



Recent advances in 3D printing with protein-based inks

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ABSTRACT

Three-dimensional (3D) printing is a transformative manufacturing strategy, allowing rapid prototyping, customization, and flexible manipulation of structure-property relationships. Proteins are particularly appealing to formulate inks for 3D printing as they serve as essential structural components of living systems, provide a support presence in and around cells and for tissue functions, and also provide the basis for many essential *ex vivo* secreted structures in nature. Protein-based inks are beneficial *in vivo* due to their mechanics, chemical and physical match to the specific tissue, and full degradability, while also to promoting implant-host integration and serving as an interface between technology and biology. Exploiting the biological, chemical, and physical features of protein-based inks can provide key opportunities to meet the needs of tissue engineering and regenerative medicine. Despite these benefits, protein-based inks impose nontrivial challenges to 3D printing, such as sufficient concentration, rheological features, and structural hierarchy observed in nature that is a source of the robust mechanics and functions of these materials. This review introduces photo-crosslinking mechanisms and rheological principles that underpins a variety of 3D printing techniques. The review also highlights recent advances in the design, development, and biomedical utility of monolithic and composite inks from a range of proteins, including collagen, silk, fibrinogen, and others. One particular focus throughout the review is to introduce unique material characteristics of proteins, including amino acid sequences, molecular assembly, and secondary conformations for designing printing inks and for controlling the printed structures. Future perspectives of 3D printing with protein-based inks are also provided to support the promising spectrum of biomedical research accessible to these materials.

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Abbreviation: 3D, three-dimensional; ECM, extracellular matrices; Ala, alanine; Gly, glycine; Ser, serine; Val, valine; Tyr, tyrosine; H-B, Herschel-Bulkley; σ_y , yield stress; LVR, linear viscoelastic region; G' , storage modulus; G'' , loss modulus; T, temperature; SLA, stereolithography; FRESH, Freeform reversible embedding of suspended hydrogels; RGD, arginine-glycine-aspartate; HRP, horseradish peroxidase; H_2O_2 , hydrogen peroxide; PCL, polycaprolactone; PLCL, polylactide-co-caprolactone; LDV, leucine-aspartic acid-valine; EDS, glutamic acid-aspartic acid-serine; PDMS, polydimethylsiloxane; hMSC, human mesenchymal stem cell.

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1. Introduction

Three-dimensional (3D) printing/additive manufacturing is transforming the landscape of conventional manufacturing and is becoming a versatile and potent tool for a myriad of biomedical and clinical applications [1,2]. 3D printing generally relies on adding materials in a layer-by-layer or voxel-by-voxel manner to generate 3D architectures. This additive strategy is in contrast to subtractive approaches such as mechanical machining, or formative ones such as molding/casting, and thus provides distinct manufacturing advantages. 3D printing enables rapid prototyping of structures customized in terms of composition and morphology, a feature desirable for precision medicine. 3D printing is also advantageous in spatially controlling the complex organization of multiple components to help recapitulate the heterogeneity of native tissues. Third, 3D printing enables tight control of structure-property relationships, providing tunable outcomes in terms of mechanical properties, as well as the degradation rate of the structures.

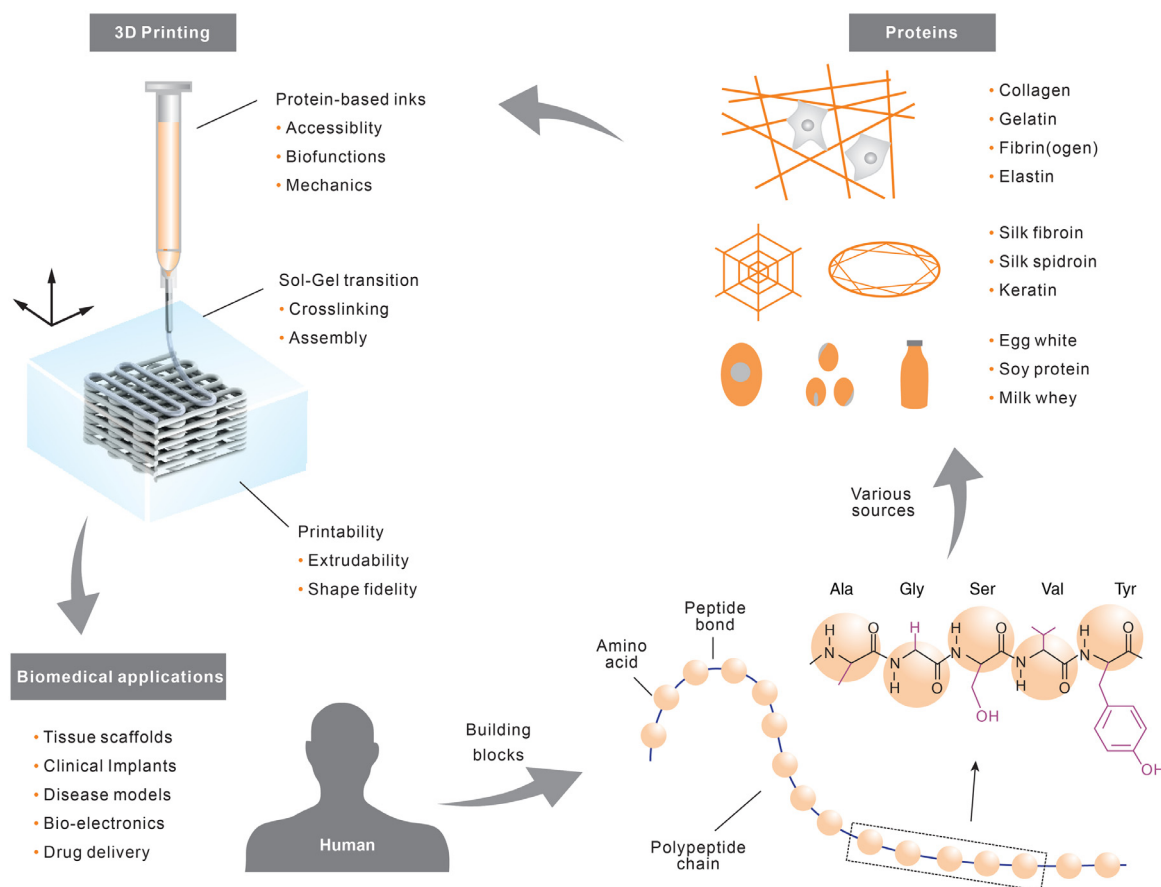


Fig. 1. Overview of protein-constituted inks for 3D printing. Proteins, as the primary building blocks of living systems including humans, are linear macromolecules/polypeptide chains consisting of amino acids held together by peptide bonds. The chemical structure of a model pentapeptide, consisting of alanine (Ala), glycine (Gly), serine (Ser), valine (Val), and tyrosine (Tyr), is illustrated. The side chain groups of each amino acid are labeled in purple. Proteins can be derived from a variety of sources as well as genetic engineering techniques. Protein-constituted inks are printed, via a sol-gel transition, into 3D structures that hold promise to address critical challenges in biomedical and clinical applications.

The key component of 3D printing is printing ink [3,4]. The performance and functionality of 3D printing are largely determined by the biochemical, rheological, and material properties of the ink. A broad spectrum of materials has been developed for 3D printing inks, including ceramics [5], metals [6,7], glasses [8,9], synthetic polymers [10,11], and nature-origin polymers [12–14], consisting of polysaccharides and proteins. Among these materials, proteins represent the major building blocks for cells, extracellular matrices (ECM), and various biological structures. The prevalence of proteins in living systems makes protein-based inks particularly alluring to construct proteinaceous scaffolds that resemble or mimic the proteinaceous composition of native tissues (Fig. 1). In comparison to other materials, the use of these protein-based compositions is expected to minimize immune reactions and promote biomaterial-host integration, as well as to degrade fully over time, generating peptides and amino acids as nutrients for the cells.

Proteins are macromolecules consisting of amino acids [15], which have 20 standard types in eukaryotes yet result in an amazingly vast diversity of protein-based materials [16,17]. Proteins can be roughly categorized as fibrous and globular proteins, depending on the capability to form fibrils. Most globular proteins form into singular entities; some globular proteins, such as lactoglobulin and insulin, can form fiber-like structures (e.g., amyloid fibrils) under certain conditions [18,19]. Some fibrous proteins are derived from the decellularized native tissues or ECM [20], including collagens, fibronectin, elastin, fibrin, and cytokeratin [21–24]. These

ECM proteins are tissue-specific and dynamically interact with cells via remodeling, degradation, and signal transduction [25], and help maintain tissue-specific functions and support the differentiation of cells [26,27]. Proteins, not from the decellularized tissues, include keratin, silk, soy, albumin, zein, and ovalbumin. The former two are fibrous while the rest are usually globular. These proteins are advantageous to be printing inks as they usually have a convenient and easy-scalable supply. For example, keratin and silk that can be harvested from the textile industrial production of wool and cocoons, respectively. Other food proteins are also available based on food infrastructure already in place.

Proteins are the structural basis of tissues with important mechanical properties, such as tendons from collagens and silks from fibroins and spidroins. For example, the specific-energy storage capacity of vertebrate tendons ($\sim 800 \text{ J kg}^{-1}$) is higher than that of high-tensile spring steel (120 J kg^{-1}) by over six-fold [28]. Also, spider dragline silk is one of the toughest materials known; its maximum toughness ($\sim 350 \text{ MJ m}^{-3}$) is higher than a range of high-performance polymers, including Kevlar 49 (50 MJ m^{-3}) and nylon (80 MJ m^{-3}) [29]. These mechanical feats are useful for regenerating native tissues and for developing new devices and systems where high-performance is desirable.

These exceptional material properties of protein-based tissues stem from a ubiquitous manufacturing strategy in biology, directed hierarchical molecular assembly. The protein molecules contain specific encoded designs, i.e., amino acid sequence information,

which guides the assembly via weak interacting forces, such as hydrogen bonding and Van der Waals [28]. The molecular assembly process is dynamically regulated by cells, via rates of biosynthesis and cellular export, enzyme-cleavage, and environmental cues, to form consistent, well-controlled structures across the hierarchical length scale under ambient and aqueous conditions. Translating these facile processing conditions to 3D printing with protein inks supports the integration of cells and bioactive molecules (e.g., cytokines and growth factors) into 3D printed proteinaceous structures, leading to hybrid living materials [30–32]. This biological manufacturing strategy is rich in information, including both molecular designs and the regulation of cell functions, in sharp contrast to the energy-intensive artificial manufacturing of polymers with organic solvents, thermal conditions or harsh chemical reactions. Such conditions usually compromise bioactivity.

Proteins have been printed into a host of 3D structures with substantial capability for tissue regeneration, disease models, and drug screening (Table 2) [33–35]. These exceptional studies mainly rely on conventional polymerization techniques, such as covalent crosslinking and organic solvents. However, the understanding of hierarchical macromolecular assembly can significantly facilitate the development of new 3D printing techniques to unleash the full potential of protein-based inks. Considerable progress has been made toward this goal, with the assembly of short peptides and polymers into functional structures [36–38]. Monolithic collagen and silk inks have been 3D printed via pH-induced assembly [39] and salt-directed assembly [40], respectively. Yet, there remains a largely untapped space of materials properties to be achieved via further understanding, and then the exploitation of proteins in 3D printed matrices for a wide range of utilities.

