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Computer-aided hydrogel synthesis for 3D bioprinting: application of design of experiments (DoE), machine learning (ML), and computational fluid dynamics (CFD)

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The hydrophilic polymeric network of hydrogels is a crucial element of bioinks, closely resembling the extracellular matrix and offering a supportive microenvironment vital for sustaining cell viability. Crucial rheological properties, especially viscosity and shear-thinning characteristics, are essential in influencing the printability and structural integrity of bioprinted constructs. Additionally, hydrogels must demonstrate suitable mechanical properties to support three-dimensional structures after printing and promote cellular proliferation and differentiation. This review highlights the incorporation of advanced methodologies such as design of experiments (DoE), machine learning (ML), and computational fluid dynamics (CFD) in the systematic optimization of hydrogel formulations for 3D bioprinting applications. For example, DoE, specifically response surface methodology, has been utilized to optimize the concentrations of essential components, such as gelatin, alginate, and methylcellulose, resulting in excellent extrusion rheological characteristics. Simultaneously, machine learning techniques are progressively employed to model and automate the optimization process, diminishing dependence on trial-and-error experimentation and expediting bioink development. This review emphasizes the importance of a balanced strategy for improving the rheological and mechanical properties of hydrogels, which may be effectively realized through the combined use of DoE, ML and CFD approaches in 3D bioprinting.

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1. Introduction

Fused deposition modelling (FDM), developed from stereolithography in 1983 and patented in 1989,^{1,2} is a form of 3D printing that can fabricate complex geometries at low cost with straightforward operational requirements. Due to its advantages, FDM has been widely used across aerospace, medical, and automotive sectors.^{3–5} 3D bioprinting is a technique that integrates additive manufacturing and biomaterial deposition and has emerged as a powerful platform for the spatially controlled fabrication of biological constructs.^{6,7} These days, 3D bioprinting expands its relevance in regenerative medicine, tissue engineering,^{8,9} drug delivery,^{10,11} and the development of implantable or organ replacements.¹²

Despite the significant promise in the biomedical field, the widespread application of 3D bioprinting remains constrained by several critical challenges. Chief among these are the difficulties in achieving high-resolution cell deposition, controlling cell distribution, or selecting the ideal bioink.¹³ An ideal bioink must exhibit appropriate physicochemical characteristics,¹⁴

including biocompatibility,^{15,16} suitable rheology,¹⁷ and mechanical properties, to support and preserve cellular viability and function throughout the printing process.¹⁸ Furthermore, achieving high printing accuracy and speed requires high resolution, which requires selecting the appropriate printing technique and a suitable bioink formulation.¹⁹

A major limitation in 3D bioprinting for tissue engineering lies in the scarcity of bioinks.²⁰ The bioinks currently in use are based on natural and synthetic polymers.²¹ While most natural polymers are cell-interacting and biocompatible,²² they frequently lack the mechanical properties needed to maintain structural integrity and withstand physical stresses *in vivo*. To address these deficiencies, hydrogels have emerged as promising candidates due to their tuneable viscoelastic properties, high water content, and ability to support printability and cellular function. Their porous networks facilitate diffusion of oxygen and nutrients, enhance cellular mobility, and promote adhesion and proliferation within the extracellular matrix.^{23,24} However, to achieve efficient hydrogel use in 3D bioprinting, it is necessary to accurately determine the



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hydrogel material formulation suitable for each specific application.²⁵

The precise selection and optimization of hydrogel formulations are critical for achieving the desired rheological properties and printability of bioinks. The design of experiments (DoE) methodologies, particularly Box–Behnken design (BBD) and central composite design (CCD),²⁶ enable simultaneous study of multiple factors,²⁷ including polymer concentration,²⁸ cross-linker type, pH, and temperature, to adjust hydrogel properties such as strength, biocompatibility,²⁹ and viscosity.^{30,31} Such multivariate optimization not only elevates reproducibility and bioink performance but also streamlines the synthesis process, reducing time and cost, thereby facilitating the translation of hydrogel-based materials into practical applications in regenerative medicine and tissue engineering.

Complementing DoE strategies, machine learning (ML) has advanced rapidly over the past two decades, demonstrating its capabilities in pattern recognition and parameter optimization across diverse manufacturing processes, including metalworking and additive printing.^{32,33} Current applications of ML in 3D bioprinting include optimizing material properties to ensure reliable printability and shape fidelity, regulating the process with the desired fiber and droplet dimensions, and refining structural design to improve cell–microenvironment interactions. Conventional ML models have been applied for prediction, classification, and anomaly detection, underscoring their potential to accelerate design cycles and reproducibility in biofabrication workflows.³⁴

In parallel, computational fluid dynamics (CFD) serves as a critical tool for evaluating key parameters in the 3D printing process, including nozzle speed, shear force, printability, and cell viability.³⁵ CFD simulations enable detailed analysis of fluid flow in the printhead and during bioink deposition, facilitating elucidation of relationships among hydrogel bulk flux, nozzle geometry, shear forces, and print path morphology.³⁶ CFD applications are not limited to nozzle

design but also simulate the interplay between bioink characteristics and nozzle design with cell viability during extrusion.³⁷ Widely used software platforms such as OpenFOAM, ANSYS Fluent, COMSOL Multiphysics, and FLOW-3D have been used in advancing these simulations.^{38,39}

This review provides a comprehensive overview of 3D bioprinting with particular emphasis on hydrogel-based bioinks, including their classification, structures, synthetic strategies, physicochemical properties, and characteristics. Additionally, the integration of DoE, ML, and CFD approaches in the development and optimization of hydrogel systems for bioprinting is critically discussed.

