inks intermix with their support materials or if inks swell or contract following printing. Both scenarios may occur with soft matter printing approaches in which inks and support materials share the same solvent that can exchange between the two phases. It would be useful to investigate whether spontaneous phase separation assists in preventing inks and support materials from mixing.

The direct correlation between measured and predicted feature size can be used to quantify the balance between total printed volume, nozzle translation speed, and feature size. Approximating the feature cross-sectional area as circular,

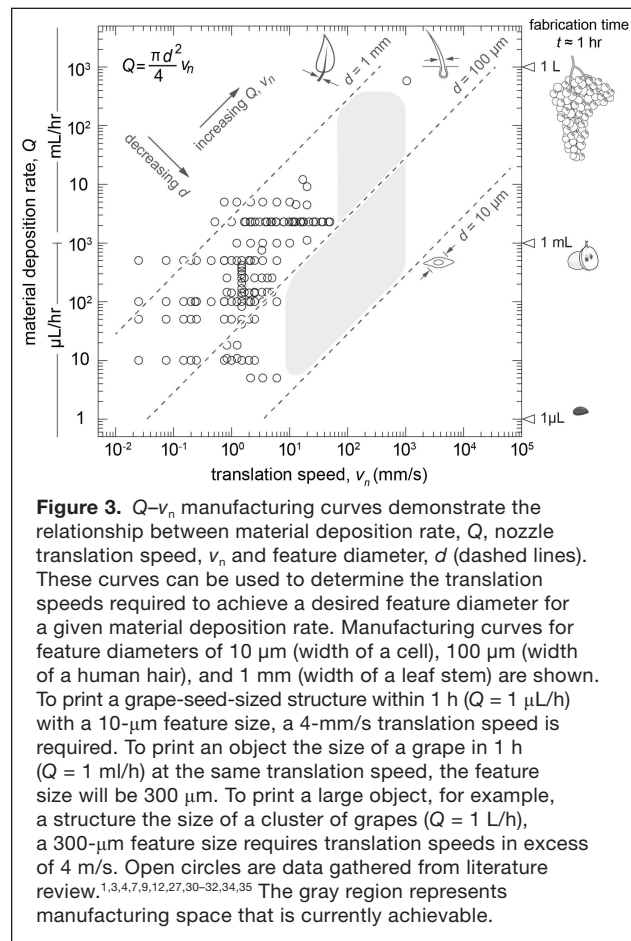


the material deposition rate can be written as $Q = \frac{\pi d^2}{4} v_n$, where

d is the feature diameter. Thus, any targeted feature diameter selects out a combination of material deposition rates and nozzle translation speeds that constitute a single manufacturing curve in $Q-v_n$ space. We plot a family of such curves corresponding to feature diameters between 10 μm and 1000 μm , along with data collected from the published literature (Figure 3).^{1,3,4,7,9,12,27,30–32,34,35} By applying a time constraint for printing an object of a given size, the speed–diameter tradeoff can be determined. For example, to print an object with 1 ml volume in 1 h, a 400 mm/s translation speed is required to achieve a feature diameter of 30 μm ; at a modest speed of 35 mm/s, a feature diameter of only 100 μm is achievable. This problem becomes unmanageable for large volumes; printing a volume of 1 L with 100- μm features in 1 h requires translations speeds exceeding 30 m/s. By contrast, 100- μm features are easily achieved at print speeds of 10 mm/s, but the required time is about 150 days. The tradeoff between time, volume, feature size, and speed illustrates the conundrum faced in tissue and organ printing, where the goal is to manufacture large objects with small features over short times.^{27,33}

Instabilities: Competition between soft matter rheology and confounding forces Inertial forces and Reynold's instabilities