According to ASTM [41], the 3D printing/additive manufacturing for polymer materials encompasses a broad range of seven technical categories, including directed energy deposition, sheet lamination, binder jetting, material extrusion, material jetting, vat polymerization. Because not all these techniques apply to protein-based inks, we only focus on rheological behavior and photo-crosslinking approaches, underpinning these distinct 3D printing techniques. Viscoelastic behavior and rheological characterization of protein inks play a central role in the process of extrusion and jetting; photo-crosslinking determines printing efficiency and the structure and mechanical performance of protein hydrogels. We also cover the recent progress of computational simulation on predicting and understanding protein assembly, as it is promising for assisting 3D printing with protein inks. Then, we review several representative protein-based inks for 3D printing, including those based on collagen, silk, fibrinogen, keratin, and several food proteins. Some protein materials have been less exploited for 3D printing than others yet demonstrate unique value in biofunctions and accessibility. The inclusion of these less-investigated protein materials adds to the comprehensive nature of this review and motivates more investigations in the future. Finally, we briefly envision several promising future directions of 3D printing with protein-based inks. Throughout the review, the emphasis is placed on monolithic inks (single types of proteins), sequence-process-property relationships, and methods to improve structural integrity and to expand functions of protein prints.

2. Technical considerations for 3D printing

2.1. Rheological characterizations

Several 3D printing techniques, such as material extrusion and jetting, involves the flow of ink. For example, the protein-based ink, often loaded in a movable syringe, is extruded by external mechanical forces through a nozzle to deposit on a surface. During the extrusion, the ink is expected to flow and after deposition, to main-

tain shape [42–45]. The printing performance, such as extrudability, filament fidelity, and sol-gel transition, are largely determined by the viscoelastic properties of the ink that are characterized by a set of steady or dynamic rheological tests.

2.1.1. Extrudability

Extrudability is defined as how efficient an ink is extruded from a dispensing nozzle (Fig. 2A). The primary measure of extrudability is viscosity, which indicates the resistance of a fluid to deform under external forces, and at the molecular level, the resistance to the sliding between molecules. Specific to protein-based inks, viscosity is relevant to concentration, molecular weight, and importantly, inter- and intra-molecular interactions that are associated with several physical factors, including temperature, ion composition/concentrations, and pH. The molecular weight and concentration of proteins are proportional to molecular entanglement, which impedes molecular sliding and thus increases viscosity. For example, the viscosity of collagen and regenerated silk fibroin solutions was associated with concentration [46] and molecular weight [47], respectively. Furthermore, the side chains of the amino acids endow a variety of secondary conformations and molecular interactions of proteins via covalent bonding, such as disulfide formation, as well as via weak bonding, including salt bridges, hydrogen bonding, and van der Waals forces. These molecular conformations and interactions also influence sliding between protein molecules and thus are relevant to the viscosity. For example, the viscosity of a partially hydrolyzed product of collagen, i.e., gelatin, is dependent on temperature because the hydrogen bonds that hold the triple-helix conformation of gelatin together are weakened by increased temperature, generally above 30 °C [48]. Also, according to previous definitions for injectable hydrogels [43], inks with a viscosity higher than 100 and less than 2000 Pa·s may be called a paste; higher than 2000 Pa·s may be called a putty. Depending on the viscosity of a protein-based ink, compressed air, pistons, and screw rods may be utilized to provide sufficient extrusion pressures.

The viscosity of most protein-based inks becomes smaller as the shear rate increases, as shown in a rheological flow curve (Fig. 2B). This shear rate-dependent phenomenon is a non-Newtonian behavior, called shear-thinning, in comparison to shear-independent flow (Newtonian) and shear-thickening (another non-Newtonian flow behavior). This shear-thinning property of the ink is desired for 3D printing, as it lowers the pressure required to extrude the ink. Shear-thinning often results from the stretching or elongation of molecules under shear, which diminishes the molecular entanglements and facilitates the sliding between molecules, decreasing the viscosity (Fig. 2A). For example, a hydrogel of recombinant spider silks, eADF4(C16) is printable because of the shear-thinning behavior [49]. The complex viscosity decreased from 40 to 4 Pa·s with the increase of oscillatory frequency. Of note, the frequency is used instead of the shear rate to describe the shear-thinning behavior, because the complex viscosity as a function of frequency approximates the viscosity as a function of shear rate, according to the Cox-Merz rule [50]. Another molecular mechanism leading to the shear-thinning is host-guest interactions [51,52], where intermolecular interactions between adamantane and β -cyclodextrin are interrupted under shear and restored upon deposition. The reversibility of the intermolecular interactions also makes the 3D prints self-healable.

The viscoelastic behavior of protein-based inks, especially the shear-thinning, can be complex. Different inks can always be fitted by several models, including the Cross, Herschel-Bulkley (H-B), Power-law, and Sisko models (Fig. 2C and Table 1). The fitting parameters allow quantitative comparison between inks. The H-B model has long and widely been used to characterize the shear-thinning behavior of inks for extrusion-based 3D printing [53,54].

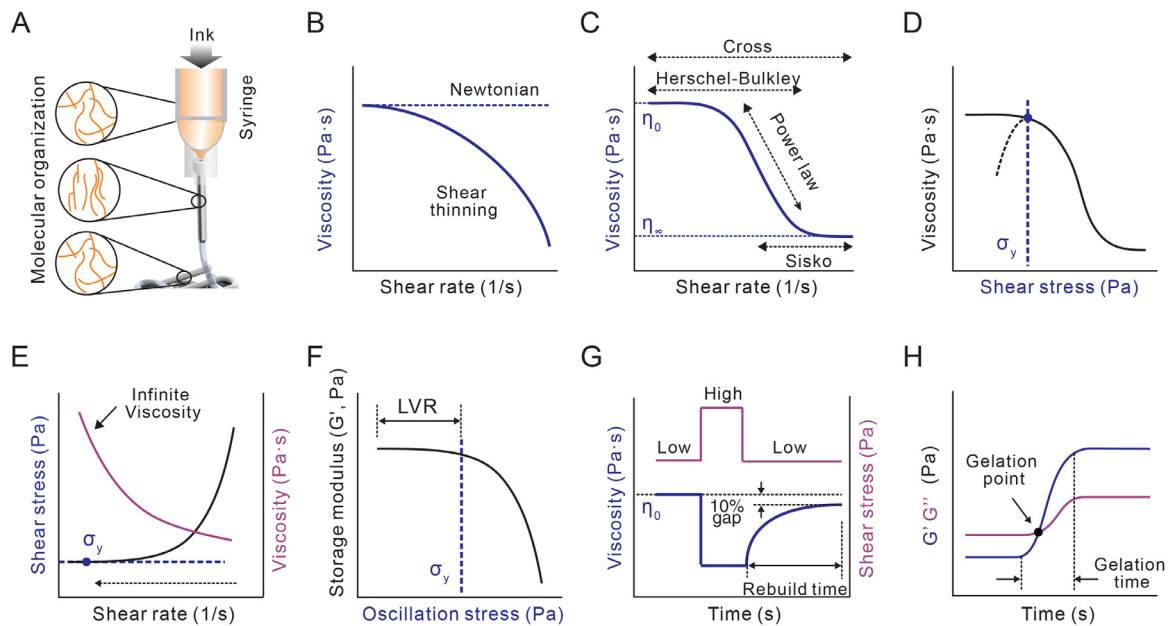


Fig. 2. Rheological characterizations of viscoelastic protein inks for 3D printability. A. Illustration of the extrusion of ink and the stretching and rebuilding of molecular organization. B. Comparison between Newtonian and non-Newtonian shear-thinning behaviors. The flow curve is plotted in viscosity as a function of shear rate. C. Illustration of four models, including the Cross, Herschel-Bulkley (H-B), Power law, and Sisko, for fitting the different viscoelastic behaviors of inks. η_0 and η_∞ indicate zero-shear and infinite-shear viscosity, respectively. D. Yield stress (σ_y) is measured by the transition point of viscosity. Dash line indicates a peak transition. E. In a steady sweep from high to low shear rate, the σ_y is measured at the low end of shear rate. F. Transition of storage modulus (G') in oscillatory sweep tests gives rise to the σ_y . D LVR indicates the linear viscoelastic region. G. Thixotropy characterization by a three-step stress sweep. The ink exhibits irreversible loss of around 10% viscosity after shearing. H. Crosslinking characterization by oscillatory time sweep. The gelation point is where G' becomes equal to loss modulus (G''). The time between two equilibrated G' values is gelation time.

Table 1
Common flow models for extrusion-based 3D printing.

Models	Equation ^a	Range
Cross	$\eta(\dot{\gamma}) = \eta_\infty + \frac{\eta_0 - \eta_\infty}{1 + K\dot{\gamma}^{n-1}}$	$\eta_0 \rightarrow \text{Shear-thinning} \rightarrow \eta_\infty$
Herschel-Bulkley	$\eta(\dot{\gamma}) = \frac{\sigma_y}{\dot{\gamma}} + K * \dot{\gamma}^{n-1}$	$\eta_0 \rightarrow \text{Shear-thinning}$
Power law	$\eta(\dot{\gamma}) = K * \dot{\gamma}^{n-1}$	Shear-thinning
Sisko	$\eta(\dot{\gamma}) = K * \dot{\gamma}^{n-1} + \eta_\infty$	Shear-thinning $\rightarrow \eta_\infty$

^a η_0 is the zero-shear viscosity (Pa·s); η_∞ is the infinite shear viscosity (Pa·s); $\dot{\gamma}$ is the shear rate (1/s); K and n are parameters related to the shape of the flow curve; σ_y is the yield stress (Pa).

In this model, the viscosity is a constant when the shear stress is smaller than the yield stress (σ_y); the viscosity starts to change when the shear stress is larger than σ_y . Yield stress implies the maximum strength of the molecular structures within the ink, and indicates the minimum shear stress to initiate the flow of ink, i.e., extrusion. Thus, for extrusion-based 3D printing, the extrusion stress needs to be higher than the yield stress of the ink. Yield stress is also important to filament fidelity and is discussed in the next section. In all models, when a factor, n approximates one, the viscosity becomes more independent of the shear rate, indicating a more Newtonian behavior, i.e., shear-independence. Besides the viscoelastic properties of the ink, extrudability is relevant to the size and geometry of the nozzle, which contributes to the hydrodynamic resistance. Nozzles with a finer gage, a longer length, and a cylinder shape, in comparison to a conical shape, lead to higher resistance and thus require a higher extrusion pressure.

2.1.2. Filament fidelity

Filament fidelity is the maintenance of the shape of the extruded filament upon deposition, which is associated with at least two viscoelastic properties, yield stress (σ_y) and thixotropy.

Insufficient yield stress may lead to collapse or sagging of printed filaments under its gravity and the weight of the above layers. The yield stress of the protein-based inks can be improved

by adding thickeners and bulking agents, e.g., gellan [55]. Depending on the magnitude of the viscosity, “apparent” yield stress of the protein-based inks can be measured from a double logarithmic plot using three methods (Fig. 2D–F) [56]. The first one is a steady stress sweep for inks that have a medium viscosity and show a well-defined transition of viscosity (Fig. 2D). Depending on the equilibrium of the shear rate, the yield stress can be obtained from either the onset point of the decrease in viscosity or the maximum peak of the viscosity. The second one is a steady sweep from high to low shear rates (Fig. 2E). The approach is usually suitable for liquids with a low viscosity that may appear as infinity for the “apparent” viscosity. The yield stress is the plateau value at the low end of the shear rate and becomes independent of the share rate. The third approach is oscillatory stress sweep (Fig. 2F). The yield stress is obtained from the onset point of the decrease in storage modulus (G'). This approach is performed mostly within the linear viscoelastic region (LVR), thus circumventing wall slippage issues and improving reproducibility. This approach is suitable for a wide range of viscosities.

Another viscoelastic property, relevant to filament fidelity, is thixotropy that describes a time-dependent process to rebuild a molecular structure after shear-induced disruption (Fig. 2A and G). Thixotropy can be experimentally characterized by a three-step shear sweep to indicate the recovered viscosity. The first step is to apply a low shear rate which corresponds to the slowly flowing ink in the syringe; then in the second step, the shear rate is dramatically increased, corresponding to the state of the ink flowing through the nozzle; in the third step, the shear rate returns to the low value, allowing the gradual building up of molecular structures and the recovery of viscosity. In Fig. 2G, the viscosity is recovered to 90% of the original value; the 10% gap may be permanent due to irreversible structural changes in the ink. For extrusion-based 3D printing, rapid rebuilding time and a large percentage of the recovered viscosity are favorable for filament fidelity. For a protein ink with collagen and gelatin nanoparticles [57], 75% storage modulus of the ink was recovered within one second. By contrast, an

Table 2
Summary of protein-based inks for 3D printed tissue scaffolds.

Proteins	Ink composition	Methods	Crosslinking	Auxiliary	Young's modulus	Tissues	References
Collagen	Monolithic (up to 24 mg/ml)	Extrusion	Physical	Supporting bath	10 kPa; 100 kPa (GLU crosslinking)	Heart	[39]
	Monolithic (up to 17.5 mg/ml)	Extrusion	Physical	Heating	~30 kPa	Meniscus	[72]
	Monolithic (up to 12 mg/ml)	Extrusion	Physical	pH	NA	cartilage	[70]
	Collagen fibrils	Extrusion	Physical	Shear-induced fiber alignment	NA	Corneal	[177]
	Composite (Gelatin; hyaluronic acid)	Extrusion; Light	Physical; Covalent	Rheology additives	NA	Liver	[57]
	Composite (gold nanorods)	Light	Physical	Near-infrared femtosecond laser	NA	Vessel	[184]
	Composite (Pluronic® F-127)	Extrusion	Physical	Rheology additives	7 kPa	Bone marrow	[71]
	Composite (Alginate)	Extrusion	Physical	Rheology additives	NA	Bone; liver	[284]
	Gelatin methacrylamide (up to 20 w/v%)	Extrusion; Light	Covalent	Photopolymerization	NA	Liver	[67]
	Gelatin methacrylamide (10 w/v%)	Extrusion	Covalent	Structural enhancement	~130 kPa	Cartilage	[285]
Gelatin	Gelatin methacryloyl (up to 20 w/v%; Gellan)	Extrusion	Physical	Rheology additives	~ 186 kPa	Cartilage	[55]
	Gelatin-methacryloyl (10 w/v%; 0.6% Collagen)	Extrusion	Covalent	Photopolymerization	NA	Cartilage	[136]
	Gelatin methacrylamide (10 w/v%)	Extrusion; Light	Covalent	Enzymatic reactions; Photopolymerization	NA	Fibroblast	[61]
	Monolithic (up to ~30 wt%)	Extrusion	Physical	Aqueous salt bath	117 MPa (wet); 2 GPa (dry)	Vessel	[40]
	Monolithic (up to ~30 wt%)	Extrusion	Physical	Methanol bath	5.6 GPa	Bone	[83]
	Monolithic (16 wt%)	Extrusion	Covalent	Enzymatic reactions	150 - 320 kPa	Meniscus	[211]
	Methacryloyl-modified (up to 30% w/v)	Light	Covalent	Digital micromirror device	15 kPa (Tensile); 40 kPa (Compressive, 20% strain)	Brain; cartilage	[237]
	Monolithic (up to 2 wt%); Composite (Gelatin)	Extrusion; Light	Covalent	Photopolymerization; Rheology additives; Sacrificial templates	~400 kPa	Cartilage	[239]
	Monolithic (Hydrogel)	Extrusion	Physical	Amino acid sequence	0.21 kPa	Fibroblast	[49]
	Composite (PEG)	Extrusion	Physical	Rheology additives	NA	Fibroblast	[225]
Silk	Composite (glycerol; Gelatin)	Extrusion	Physical	Rheology additives	1.4 kPa	Bone	[216]
	Composite (GelMA; PEGDMA)	Extrusion; Light	Physical; Covalent	Photopolymerization; Enzymatic reactions	~40 kPa	Heart	[243]
	Composite (Fibrin/Gelatin)	Extrusion	Physical; Covalent	Rheology additives	~16 kPa	Cartilage	[217]
	Composite (Gelatin)	Extrusion	Physical	Rheology additives	~70 kPa (Compressive, 20% strain)	Cartilage	[218]
	Monolithic (ca. 1 mg/ml)	Extrusion	Physical	Ionic bonding	NA	Skin	[253]
	Composite (Alginate; Methylcellulose)	Extrusion	Physical	Rheology additives	NA	Bone	[255]
	Composite (PCL; PLCL)	Extrusion	Physical	Structural enhancement	1 MPa	Urethra	[252]
	Composite (Hyaluronic acid; Polyvinyl alcohol)	Extrusion	Physical; Covalent	Rheology additives; Enzymatic reactions	NA	Nerve	[254]
	Composite (Furfuryl-gelatin)	Extrusion; Light	Covalent	Photopolymerization; Rheology additives; Enzymatic reactions	NA	Heart	[286]
	Composite (PEG; alginate)	Extrusion	Physical; Covalent	Ionic bonding; Enzymatic reactions; Photopolymerization	Ca. 0.9 - 3 MPa	Cartilage	[257]
Fibrin(ogen)	Composite (Gelatin)	Extrusion	Physical; Covalent	Heating; Enzymatic reactions	1 kPa	Vessel	[251]
	Monolithic (up to 6% w/v)	Light	Covalent	Photopolymerization	~15 kPa	Fibroblast	[113]
	Composite (Lignin; guar gum; glycerol)	Extrusion	Physical	Rheology additives	NA	NA	[268]
	Composite (glycerol; dithiothreitol)	Extrusion	Covalent	Rheology additives	1 kPa; ~4 kPa (EDC crosslinking)	NA	[272]
	Composite (polyurethane)	Extrusion	Physical; Covalent	Rheology additives	NA	Fibroblast	[270]

GLU: glutaraldehyde; EDC: 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide; PEGDMA, Poly(ethylene glycol) diacrylate.

ink of alginate and methylcellulose showed recovery of ~56% viscosity after 30 s [58].

2.1.3. Sol-gel transition

The sol-gel transition in 3D printing, from liquid to solid phases, is often associated with the crosslinking of protein molecules. The crosslinking is characterized by the ratio between storage (G') and loss (G'') moduli in an oscillatory test (Fig. 2H). G' and G'' describe the elastic (solid-like) and viscous (liquid-like) properties of the ink, respectively. When G' becomes larger than G'' , this signifies the sol-gel transition. Where G' equals G'' is usually called a gelation point in a sol-gel transition. An equilibrium value of G' and the time required to reach the equilibrium can roughly indicate the density and rates of the crosslinking. A variety of crosslinking mechanisms have been used for making protein-based inks printable and for enhancing the printability. The mechanisms include chemical crosslinking via covalent bonding and molecular assembly via weak bonding. It should be noted that multiple crosslinking mechanisms are often used to enhance printability and strength [59,60].

Chemical crosslinking of protein molecules has been achieved by using enzymes, amine-reactive chemicals, and light-based polymerization. Enzymes can crosslink amino acids side chains in the protein inks. For example, transglutaminase crosslinks lysine and glutamine [61,62]; horseradish peroxidase can crosslink tyrosine groups [63]. Microbial transglutaminase was used to partially crosslink gelatin inks with improved viscosity for optimizing printability [61]. In contrast to the enzymes that form covalent bonds, another enzyme, thrombin cleaves fibrinogen to form fibrin that can form assemblies for 3D printing [64]. As an amine-reactive chemical, glutaraldehyde [65] has been used to stabilize the prints of gelatin [66] and collagen [39] after printing. However, the toxicity of glutaraldehyde may be an issue for subsequent cell-involved applications [67]. Another chemical crosslinking mechanism for 3D printing is light-based polymerization. The detailed mechanisms of light-based polymerization will be discussed in Section 2.2.

Molecular assembly via weak bonding is an appealing trait for printing protein-based inks. The secondary conformations of proteins, such as helices and β -sheets, are both stabilized via hydrogen bonds, while also serving as physical crosslinking sites to connect molecules and to form gel networks (Fig. 3). The molecular basis of collagen to form α -helices is the sequence motif, XYG, where G is glycine at every third residue, while X and Y are most frequently the amino acids, proline and hydroxyproline [68]. Gelatin is a degraded product of collagen [69]. Below a critical temperature, gelatin molecules form a partial triple-helical conformation via hydrogen bonds, leading to a sol-gel transition; above this temperature, the hydrogen bonds are weakened, and the helices are lost, leading to a gel-sol transition. This well-defined property has made gelatin a versatile additive for formulating an array of composite protein inks. Similar to gelatin, collagen is able to assemble into gels under pH [70] and temperature [71].

It should be noted that the secondary conformation (here the helical conformation) is capable of initiating a sol-gel transition for 3D printing but often leads to mechanical weakness in comparison to that of native tissues. It is owing to insufficient control over the molecular assembly and, consequently, the lack of native structural hierarchy. For example, the compressive modulus of a meniscus print with 20 mg/ml collagen was around 30 kPa, less than half of the lower limit of the native meniscus [72]. A common way to improve the mechanical strength after molecular assembly is chemical crosslinking. The Young's modulus of a collagen print is created from around 10 kPa to around 100 kPa after glutaraldehyde treatment [39].