An obvious approach for achieving fine features in manufacturing large objects over short times is to rapidly translate printing nozzles. However, inertial instabilities may arise at high speeds, generating unpredictable flows that reduce printing quality (Figure 4a). Instabilities controlled by competing viscous and inertial forces can be predicted by



estimating the Reynold's number. For a cylindrical nozzle of diameter d , dragging through a fluid with density ρ and viscosity η at speed v , the Reynold's number is given by $Re = \rho v d / \eta$. For flow around a needle translating through a support material, recirculation in the wake emerges at around $Re = 10-15$,^{36–38} setting an upper bound on the speeds and needle sizes suitable for quality printing. High-speed printing within a sacrificial microgel support was performed at $v = 1.05 \text{ m/s}$, where recirculating instabilities emerged at Re between 3.7 and 17, consistent with unstable flow around a cylinder.^{27,38} In these tests, the ink viscosity controlled the Reynold's number at the needle tip.²⁷ As high-speed printing methods emerge, a combination of small needle diameters and high fluid viscosities will help avoid inertial instabilities, increasing print quality.

Dynamic and static crevasse formation

When printing at high speeds, the reflow of support material in the nozzle's wake may be slow, opening a transient crevasse that follows behind the printing nozzle (Figure 4a). This transient crevasse instability arises from the competition between the hydrostatic pressure that drives fluid reflow, $\rho g h$, and the viscous stresses resisting reflow, $\eta v / d$. In this dynamic equilibrium, v is both the reflow speed and nozzle translation