Besides gelatin/collagen, another example is silk proteins that form a β -sheet conformation [73–75] to crosslink molecular chains

physically for a sol-gel transition (Fig. 3) [76]. The sequence motif of silk fibroin is GAGAGS that stacks together to form β -sheets. A range of processing conditions can induce the β -sheet conformation, including organic solvents [77], salts [78], heating [79], pH [80], mechanical shearing [81], and ultrasonication [82]. However, not all sol-gel transitions satisfy the dynamics required by 3D printing. The most widely used sol-gel transition for 3D printing with silk proteins is by organic solvents, such as methanol and isopropanol [83,84]. Recently, a de novo aqueous salt bath has been developed for 3D printing with silk proteins [40]. This approach directs the assembly of silk fibroin into β -sheets, but also helps to align the molecular chains longitudinally for improved mechanical performance. This printing approach is discussed in Section 3.2.

2.2. Photo-crosslinking approaches

Light-based polymerization usually allows high resolution and speed of 3D printing. The resolution is affected by multiple factors but theoretically can be as small as the diffraction limit of the light used in the process. The scanning of the laser and the projecting of light are faster than the mechanical movement of dispensing nozzles. In addition, light-based 3D printing uses a reservoir to hold the photo-polymerizable ink during the printing, which can eliminate some technical issues in extrusion-based methods, such as nozzle clogging and harmful shear stress to cells [35,85]. However, the configuration of an ink reservoir in light-based 3D printing usually restricts the use of multiple inks, as it is challenging to rinse one ink and fix the print before adding the next ink [86–89]. By contrast, extrusion-based 3D printing is advantageous in multi-material printing, due to the convenient and rapid exchange of inks between multiple printheads [90–92]. Light-assisted 3D printing can be roughly categorized as laser scanning and light projection, depending on the light source. The first category includes two-photon polymerization [93,94] and stereolithography (SLA) [95], which employ lasers to scan protein-based inks in a voxel-by-voxel manner for (de)polymerization. The second one uses light projection via either a micro-mirror array or a dynamic mask to cure proteins in a layer-by-layer manner [96,97], including digital light processing (DLP). Despite the broad range of light-assisted 3D printing techniques, the underlying photochemistry is general. In this section, we focus on three representative mechanisms of photo-polymerization of proteins [98–101], including acrylate free-radical polymerization, thiol-ene click reaction, and photo-oxidation, which have demonstrated utility with protein-based inks. Other photo-polymerization mechanisms suitable for protein inks such as nitrogen radical reactions, were discussed elsewhere [102].

2.2.1. Free-radical

Free-radical photopolymerization of acrylate groups is widely used for 3D printing with protein-based inks, especially gelatin (meth)acryloyl (GelMA), due to the high reaction efficiency [103,104]. Acrylate groups are not present in native proteins, so chemical immobilization is needed. In the presence of photo-generated radicals, (meth)acryloyl groups are polymerized with each other in a chain-growth manner (Fig. 4A). A cross-linker is not necessary for the polymerization, but its involvement may provide finer control over the photopolymerized molecular network of proteins. One drawback of the free-radical photopolymerization process is the inhibition of soluble oxygen, which can be addressed by using a nitrogen purge gas but may interfere with the encapsulation of cells if this is a consideration.

The efficiency of the free-radical photopolymerization is also associated with photo-initiators that produce free radicals in response to light. The photoinitiators for protein-based inks should be water-soluble and biocompatible [105–107]. Three broadly used

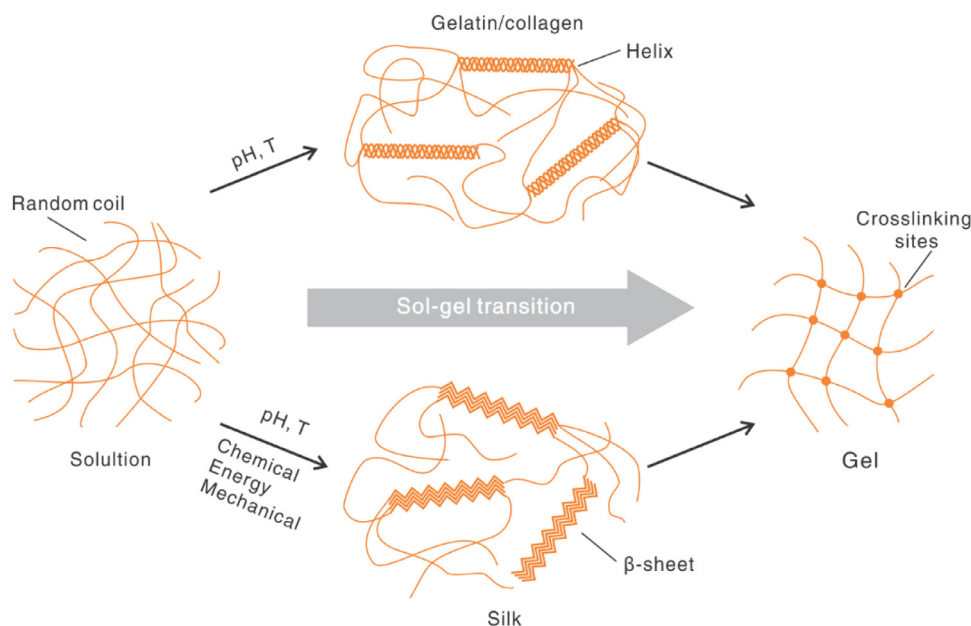


Fig. 3. Crosslinking mechanisms of proteins via secondary conformation leading to a sol-gel transition in 3D printing. For gelatin/collagen, random coil is transitioned into helices by pH and temperature (T). For silk, random coil is transitioned to β -sheets under several conditions, including chemical (salts, pH, and solvents), energy (heating and ion beam), and mechanical (vibration and sonication) inputs.

photo-initiators with UV light absorbance are 2,2-azobis (2-methyl-N-2-hydroxyethyl) propionamide (VA-086) [67], 2-hydroxy-1-[4-(2-hydroxyethoxy) phenyl]-2-methyl-1-propanone (I2959), and lithium phenyl-2,4,6-trimethylbenzoylphosphine (LAP) (Fig. 4A). LAP exhibits a 10-fold faster gelation time than I2959 [105]. The photo-initiators with visible light absorbance include Rose Bengal [108], Eosin Y [109–111], and flavin-monomonucleotide [112–115]. The use of visible light, in place of UV light, reduces potential damages to cells, especially to DNA [116]. Flavin-monomonucleotide is a water-soluble variant of riboflavin and has been used with unmodified proteins, such as collagen [70,117,118] and silk [112]. However the riboflavin-photocured protein structures are rather weak. For example, the storage modulus of 8 mg/ml collagen ink was about 20 Pa, while the pH-induced assembly was around 1700 Pa [70].

2.2.2. Thiol-ene

Another photopolymerization mechanism is the thiol-ene reaction (Fig. 4B) [119]. This reaction couples two complementary groups, a C=C double bond/ene and a mercapto/thiol group. The thiol-ene reaction is a well-known example of click chemistry, which features fast kinetics and high efficiency [119]. Because of these benefits, photoinitiator concentration and the time of light exposure can be reduced, which favors cells in terms of cytocompatibility. The thiol-ene reaction, in comparison to acrylate systems, is not limited by soluble oxygen and works in a step-growth manner, which is promising for more homogeneous polymer networks and a better filament fidelity.

The thiol-ene reaction may be beneficial for some proteins rich in cysteines with a thiol sidechain, such as soy proteins [120], because the chemical modification is not required. For other protein molecules without abandoned thiol side chains, chemical modification with “ene” groups is needed. Common ene groups include norbornene, vinyl esters, and allyl ethers [121,122]. Norbornene provides the highest reaction rates, as ring strain leads to a “spring-relief” behavior [122,123]. Less commonly, proteins were chemically modified with thiol groups and crosslinked with multidentate “ene” crosslinkers [124,125]. Deep-UV light, at a low wavelength of 254 nm, is able to activate the thiol group and carry out the thiol-ene reactions in the absence of photoinitiators (Fig. 4B)

[126]. The elimination of photoinitiators is another benefit of the thiol-ene reaction.

2.2.3. Photo-oxidation

A third mechanism is photo-oxidation, which crosslinks the side chain groups of native amino acids, such as tyrosine and tryptophan. This approach is particularly appealing to protein-based inks, as a wide range of proteins have considerable content of tyrosine, including recombinant resilin (~6.7%) [127], titin immunoglobulin domains (~6.7%) [128], fibrinogen (β chain, 4.9%) [129], gelatin (0.9%) [130], fibronectin (4%) [131], collagen (1%) [131], fibrin (~4%) [131], and silk fibroin (5.3%) [132,133]. The percentage in the brackets indicates the mole percentage of tyrosine. Besides, the tyrosine side chain can be incorporated into proteins via chemical modification for higher tyrosine content [132,134]. The higher content of tyrosine results in a higher density of crosslinking, thus leading to additional ranges of mechanical performance.

The photo-oxidation process was illustrated with an example of the widely used ruthenium (II) trisbipyridyl ($[\text{Ru}(\text{II})(\text{bpy})_3]^{2+}$)/persulfate system (Fig. 4C) [135]. $[\text{Ru}(\text{II})(\text{bpy})_3]^{2+}$ has a high molar attenuation coefficient of $14,600 \text{ M}^{-1} \text{ cm}^{-1}$ at 450 nm. Under visible light, Ru(II) and persulfate are photolyzed into Ru(III) and sulfate radicals, respectively. Ru(III) is an oxidant that can oxidize tyrosine residues to give rise to a tyrosine radical intermediate. The tyrosine intermediate forms are to couple with nearby tyrosine groups, which is further stabilized by the persulfate radical and the removal of a hydrogen atom. In comparison to a free-radical acrylate reaction based on I2959, the Ru/persulfate photo-oxidation system showed less oxygen inhibition and maintained higher cell viability even with increased concentrations and light intensity. For example, with light intensity increased from 3 to 100 mW/cm^2 , I2959 decreased cell viability from 90% to around 45%, while Ru/persulfate maintained cell viability at around 90% [136].

2.3. Computational simulation for protein assembly

Computational simulation has been widely used for designing tissue scaffolds and biomedical devices with regard to mechan-

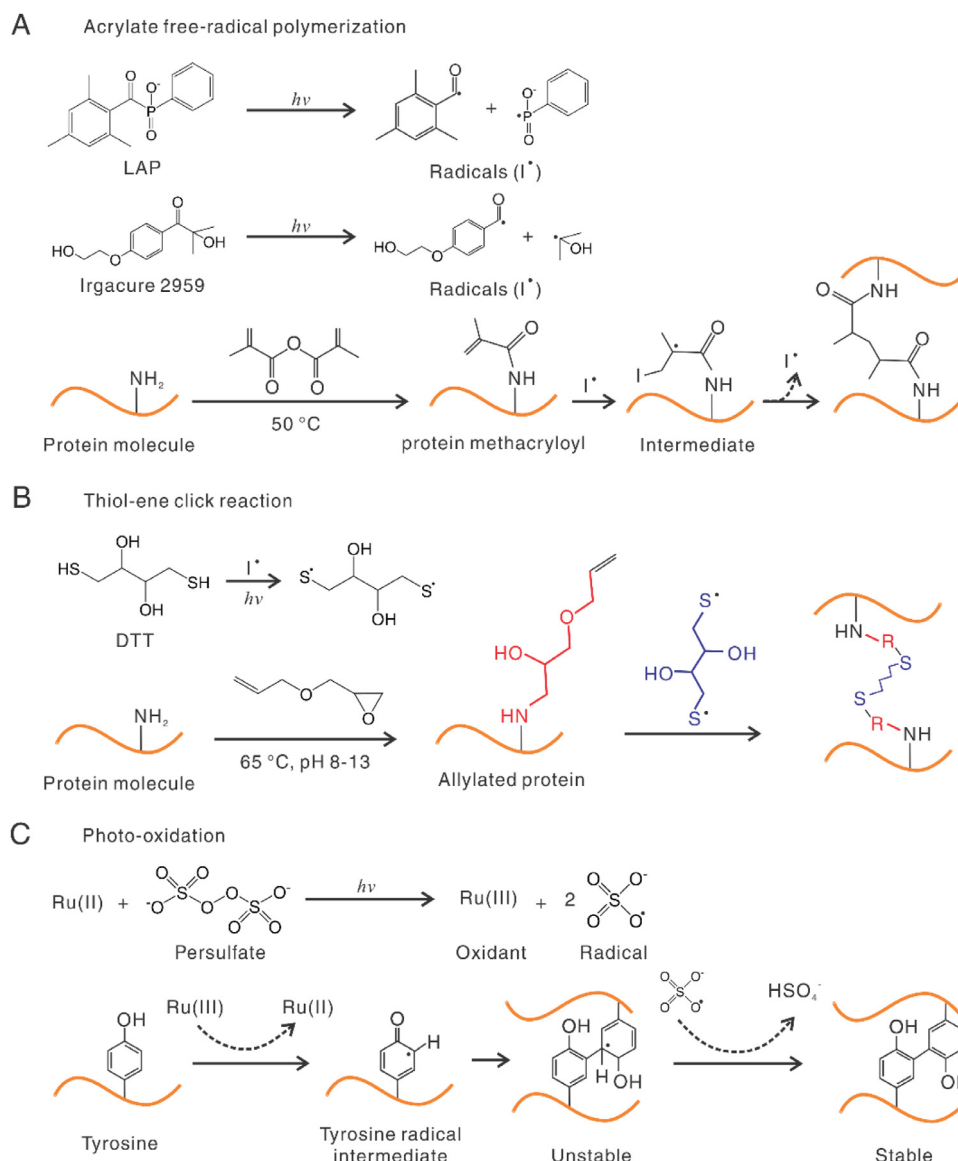


Fig. 4. Three representative photo-crosslinking mechanisms for 3D printing with proteins-based inks. A. Acrylate free-radical photo-crosslinking. Photo-initiators, such as LAP and Irgacure 2959, produce free-radicals under UV or visible light. Protein molecules are modified with acrylate groups at primary amine and hydroxyl groups. Free radicals lead to chain polymerization between methacryloyl groups from one (intra-) or different (inter-) molecules. Chain propagation is not shown for clarity. B. 1,4-dithiothreitol (DTT) is activated by UV or photoinitiators and reacts with C=C bonds, i.e., “ene” groups, for crosslinking. R in red color represents the immobilized group in the previous step. C. ([Ru(II)(bpy)₃]²⁺)/persulfate-based photo-oxidation of tyrosine residues via the formation of dityrosine bonds. The unstable intermediate is fortified by removing a hydrogen atom.

ics [137–139], mass transport [140–142], and shear stress [143]. The simulations can predict experimental results and thus reduce costly and lengthy experimental iterations [144,145]. Computational simulation increasingly plays important roles in 3D printing to predict flow behavior within nozzles [40,146,147] and the morphology of the deposited inks [148]. Specific to optimizing protein-based inks, computational simulation is also a promising tool to elucidate the molecular basis for assembly, mechanical performance, and degradation [149].

The assembly of large proteins, instead of short peptides, is often challenging for traditional molecular dynamics (MD) simulations, owing to the long folding times on the scale of seconds. To address this issue, replica exchange MD (REMD) simulation, combined with experimental characterization, has been developed to simulate large proteins with 698 residues, tropoelastin (~60 kDa) (Fig. 5A) [150]. This approach demonstrated intramolecular assembly in water from a fully extended chain of tropoelastin into elastic

fibers. Three mutations (R515A, D72A, and G685D) were simulated to change molecular conformation as well as assembly propensity, providing important insights into sequence-structure relationships. Furthermore, coarse-grained MD (CGMD) suggested that the hierarchical assembly of tropoelastin was driven by specific hydrophobic domains [151]. The interaction between hydrophobic domains reorients surrounding water molecules, and in turn, gives rise to multi-molecule coacervation. While the simulated results were consistent with experimental observations, several co-factors in the assembly process, including integrins, glycosaminoglycans, and environmental cues, such as pH and salts, can be included in the future to further mimic in vivo conditions [151].

Dimers of amino-terminal domains play an important role in the assembly of spider silks. Previous experimental results suggested a high concentration of sodium chloride (up to 300 mM) stabilizes the monomer of the N-terminus of the silk during storage in the glands of the animals prior to spinning the web, while

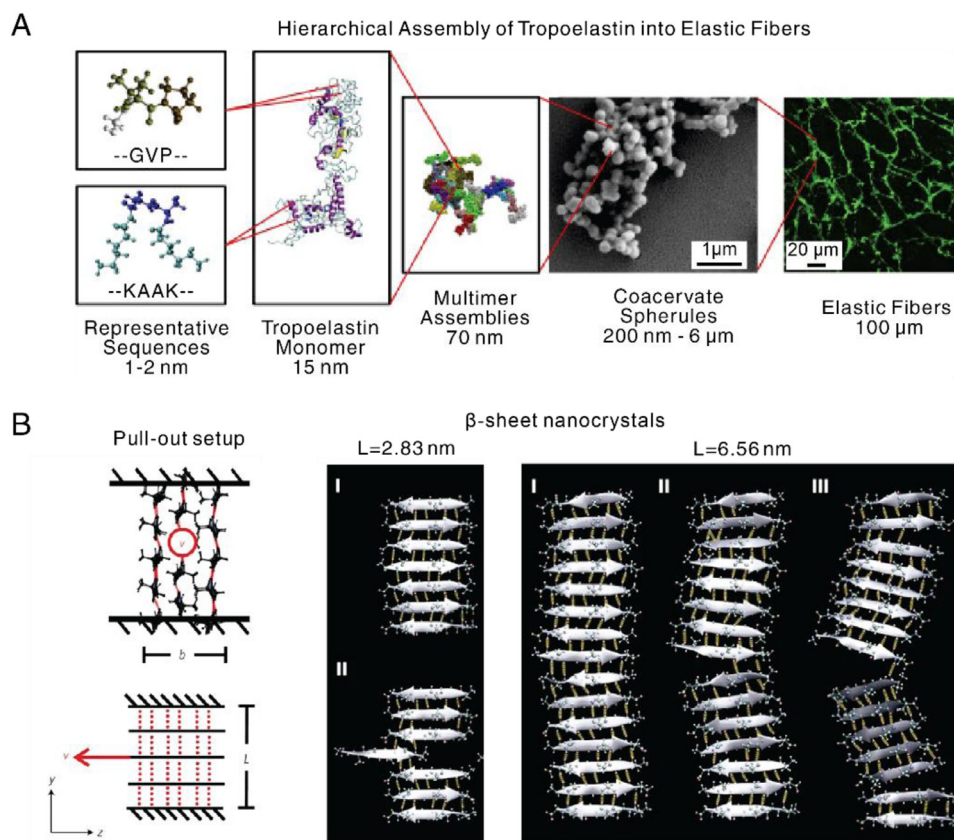


Fig. 5. Computational simulations for proteins' structures. A. Tropoelastin, as a basic unit of elastic fibers, consists of alternating GVP-rich hydrophobic domains and hydrophilic K (lysine)-containing cross-linking domains. Tropoelastin interacts with each other to form multimer assemblies, spherules, and eventually microfibers. [151], Copyright 2019. Reproduced with permission from the Elsevier. B. Pull-out configuration to characterize fracture resistance of β -sheet nanocrystals. A central strand of the middle sheet is pulled out with constant velocity (indicated by the red arrow), and the top and bottom strands are restrained. Red dash line and black line indicate hydrogen bonding and amino acid chain, respectively. Nanocrystal with short size ($L=2.83$ nm) shows a stick-slip motion, due to uniform shear loading, preserving high strength; long size ($L=6.56$ nm) leads to a crack-like flaw and significant bending, thus decreasing the strength. [164], Copyright 2010. Reprinted with permission from the Springer Nature.

the absence of sodium chloride promotes the formation of dimers for molecular assembly and fibrillogenesis during web-spinning [152,153]. This finding is consistent with the observation that along the spinning gland, from silk storage to spinning, the concentration of sodium chloride decreases [154]. MD simulation has been used to support further the influence of salt on the N-terminus and these sol-gel transitions of silk proteins. A high concentration of sodium chloride, up to 0.5 M, weakened salt bridges between the N-terminal domains by 28.6%, resulting in stabilized monomers [155].

The amino acid sequences contain essential "information" that directs protein assembly. These sequence-assembly relationships have been investigated by CGMD [156], artificial intelligence (AI) [157,158], and de novo folding calculations [159,160]. CGMD predicted the size of aggregations of two recombinant silk peptides, H(AB)₁₂ and H(AB)₂, with different numbers of domains or blocks, i.e., polymer chain length [156]. The simulations showed that longer chain length led to larger micelle-like structures, which was experimentally confirmed by atomic force microscopy (AFM) and dynamic light scattering (DLS). H(AB)₁₂ with a molecular weight of 43.7 kDa showed an average diameter of 25 nm; H(AB)₂ with 11.6 kDa showed a diameter below 2 nm. Furthermore, MD simulations identified the roles of specific amino acids in the response of proteins to environmental stimuli, such as temperature [161]. In recombinant silk-elastin-like proteins (SELPs), the X in the sequence of (GVGVP)(GXGVP)(GVGVP) has been substituted with valine (V) with a hydrophobic methyl side chain and lysine (K) with a positively charged amino group. The former increases the temperature for the thermal transition with folding via hydrophobic in-

teractions; the latter suppressed this structural transition and the response.

Besides simulating the assembly of proteins, computational simulation has elucidated the molecular basis of the mechanical strength of protein-based materials at the macroscale [162,163]. All-atom MD simulations indicated that the strength of silk materials mainly results from the appropriate size of β -sheet nanocrystals (2–3 nm) (Fig. 5B) [164,165]. β -sheet nanocrystals, longer than 3 nm, tend to bend, thus decreasing pulling strength. Mesoscale dissipative particle dynamics (DPD) simulations identified that intermediate hydrophobic/hydrophilic block ratios determined the optimal combination of protein solubility, size of assembled structures, and polymer network topology, which in turn leads to high-performance silk-based materials [166]. Simulations also showed the scale-up of nanoscale properties of silk repeat units to generate macroscopic silk fibers with outstanding mechanical properties, despite the presence of cavities, tears, and cracks [167].