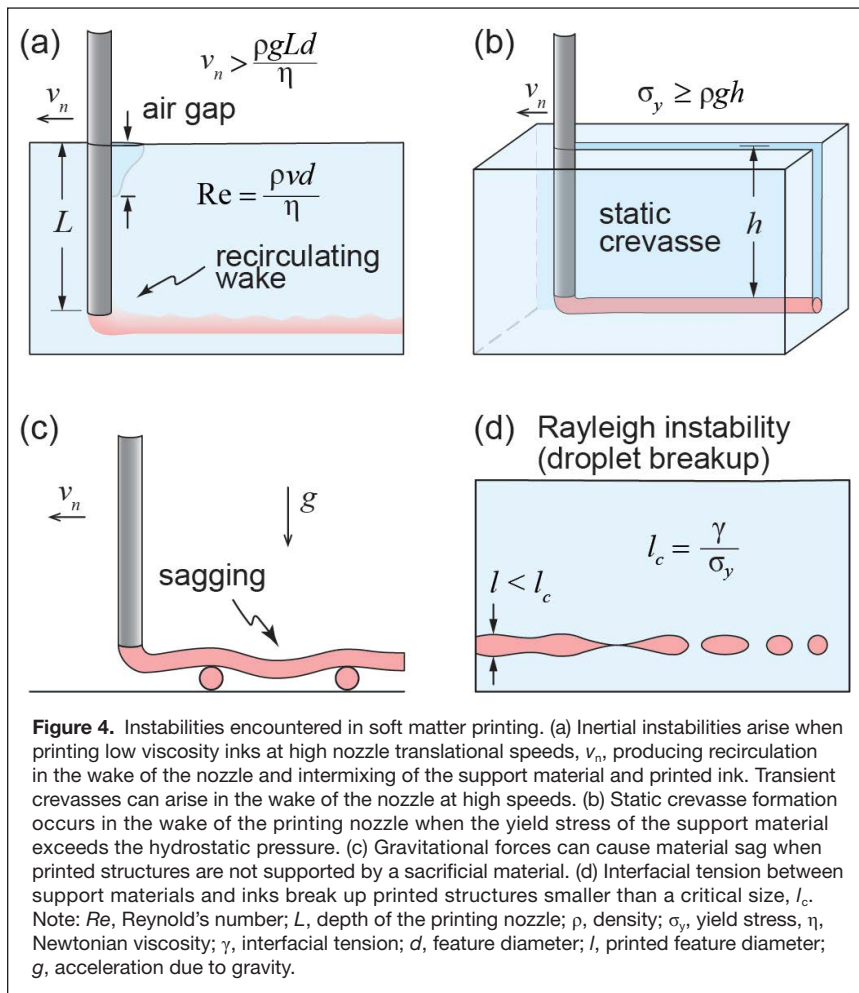


Figure 4. Instabilities encountered in soft matter printing. (a) Inertial instabilities arise when printing low viscosity inks at high nozzle translational speeds, v_n , producing recirculation in the wake of the nozzle and intermixing of the support material and printed ink. Transient crevasses can arise in the wake of the nozzle at high speeds. (b) Static crevasse formation occurs in the wake of the printing nozzle when the yield stress of the support material exceeds the hydrostatic pressure. (c) Gravitational forces can cause material sag when printed structures are not supported by a sacrificial material. (d) Interfacial tension between support materials and inks break up printed structures smaller than a critical size, l_c . Note: Re , Reynold's number; L , depth of the printing nozzle; ρ , density; σ_y , yield stress, η , Newtonian viscosity; γ , interfacial tension; d , feature diameter; l , printed feature diameter; g , acceleration due to gravity.

speed, d is the nozzle diameter and gap width, h is the crevasse depth, ρ and η are the density and viscosity of the support medium, respectively, and g is the acceleration due to gravity. These transient crevasses were reported for printing into a microgel support material,²⁷ though in principle, they can arise with most soft matter support materials. These transient near-surface crevasses are unlikely to reduce printing quality until the crevasse grows to meet the nozzle tip and material deposition occurs at the air–support interface. Therefore, v should be kept below $\rho g L d / \eta$, in which L is the submerged depth of the printing nozzle. An alternative to limiting v is to increase L with a longer needle, although the resulting increased needle deflection will reduce printing accuracy.

Even in the limit of zero nozzle translation speed, crevasses can emerge (Figure 4b). These static crevasses are produced when the hydrostatic pressure at a depth, h , is less than the yield stress of the support material, described by $\rho g h \leq \sigma_y$. In many viscous liquids and materials with low yield stresses, these static crevasses do not arise.^{1,7,9,10,22,27,29} When they do emerge, crevasses can be filled with a secondary support liquid that continuously flows into the gap.³⁹

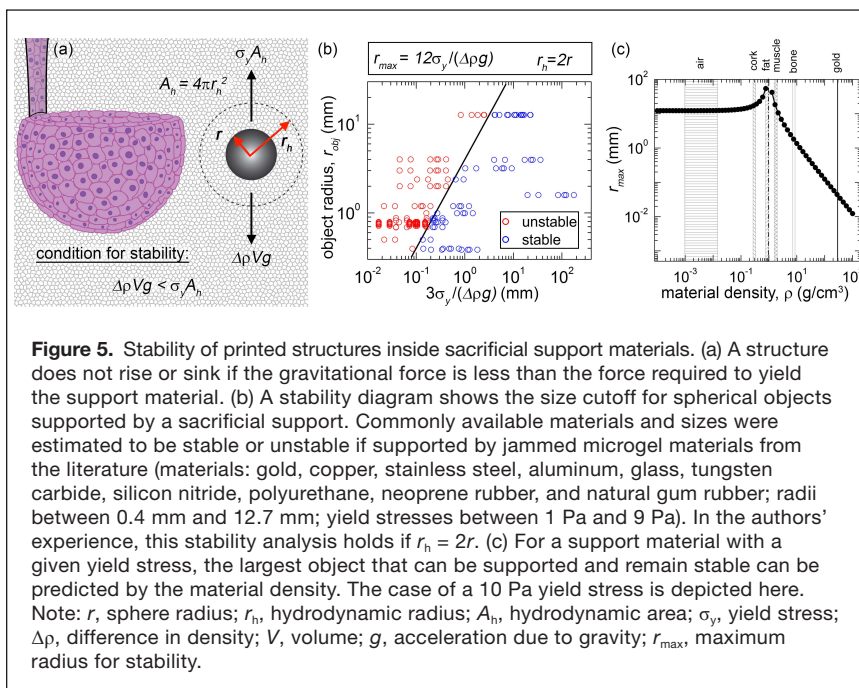
Gravitational instabilities

Gravity often causes printed structures to sag; a traditional method for overcoming sag is to simultaneously print sacrificial support structures and permanent structures (Figure 4c).^{2,6,11,40,41} The support must solidify rapidly after printing and wash away gently without damaging the permanent structure. This process was demonstrated with support materials, including poly(ethylene) glycol (PEG), poly(ϵ -caprolactone) (PCL), and alginate.^{2,11,41} Sacrificial inks that become rigid have also been developed; pre-printed sacrificial structures can be submerged into a liquid precursor of a permanent material that is cured. The sacrificial ink is then removed, leaving channels that can be lined with living cells, for example.^{21,32,34,35,42–46}