3. Protein-based inks and applications

3.1. Collagens

Collagens are a family of around 29 variants. The most well-known fibrillar collagen is type I, which accounts for ~90% of all collagens in the human body and constitutes the main structural component in most connective tissues, skin, and bone of mammals [168,169]. Type I collagen is composed of three left-handed peptide chains that wind-up together to form a right-handed superhelix. The biological significance of type I collagen prompts the use of

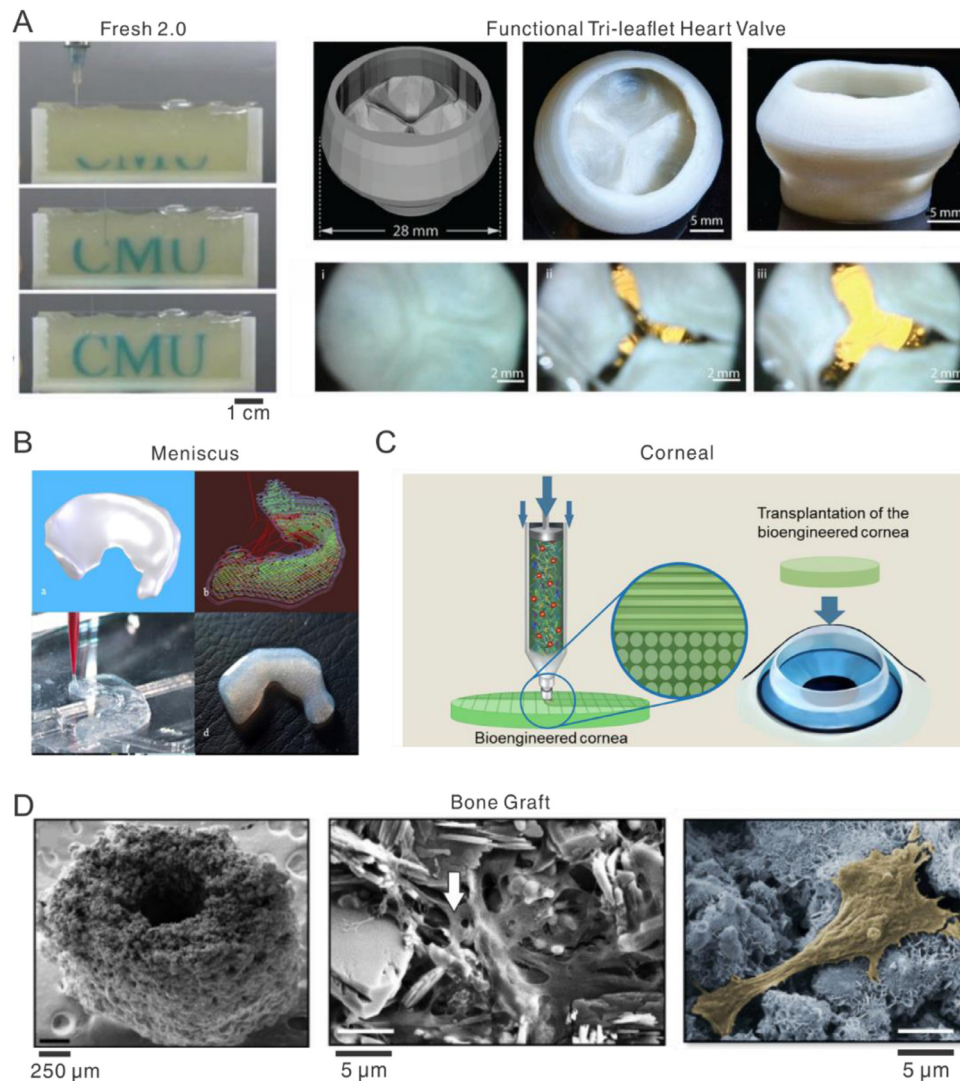


Fig. 6. Representative 3D prints with collagen-based inks. A. Left, time-lapse images of the 3D printing of the letters “CMU” dyed in cyan in a creamy suspension of gelatin microparticles with a diameter of 25 μm . FRESH means freeform reversible embedding of suspended hydrogels. Right, a 3D printed functional tri-leaflet heart valve, mechanically enhanced by glutaraldehyde crosslinking. The opening of the valve was observed under a flow. [39], Copyright 2019. Reprinted with permission from the American Association for the Advancement of Science. B. Sheep meniscus analogy printed by using 17.5 mg/ml collagen and bovine meniscal fibrochondrocytes. Top left, CT scan of native meniscus; top right, print path; bottom, 3D printed meniscus during and after printing. [72], Copyright 2016. Reprinted with permission from the American Chemical Soc. C. Illustration of 3D printing of transparent cornea with aligned collagen fibrils. The alignment results from extrusion-caused shear stress during printing. [177], Copyright 2019. Reprinted with permission from IOP publishing. D. Left, 3D printed bone graft represents the geometry of the murine femoral mid-diaphysis. Middle, collagen, indicated by a white arrow, dispersed among bound calcium phosphate crystals in the 3D printed structure. Right, a representative cell spreading on the 3D printed structure is healthy. [179], Copyright 2014. Reproduced with permission from the Elsevier.

this protein as inks for 3D printing [170–172]. Type I collagen is commonly harvested from rat tail, bovine skin, and tendons [173].

Monolithic collagen inks have been 3D printed for constructing a range of tissue equivalents, including heart, meniscus, and cornea (Fig. 6). One outstanding printing approach is the freeform reversible embedding of suspended hydrogels, or FRESH (Fig. 6A) [39,174]. A suspension of gelatin microparticles was employed to support and maintain the shape of the extruded collagen filament until the gelation of collagen was complete. In FRESH 2.0, the size and distribution of gelatin microparticles were finely controlled, resulting in an improvement of printing resolution from $\sim 200\mu\text{m}$ to $\sim 20\mu\text{m}$. Optimized pH (7.4), a high concentration (up to 24 mg/ml), and glutaraldehyde fixation [175] were also adopted to improve printing performance. For example, a 3D printed tri-leaflet heart valve showed Young’s modulus of around 110 kPa. The mechanical strength enabled cyclic opening and closing of the valve leaflets and a maximum transvalvular pressure of around

40 mmHg, above normal physiologic pressures for the tricuspid ($\sim 25\text{ mmHg}$) and pulmonary valves ($\sim 10\text{ mmHg}$) [39].

Collagen inks were also used to print a 3D meniscus by adopting a sharp transition of temperature that expedite the gelation rate (Fig. 6B) [72]. The temperature transition is achieved using an infrared heat lamp to keep the printing surface at 37°C and an ice pack to cool the printing nozzle and syringe. The sharp change of temperature expedites the gelation of collagen and avoids the collapse and spread of filaments. The heating of the printing surface, however, led to premature gelation and syneresis of 20 mg/ml collagen inks, thus limiting the highest concentration available for printing. Nevertheless, 17.5 mg/ml collagen inks led to a maximum modulus of 25 kPa, almost half of the native meniscus. A follow-up study explained the relationship between rheological behavior and the density of chondrocytes in the collagen inks [176]. The chondrocytes with a high cell density up to 100 million/ml remained viable through 14-day cultures.

Besides water-soluble collagen molecules, a suspension of monolithic collagen fibrils was used for 3D printing of transparent cornea stroma (Fig. 6C) [177]. The collagen fibrils were obtained from decellularized corneal tissues (Co-dECM) and were around 50 nm in diameter and 2000 nm in length. The shear-induced alignment was adopted primarily by manipulating the gage of the printing nozzle to recapitulate the spatial organization of collagen fibrils in the cornea. The alignment of collagen fibrils increased with increased shear stress, resulting from the decrease of the nozzle diameter. However, a 25 G nozzle, in comparison to a finer 30 G nozzle, maintained the phenotype of keratocytes better, with fewer formation of myofibroblasts. Myofibroblasts are problematic due to the loss of transparency [178]. The 3D printed cornea analogy exhibited densely packed organization of collagen fibrils and visual transmittance comparable to the human cornea and better than Co-dECM hydrogels.

Composite collagen inks have also been developed for enhancing structural integrity and for adding extra biofunctions. The common additives include calcium phosphate [179], gelatin [180–182], GelMA [67,183], gold nanorods [184], and alginate [59,185–187]. In addition, the inherent bioactivity of collagen, largely resulting from the collagen-specific amino acid sequence, was harnessed for bone regeneration [179]. Collagen at a low concentration of 1–2 wt% was blended with other components, including 0.25 wt% Tween 80 and calcium phosphate, to print bone graft substitutes with improved flexural strength and cell viability (Fig. 6D).

3.2. Silk proteins

Silk proteins are appealing for 3D printing inks [84,188–190], because of the strong analogy between native silk spinning and extrusion-based printing processes, as well as other material properties. Silk is spun from a gland via a duct, which is similar to the extrusion of printing inks from a syringe through a nozzle [188]. Furthermore, the spinning speed of silks is around 20 mm/s [191], comparable to that of extrusion-based 3D printing and much faster than the weeks- and months-long growth of other natural protein structures, such as wool and tendons. This analogy implies a natural source of inspiration for developing extrusion-based 3D printing techniques with ambient processing conditions and exceptional material performance.

The translation of silk-spinning into 3D printing requires the understanding of the underlying molecular mechanisms [29,192,193], especially those governing molecular assembly and phase-transitions. Silks are spun by a variety of animals, including insects and spiders. However, all silk proteins, despite species-specific peculiarities, share similar design principles at the molecular level. Silk protein consists of a long central segment, flanked by terminal domains; the central core includes short, repeat hydrophobic regions interlaced with hydrophilic domains [194]. For example, the representative repeated hydrophobic regions of spiders and silkworms are often AAAAAA and GAGAGS, respectively. Here, A is alanine, G is glycine, and S is serine. The hydrophobic regions of silk proteins are folded into crystalline β -sheets that underpin the high stiffness and strength of silk fibers [164]. The terminal domains regulate the formation of micelle-like structures, which enable the storage of silk proteins (>25 wt% in aqueous solutions) in the gland prior to spinning [195,196]. Furthermore, the spinning conditions, such as salts, pH, and water content, have critical roles in mediating the hierarchical molecular assembly that underlies the phase transition from water-soluble silk protein to insoluble fibers [197,198]. All of these biochemical, mechanical, and processing properties of silk proteins are valuable sources of inspiration for the development of 3D printing techniques with silk proteins or silk-inspired proteins and polymers [40,54].

Silk-based inks are often regenerated from silkworm cocoons [199,200] or produced by recombinant DNA techniques with the spider or silkworm motifs [201,202]. In particular, sericulture provides a mass supply of cocoons for textile industries, thus beneficial for scaling up silk-based inks from silk fibroin, the core protein of domestic *B. mori* silkworm cocoons. Recombinant DNA approaches provide flexibility to manipulate amino acid sequences based on silks to bring new functions. The native silk obtained directly from the spinning gland may also be used as the ink for light-based patterning [203]. It should be noted that regenerated or recombinant silk proteins may possess different properties from that of native silks due to changes in molecular weight, concentration, amino acid sequence, and molecular conformation. These differences need to be considered in designing and developing 3D printing processes with regenerated and recombinant silk proteins.

Monolithic silk protein solutions have been generated for extrusion-based 3D printing. A concentrated solution of regenerated silk fibroin (28–30 wt%) was printed into a methanol bath (85%) for constructing bone scaffolds [83,204]. However, the methanol bath aggregates silk molecules in a random manner, leading to the brittleness of 3D printed silk structures. Monolithic silk ink was also 3D printed into a bath consisting of synthetic nanoclay (Laponite) and polyethylene glycol (PEG) [205]. The bath mechanically supports the extruded silk ink and physically crosslinks silk molecules via electrostatic interactions and osmotic pressure-induced water removal [206]. The resulting mechanical properties, however, were weak. Hydrogels of recombinant spider silk protein, eADF4(C16) were used for 3D printing cell-laden lattice structures (Fig. 7A) [49,207]. The hydrogels were formed by thermal gelation (at 37 °C) of the eADF4(C16) via physical crosslinking. The cells encapsulated in the hydrogel remained viable for at least one week. RGD sequences can be added to the eADF4(C16) to improve cell adhesion, an advantage in recombinant silk proteins, in comparison to chemical coupling [208]. Adding RGD in the sequence of eADF4(C16) also increased the elastic modulus from 20 Pa to 200 Pa.

A *de novo* aqueous salt bath for 3D printing with monolithic silk inks was developed to improve the mechanical strength of the structures while also avoiding the use of organic solvents in the process [40]. The bath consisted of 0.5 M dipotassium phosphate and 4 M sodium chloride to recapitulate solvent conditions found in the native spinning glands of both spiders and silkworms, such as ion composition and pH. Besides, the high osmolarity (around 9.5 M) of the bath removes water from the silk ink to mimic the decreased water content along the native spinning gland where the silk is pre-assembled prior to spinning into the air. By using the biomimetic salt bath, silk fibroin is directed to physically assemble at hierarchical length scales [40], as shown by the longitudinal alignment of nanofibrils. The hierarchical assembly also improves the printability of the monolithic silk ink, so fine resolution (about 100 μ m), complex geometry, and superior mechanical performance were achieved. Specifically, a Y-shape microfluidic channel/vascular graft with a burst pressure of about 300 kPa was printed in the absence of sacrificial and supporting materials (Fig. 7B). Also, an overhanging filament of monolithic silk fibroin was printed with a high aspect-ratio of ~ 375 , in comparison to previous results of ~ 20 (polyelectrolyte ink) [53], ~ 33 (Carbomer-laden hydrogel) [209], and ~ 1 (silk fibroin) [63,210]. The overhanging filament is only supported by its two ends (or by one end during printing), thus representing a measure of the printability. In addition, the mechanical performance of silk prints was dramatically improved. The tensile strength, toughness, and Young's modulus of the 3D printed silk filaments in the wet state were around 39 MPa, 37 MJ m⁻³, and 117.37 MPa, respectively, superior to other 3D printed protein materials by orders of magnitude, and compa-

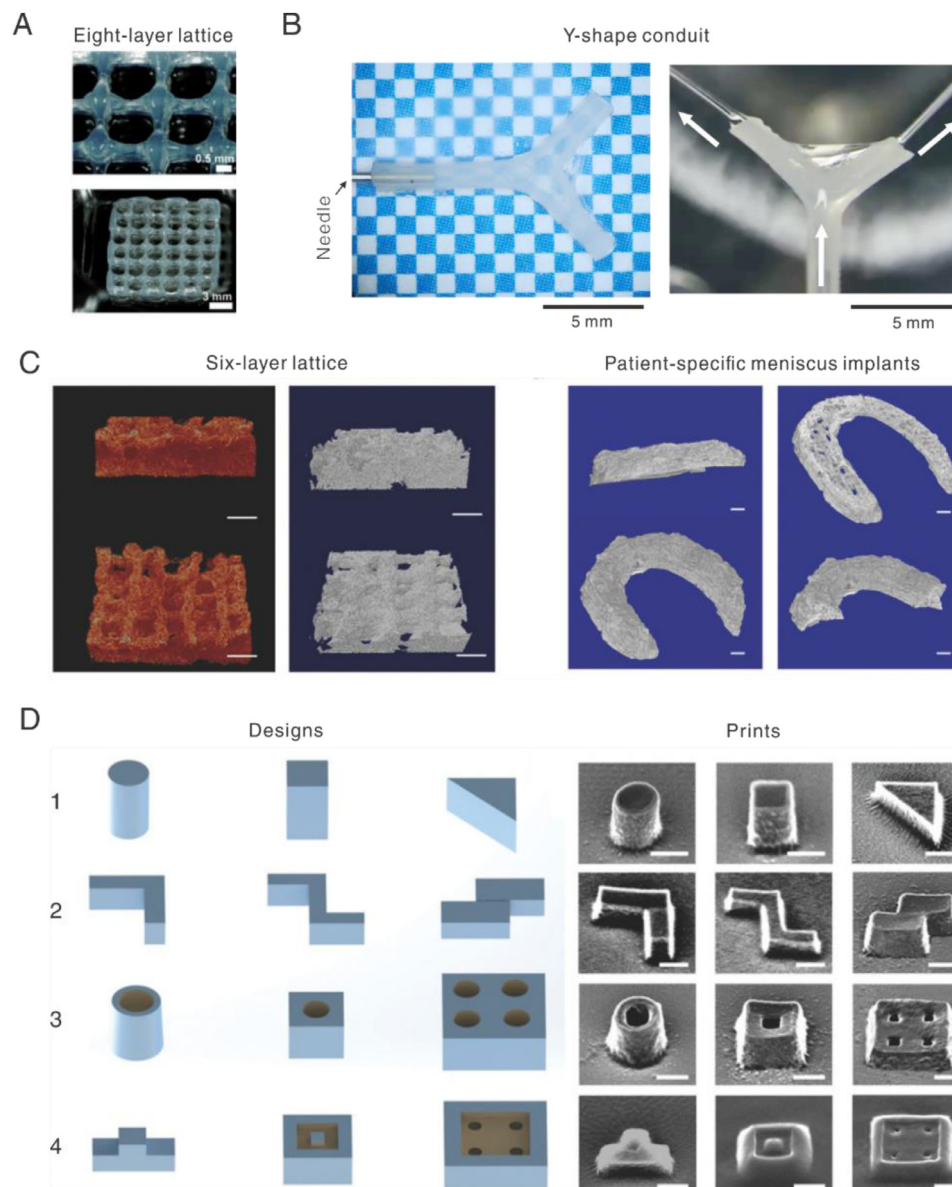


Fig. 7. Typical 3D prints with silk-based inks. A. Eight-layer lattice scaffold printed with recombinant spider silk. [49], Copyright 2015. Reprinted with permission from the John Wiley and Sons. B. Y-shape conduit/vascular graft printed with regenerated silk fibroin without using sacrificial or supporting materials. Water jets from the silk conduit, indicated by white arrows, demonstrate the remarkable strength and perfusability. [40], Copyright 2020. Reprinted with permission from the John Wiley and Sons. C. Micro-CT images of enzyme-crosslinked silk prints after freeze-drying. Scale bars: left, 1 mm; right 2.5 mm. [211], Copyright 2017. Reprinted with permission from the John Wiley and Sons. D. Left, a set of designs; right, corresponding SEM images of silk structures by ion and electron beam lithography. The scale bar is 500 nm. [240], Copyright 2018. Reprinted with permission from the John Wiley and Sons.

nable to synthetic polymers [40]. Of note, the mechanical performance was achieved by molecular assembly and in the absence of crosslinking chemicals.

Silk inks can be 3D printed using enzymatic reactions for a human meniscus scaffold [211] (Fig. 7C) and for microalgal-laden hydrogel structures [63]. Horseradish peroxidase (HRP) and hydrogen peroxide (H_2O_2) can crosslink silk via the formation of dityrosine bonds [212]. The enzymatically crosslinked silk meniscus scaffolds were post-processed via freeze-drying to be porous. Micro-CT images showed a porosity of about 59.2% and a pore size of about 224.4 μm (Fig. 7C). The meniscus silk scaffold also exhibited shape recovery after a compression strain of 80% and a storage modulus close to the native meniscus. These mechanical and morphological properties are desired for meniscus regeneration [213]. The cytocompatibility of enzymatic crosslinked silk prints was confirmed by the viability of human adipose-derived stem cells [211] and oxy-

gen production by microalgae encapsulated in the 3D printed silk structures [63].

Besides monolithic silk inks, a range of functional additives has been used to formulate composite silk inks for extrusion-based 3D printing. A major portion of additives were thickeners or bulking agents to improve printability and structural integrity, which include agar [214], gelatin [214–221], hydroxypropyl methylcellulose [63,222,223], PEG [224,225], glycerol [216], and Konjac gum [226]. Another category of additives was used to add specific biofunctions to silk inks, including tricalcium phosphate [219], extracellular matrix components [227,228], hydroxyapatite [204,229], and bioactive glass [230]. Some components have been used to formulate self-curing silk inks [231,232], including hydroxypropyl methylcellulose [233], Eri silk/gelatin [218], and polyol/gelatin [214]. Silk was also used to enhance the mechanics of other printing inks, such as chitosan [234,235] and soy protein isolate [236].

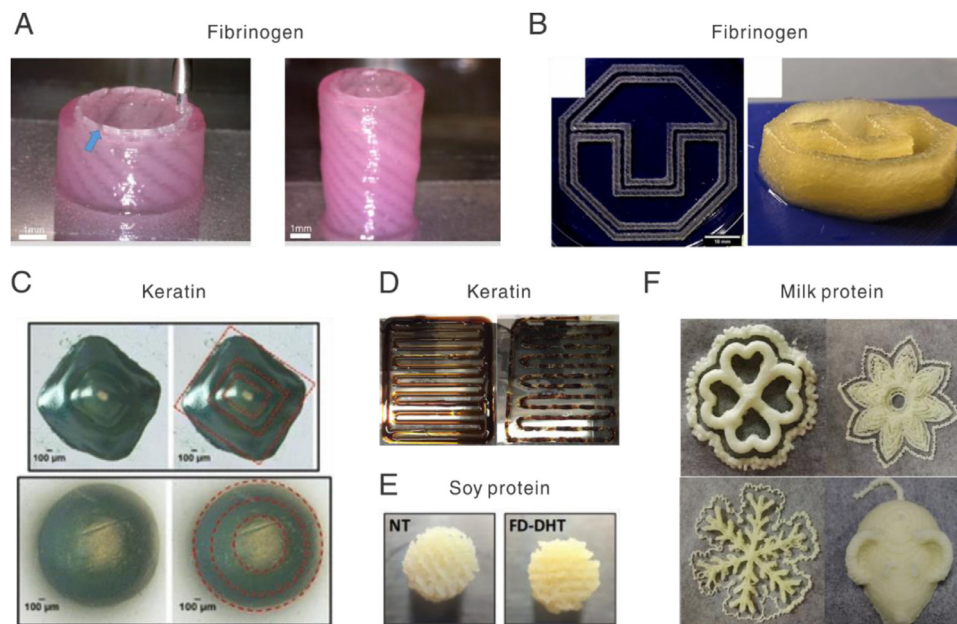


Fig. 8. Representative 3D prints of fibrinogen, keratin, and food protein-based inks. A. Urethra scaffold printed with fibrinogen-based inks. Blue arrow indicates the PCL/PLCL blend. [252], Copyright 2017. Reproduced with permission from the Elsevier. B. 3D logo printed with a composite ink of fibrinogen, alginate, and methylcellulose. [255], Copyright 2020. Reproduced with permission from the American Chemical Soc. C. Pyramid structures were printed with ketratin-based inks by light-based 3D printing. The red dash lines outline the boundary of each layer in cube or cylinder shapes. [113], Copyright 2017. Reprinted with permission from the Springer Nature. D. Printing results of a keratin-lignin (4:1) composite ink. The left and right traces were printed at 60 and 95 min after ink preparation. [268], Copyright 2020. Reprinted with permission from the Elsevier. E. Porous scaffolds printed with soy protein. NT means non-treatment with a diameter of 6.5 ± 0.19 mm; FD-DHT means freeze-dried and dehydrothermal treatment with a diameter of 6.57 ± 0.14 mm. [273], Copyright 2013. Reproduced with permission from the Elsevier. F. Various patterns and structures printed with milk proteins doped with whey protein isolate and other additives. [274], Copyright 2018. Reprinted with permission from the Elsevier.