Gravitational instabilities can also be overcome by 3D printing directly into a sacrificial support material, reducing buoyancy forces by matching the densities of inks with sacrificial liquid baths.^{12,30,31} Density matching requirements can be estimated by equating the destabilizing buoyancy force of a 3D printed sphere to the resisting Stoke's drag, given by $\Delta \rho V g = 6 \pi \eta r v$, where $\Delta \rho$ is the density mismatch, V is the sphere volume, g is the acceleration due to gravity, η is the support liquid viscosity, r is the sphere radius, and v is the speed at which the sphere rises or sinks.⁴⁷ To prevent a $d = 1$ mm sphere from rising in 1 h by its own diameter through an aqueous solution

($\eta \approx 1$ mPa s), the printed structure must have a density within $4.5 \times 10^{-4}\%$ of water. Thus, even using viscous support materials, alternatives to density matching may better reduce the role of buoyancy forces in soft matter printing.

An alternative to density matching is to print directly into sacrificial support materials having a finite yield stress at long time scales, such as jammed granular materials and polymer networks.^{1,7,9,10,22,27,29,39} This approach provides gravitational stability with dramatically reduced limitations on material density. The stability of an object supported by a material with yield stress, σ_y , can be predicted by comparing the gravitational force to the yielding force, given by $\Delta \rho V g < \sigma_y A_h$, where $\Delta \rho$ is the difference between printed and support material densities, V is the printed object's volume, g is acceleration due to gravity, and A_h is the hydrodynamic surface area (Figure 5a). For 3D printing spheroids, this stability condition predicts a maximum radius for stability, $r_c = \beta^2 \sigma_y / \Delta \rho g$, where β is the hydrodynamic radius scaling coefficient given by $r_h = \beta r$ (Figure 5b). For a soft matter printing support material of known yield stress, a chart can be generated to estimate the size limits on stable printing with inks of different densities (Figure 5c). A support material will stabilize a larger object made from, for example, low-density fatty tissue than from higher-density bony material.



Interfacial forces and the Rayleigh–Plateau instability

Immiscibility between support materials and printed inks provides a diffusive barrier, prevents their mixture, and promotes smooth surfaces with consistent cross-sectional areas. However, these interfacial forces can destabilize structures through a mechanism similar to the Rayleigh–Plateau instability (Figure 4d). The balance between destabilizing interfacial forces and stabilizing yield stress creates a minimum stable feature size, l_c , given by $l_c = \gamma/\sigma_y$, where γ is the interfacial tension between the support material and ink, and σ_y is the support material's yield stress.^{48–51} Instabilities arising from interfacial forces are overcome when printing large features into materials with large yield stresses.⁷

An additional concern when printing with immiscible pairs is the accumulation of nanoparticles or microparticles at the ink–support interface.⁵² Similar to Pickering emulsions, in which micro- or nanoparticles stabilize droplets and prevent coalescence by accumulating at the droplet surface, filler material in the printed ink will accumulate at the interfaces between printed features and immiscible support material.^{53,54} This Pickering effect can reduce adhesion between printed filaments.⁷ One potential solution is to develop new support materials that remain immiscible with inks, but exhibit a drastically reduced interfacial tension.⁹

Thixotropic instabilities: A new opportunity in soft matter 3D printing

A characteristic of most sacrificial support materials is the tendency to rapidly recover zero-shear rate rheological properties after yielding or shear thinning (see the “Supplementary material” section). This recovery upon the removal of stress arises from differing underlying processes in different materials,

yet is broadly called thixotropy; the corresponding time scale is the “thixotropic time.” When 3D printing into sacrificial support materials, returning to the same location before thixotropic relaxation has occurred may reduce printing quality. Thixotropy in sacrificial support materials were reported and commented on previously,^{1,9,22} however, additive manufacturing path-planning principles that leverage this rheological property have not been developed. We envision new nozzle path-planning protocols that prohibit revisiting locations until the thixotropic time has elapsed, ensuring that the support material is always in the same rheological state, locally, when printing.

Conclusion and outlook

One goal has dominantly motivated the invention of new materials and methods in soft matter manufacturing—to 3D print functional tissues and organs for implantation. The manufacturing charts produced here corroborate the current view that soft matter printing technology

is still far from reaching this goal,³³ which may be achieved more quickly by establishing manufacturing principles that help to predict the performance of new materials and methods. In the immediate term, current levels of manufacturing speed, precision, accuracy, and rheological stability provided by soft matter 3D printing methods and materials are sufficient to make significant impacts in mesoscale biomanufacturing applications. For example, microscale to millimeter-scale structures made from living cells and extracellular matrix materials can now be 3D printed rapidly and reproducibly.^{10,21} These structures can be used to discover new drugs, screen for compound toxicity, and study new therapeutic approaches before conducting animal or patient trials. Moreover, these approaches can be used to perform basic research on cell behavior in 3D printed microtissues, which is needed to elucidate the principles that will one day enable large-scale tissue and organ fabrication.

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Three-dimensional printing with sacrificial materials for soft matter manufacturing

Christopher S. O'Bryan, Tapomoy Bhattacharjee, Sean R. Niemi, Sidhika Balachandar, Nicholas Baldwin, S. Tori Ellison, Curtis R. Taylor, W. Gregory Sawyer, and Thomas E. Angelini

Rheological concepts

Jammed granular particles,¹⁻⁵ entangled polymer solutions,⁶ micelles packed into solid-like phases,^{7,8} and polymer networks with reversible bonds^{9,10} are among the materials that have been investigated for the manufacture of soft structures. These different categories of soft matter exhibit unique and complex frequency-dependent behaviors, however, they are often described by classical viscoelastic models within bounded frequency ranges. While simple classical models do not perfectly capture the complex behavior of these materials, they can provide a framework for predicting performance in three-dimensional (3D) printing applications. The two simplest frameworks for describing the rheology of viscoelastic materials are the Maxwell and Kelvin–Voigt models.¹¹⁻¹⁴

Maxwell materials:

Low-frequency fluids/high-frequency solids

Soft matter printing materials that behave like solids at high frequencies and liquids at low frequencies may be described by the Maxwell model (Figure S1a). Even in the limit of zero applied stress and zero strain, these materials are fluids over long time scales and are often referred to as “Maxwell fluids.” The stress relaxation curve of these materials is modeled by a spring and dashpot arranged in series, given by $G(t) = G_0 e^{-t/\tau}$, where $G(t)$ is the ratio of stress to strain as a function of time, often called the stress relaxation modulus. The frequency-dependent elastic shear modulus (G') and viscous shear modulus (G'') of Maxwell materials can be obtained by Fourier transforming the stress relaxation curve and are given by:

$$G'(\omega) = G_0 \frac{(\omega\tau)^2}{1 + (\omega\tau)^2} \quad \text{and} \quad G''(\omega) = G_0 \frac{(\omega\tau)}{1 + (\omega\tau)^2}, \quad (1)$$

where G_0 is the high-frequency elastic modulus, ω is the frequency, and τ is the characteristic structural relaxation time. Generally, materials exhibiting thermally driven relaxations,

including spontaneous bond-breaking, particle rearrangements, or polymer entanglements, will behave like Maxwell fluids at long time scales. In soft matter 3D printing, materials that appear to behave like Maxwell fluids at low frequencies include concentrated micelles^{15,16} and polymer networks with weak reversible bonds.^{9,10} Micelles have a counterintuitive pairing of structure and rheology; they exhibit fluid-like behavior at low frequencies even when concentrated into packings with crystalline symmetry.^{15,16}