Light-based 3D printing with silk-based inks is mostly based on the photo-polymerization mechanism [95,210,237–243], as discussed in Section 2.2. The polymorphic nature of silk proteins has also been exploited [244] as a new mechanism to form pseudo-3D and 3D structures via electron-beam and laser irradiation [245,246]. In particular, irradiation can change the solubility of silks. A set of typical silk structures via the combination of ion and electron beam lithography is shown in Fig. 7D [240]. The first row shows simple geometries; the second row shows combinations of simple geometries; the third row shows both nanosintered (embossing) and nanosculptured (debossing) geometries; and the last row shows geometries with two levels of nanosintering and nanosculpturing, respectively.

3.3. Fibrinogen/Fibrin

Fibrinogen and fibrin are essential for hemostasis, play important roles in many pathophysiological processes, and are promising biomaterials in tissue engineering [247–249]. Fibrinogen has an autologous source that is a unique advantage, in comparison with other protein materials [250]. Fibrinogen is produced in the liver and circulates through blood vessels, and thus can be derived from patients' plasma albeit in limited quantities.

Fibrinogen consists of three pairs of polypeptide chains (α , β , and γ) with a total molecular weight of about 340 kDa. Fibrinogen is cleaved by thrombin, a highly specific serine protease, to remove fibrinopeptides (A and B) to form fibrin monomer (α , β , and γ). The fibrin monomer has two complementary domains at the central region and chain ends, which are called “holes” and “knobs”, respectively. Via interactions between the holes and knobs, fibrin monomers form fibrin oligomers and protofibrils that further assemble, laterally and longitudinally, to form a network of branched and interwoven fibrin fibers, i.e., fibrin gels or clots. The chemical and mechanical stability of the fibrin gel are enhanced

by calcium-mediated ionic binding, and a plasma transglutaminase, Factor XIIIa, via covalent crosslinking between glutamine and lysine [248].

The understanding of the molecular mechanisms of fibrinogen assembly/ polymerization underpins the 3D printing with fibrinogen-based inks. For example, thrombin was used as a printing bath to trigger the phase-transition of fibrinogen solution after extrusion [64,251,252]. Factor XIIIa and calcium ions were also frequently used to enhance printability [253,254]. However, it remains challenging to use fibrin(ogen) alone for printing, owing to inadequate structural stability, and poorly defined geometry. Thus, structural enhancements of fibrinogen-based inks, by adding thickeners and multiple crosslinking mechanisms, are often necessary.

Hyaluronic acid (HA) and polyvinyl alcohol (PVA) were added to fibrinogen inks to increase printability [254]. Moreover, extrusion-induced shear stress aligned fibrin fibers, which directs the growth of neurites and Schwann cells. Other thickeners for fibrinogen-based inks include 3 w/v% alginate, 9 w/v% methylcellulose [255], and gelatin [251,256] for bone and vascular bioengineering. The printed fibrinogen-gelatin vascular graft, after two-month culture with primary neonatal human dermal fibroblasts, demonstrated a burst pressure of about 1100 mmHg, roughly 52% of that of the human saphenous vein. A PEG-alginate supporting bath was used for the printed fibrinogen inks to make cartilage-like tissues with controlled mechanical properties [257]. The fibrinogen prints supports the growth of human mesenchymal stem cell (hMSC) spheroids. Also, a blend of polycaprolactone (PCL) and polylactide-co-caprolactone (PLCL) was used, as an internal structural framework, to strengthen the fibrin gel to approach the mechanical properties of the rabbit urethra (Fig. 8A) [252]. Another interesting work blended fibrinogen with multiple additives, including alginate, genipin, and chitosan, and adopted multiple crosslinking mechanisms (Fig. 8B) [60]. This approach has been exploited to print a glioblastoma tumor model with U87MG cell line for drug screening [258].

3.4. Keratin

Keratin, commonly derived from wool, hair, and feathers, has been used for tissue scaffolds and drug delivery [259–261]. One benefit of keratin is the presence of cell-binding amino acid motifs, including leucine-aspartic acid-valine (LDV), glutamic acid-aspartic acid-serine (EDS), and arginine-glycine-aspartic acid (RGD) [262,263], which promote cell adhesion and proliferation. Keratin has two pentapeptide repeats, CCXPX and CCXS(T)S(T), where C is cysteine, P is proline, S is serine, and T is threonine [264]. The sequence motif enables keratin to form α -helix and coiled-coil architectures that transform to β -sheets in either an irreversible or reversible way. The reversible transformation between coiled-coil and β -sheets for 3D printing shape-memory Keratin structures was recently reported [265].

Keratin was oxidized to reduce the formation of disulfate bonds for preparing inks for light-based 3D printing (Fig. 8C) [113,266]. 3D printed keratin scaffolds have been used to load a drug, halofuginone, to inhibit tissue contraction for the treatment of dermal burn wounds [267]. The post-processing of the keratin prints, such as lyophilization and sterilization, was investigated toward scalable production and shelf-storage. Besides formulating photocurable inks, keratin was blended with lignin and other additives, such as Guar gum, for extrusion-based 3D printing (Fig. 8D) [268]. The blending of keratin and lignin forms copolymers via a range of molecular interactions, for example, between keratin amide and lignin aryl hydroxyls, evidenced by solvent solubility and thermal stability.

3.5. Food proteins

Soy, egg white, and milk proteins are common food ingredients and have been formulated into composite inks for 3D printing [269]. Soy protein-based ink has been developed by blending with thickeners, such as polyurethane [270], gelatin [271], and glycerol [272]. A reagent to inhibit disulfide bonds, dithiothreitol, was added into the soy ink to improve extrudability and printability [272]. The 3D printed soy scaffolds, when used as *in vivo* implants, showed immune-compatibility and macrophage-mediated degradability (Fig. 8E) [273]. For making printing inks, milk protein concentrate and egg white protein were supplemented with a range of additives, including gum and glycerol [274], gelatin [275,276], cornstarch [275,276], sucrose [275,276], egg yolk [277], and rice flour [277]. Several representative 3D prints of milk proteins and whey protein isolate (5:2) are shown in Fig. 8F. It should be noted that these additives adjust rheological behaviors for printing while also optimize the texture and the oral processing of the prints that are edible. For example, the addition of whey protein isolate into the milk protein inks altered microstructures of the 3D printed structures, and thus increased the springiness, cohesiveness, and chewiness, but decreased the gumminess [275]. Furthermore, egg white protein has been developed as a highly transparent and conductive liquid, which was used as an ink for 3D printing for bioelectronics and as a triboelectric nanogenerator [278]. In this study, strong alkaline solutions were used to hydrolyze the egg white proteins, leading to higher conductivity (20.4 S/m). Because both egg white and milk proteins have been useful for 3D cultures of cells and angiogenesis assays [279–283], the ink based on these proteins holds promise for 3D printed tissue scaffolds.

4. Conclusions and perspectives

3D printing with protein-based inks takes advantage of both the material benefits of proteins and the versatile fabrication capability of 3D printing, which has shown the potential to change the landscape of biomedical and clinical research. A wide range of

protein materials, nature-derived and recombinant, have been used as essential components of inks for 3D printing with considerable biomedical and clinical interests. 3D printing with protein-based inks brings a new level of the capability to construct structures that underpin disease pathogenesis and contribute to regenerative medicine, as well as to support drug delivery and screening. Despite the substantial promise and rapid progress in recent years, 3D printing with protein-based inks is still in its infancy, with many challenges that require interdisciplinary endeavors between polymer science, molecular biology, rheology, physical chemistry, computational simulations, microfluidics, and biotechniques.

In particular, one major challenge is the weak mechanical strength of most protein prints [287], especially for monolithic protein inks in comparison to their natural counterparts. This issue of weak strength also limits structural integrity and shape complexity, thus restricting the broad utility of protein prints. Except for the 3D printing of silk fibroin using bioinspired aqueous salt solution [40], monolithic protein inks are rare to print overhanging filaments and perfusable conduits. Some techniques have been developed to address this issue, including covalent crosslinking [39], suspension baths [39,174], and the addition of structural enhancing materials [22,288]. However, non-protein materials and potentially toxic conditions, even approved by the FDA, may impair the merits of protein materials.

Thus, it is reasonable to understand the fundamental molecular mechanisms of high-performance protein tissues for developing better 3D printing with protein inks. These mechanisms embrace the encoded sequence information, hierarchical structures of protein tissues, and more importantly, the processing principles to govern macromolecular assembly. To recapitulate the assembly process in nature, but here in a 3D printing process, is a route to capture the unique and important mechanical properties of protein tissues.

Besides their unique assembly properties as macromolecular building blocks, various protein materials possess other polymeric properties which are valuable to exploit for 3D printing. For example, several proteins, including silk [75], suckerins [289], zein [290,291], and keratin [268], are thermoplastic. These proteins can be extruded and molded with appropriate thermal and pressure processes. The thermoplastic properties of these proteins represent a promising mechanism for fused deposition modeling (FDM) widely used for 3D printing with traditional thermoplastics [292] and more recently, liquid crystal polymers [10,293]. The benefits of the thermoplastic approach include the long shelf-storage of dry protein-based inks and the streamlined printing process for scale-up.

The functionality of 3D prints has been growing in emphasis [294], with protein-based inks particularly promising toward this goal. Protein inks, instead of synthetic materials, such as PEG, polydimethylsiloxane (PDMS), and PCL, bring new opportunities to responsive actuators [295,296], *in vivo* 3D printing [297,298], as well as to the construction of microphysiological systems and tissue-chips [299]. Moreover, 3D printed protein structures have potential in bio-optoelectronics with improved capacity to interface between biology and devices, especially for on-skin applications and *in vivo* implantation [300–303].

Declaration of Competing Interest

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

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