Kelvin–Voigt materials:

Low-frequency solids/high-frequency fluids

In direct contrast to Maxwell fluids, soft matter printing materials that exhibit elastic behavior at low frequencies and viscous behavior at high frequencies may be described by the Kelvin–Voigt model (Figure S1b). In the Kelvin–Voigt model, the elastic and viscous shear moduli are given by $G'(\omega) = G$ and $G''(\omega) = \eta\omega$, where G is a frequency independent shear modulus and η is a simple Newtonian viscosity. These moduli can be obtained by Fourier transforming the stress relaxation curve of a spring and dashpot in parallel.^{11,12} The Kelvin–Voigt model captures the elastic modulus of soft materials that do not spontaneously restructure under thermal forces, such as covalently cross-linked hydrogels or soft jammed granules (particle diameter larger than 1 μm). However, these materials generally exhibit non-Newtonian dissipation, so the model's G'' scaling law often fails to describe material behavior. In soft matter printing, jammed granular microgels have been shown to exhibit the frequency-independent elastic modulus of the Kelvin–Voigt model and behave dominantly like elastic solids, even in the low-frequency limit.^{1,3,4,17,18}

Applying rheological models to soft matter printing materials

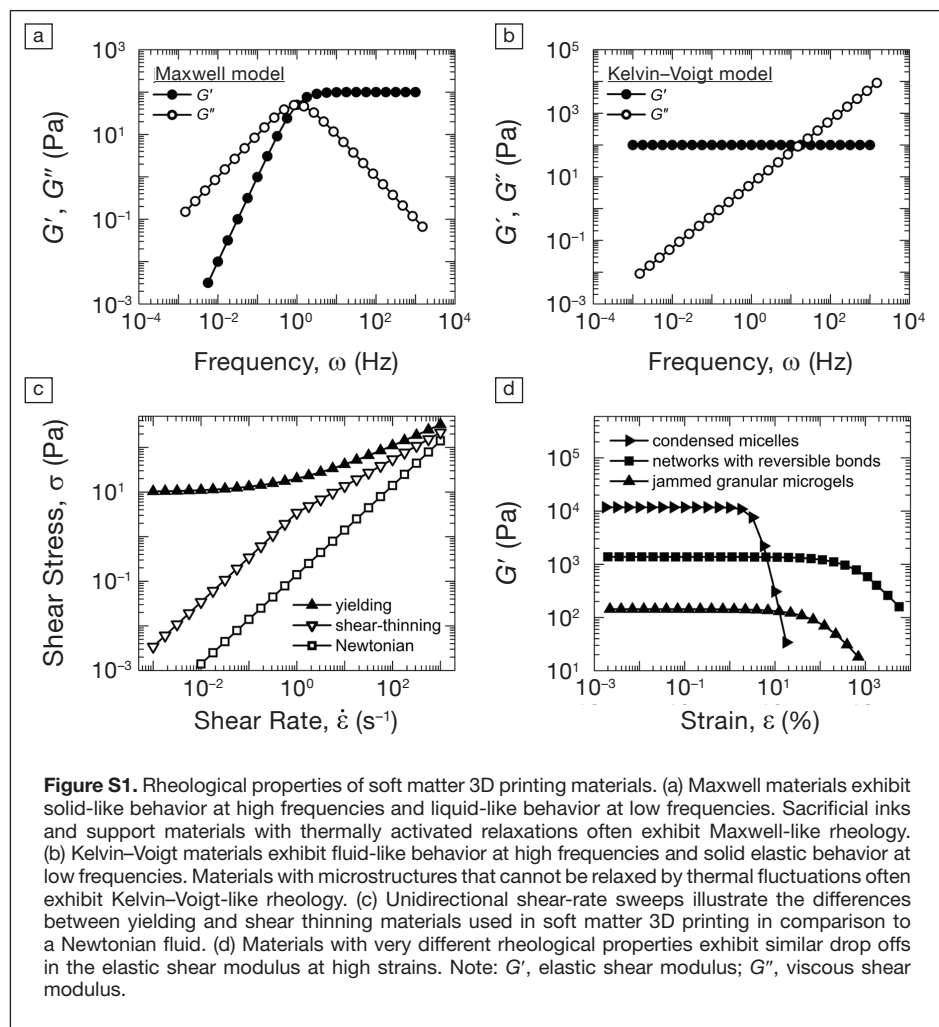
The dramatically different low-frequency behaviors of these two classes of material are reflected in their responses to applied strain in the zero-frequency limit. The zero-frequency

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limit can be probed by performing unidirectional shear-rate sweeps in which stress is measured as a function of shear rate (Figure S1c). For example, at low shear rates, jammed granular microgels bear shear-rate-independent stresses and are dominated by elasticity; at high shear rates, these materials fluidize and are dominated by viscous losses.^{17,19} This behavior is captured by $\sigma = \sigma_y + k\dot{\epsilon}^p$ where σ is the shear stress, σ_y is the yield stress of the material, $\dot{\epsilon}$ is the shear rate, and p is a dimensionless constant; $p = 1$ corresponds to a Bingham plastic and $p < 1$ corresponds to a Herschel–Bulkley material.^{17,20,21} In contrast, materials well described by the Maxwell model, including polymer melts, entangled polymer solutions, polymer networks with reversible bonds, and concentrated micelles, are dominantly fluid-like at low frequencies, and therefore, do not exhibit a shear-rate-independent shear stress at low shear rates.^{10,22} Shear-rate

sweeps on several of these materials reveal shear-thinning behavior, in which stress scales as $\dot{\epsilon}$ at low shear rates and $\dot{\epsilon}^p$ at high shear rates, where $p < 1$ (Figure S1c).

In the published literature, when new materials and methods for soft matter printing applications are reported, the yielding of a material is often characterized by measuring how the elastic shear modulus, G' , varies with applied stress or strain at a single oscillatory frequency. In these tests, G' is most always found to be a constant at low levels of stress or strain and to drop dramatically at high stresses or strains. The threshold stress at which the elastic shear modulus begins to drop is used to approximate a yield stress. Most complex fluids, including jammed granular materials, colloidal glasses, concentrated polymer solutions, polymer melts, and polymer networks with reversible bonds, exhibit this transition (Figure S1d). The preceding discussions of Maxwell fluids, Kelvin–Voigt solids, yielding, and shear thinning, demonstrate that the underlying mechanisms controlling this general “thinning” behavior can differ significantly from material to material. Thus, caution must be taken when interpreting measurements of G' versus stress or strain; the performance of a material



in 3D printing applications are not easily inferred from such measurements. A more tractable characterization is attained from measuring frequency-dependent moduli and stress as a function of shear rate.

Methods

All literary searches conducted for this review were performed in Google Scholar using combinations of the following search terms: 3D printing, 3D bioprinting, biofabrication, soft matter printing, hydrogel printing, embedded printing, direct write, sacrificial material, sacrificial support, sacrificial scaffold, sacrificial inks, biomaterials, self-healing, granular materials, yield stress materials, liquid like solids, tissue engineering, rheology. All graphical representations of data presented here were developed specifically for this review article by the authors using data gathered during literary searches; citations are included for the sources of the presented data in the figure and figure captions as appropriate.

All papers that contain the necessary data related to printed feature size, material deposition rate, and tangential velocity of the nozzle were included in Figures 2 and 3 (in the main

article). When presented in the text or in a tabular format, the material deposition rates, feature sizes, and tangential velocities were copied directly; when presented in a graphical format, these variables were measured from the given axes or scale bars. The cross-sectional area of printed structures was determined using $A = \pi ab/4$, where a is the width and b is the height of the printed structure. When only a single dimension was given, the cross-sectional area was approximated by $A = \pi d^2/4$. Those papers that did not provide sufficient information to determine one or more of the required variables were excluded from Figures 2 and 3 (in the main article).

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