2. Overview of 3D bioprinting

Traditionally, biological research has been conducted using two-dimensional (2D) rigid platforms such as glass slides, tissue culture plates or animal models. Nonetheless, these methods face many limitations, particularly in accurately replicating and observing physiological characteristics. To enhance the effectiveness of biofabrication techniques, researchers have advanced toward mimicking biological systems through various enabling technologies, with 3D bioprinting emerging as a key approach.⁴⁰ This technique is described as “the application of computer-assisted processes to design and construct living and non-living components in predetermined 2D or 3D configurations, forming engineered biological constructs”.⁴¹

2.1 Bioinks – hydrogels

Biofabrication is an evolving research discipline focused on generating tissue constructs with hierarchical structural organization. Traditional biofabrication approaches include particulate leaching, lyophilization (freeze-drying), electrospinning, and microfabrication.⁴² While each of these methods can produce 3D constructs from a variety of biomaterials, they typically suffer from limited reproducibility and fabrication flexibility. Recently, 3D bioprinting has emerged as an innovative biofabrication technique, significantly enhancing the control and reproducibility of tissue construct fabrication through automated deposition processes.^{17,43} In essence, 3D bioprinting enables the production of 3D tissue constructs with pre-defined architectures and geometries that incorporate biological materials and/or living cells (collectively termed bioinks) by synchronizing the deposition and crosslinking of bioinks with the movement of a mechanical stage.⁴⁴ The principal 3D bioprinting modalities include inkjet printing, extrusion-based, and laser-assisted bioprinting,¹⁷ as shown in Fig. 1.

Inkjet bioprinting is a non-contact printing technique capable of depositing bioink in picolitre-scale droplets (1–100 pL) onto substrates with micrometer-scale resolution.⁴⁵ During operation, droplets with diameters of approximately 10–50 μm are rapidly generated *via* either thermoelectric or piezoelectric actuation and expelled through a small orifice at the tip of the reservoir.^{46,47} In thermal inkjet printing, a heating



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gel materials for 3D-bioink applications. His recent work integrates materials synthesis, response surface methodology, nanostructured platforms, and physicochemical characterization to support reproducible development of advanced biomedical materials.



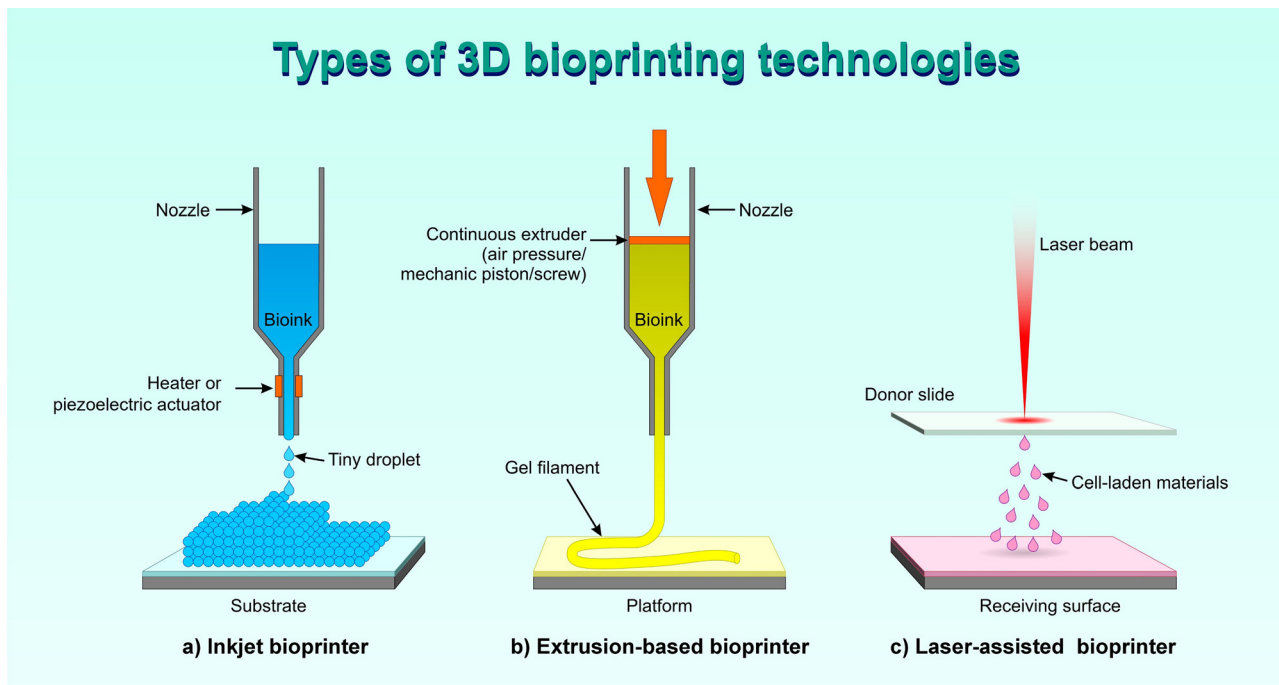


Fig. 1 Types of 3D bioprinting technologies. (a) Inkjet/biodroplet printing. (b) Extrusion-based bioprinting. (c) Laser-assisted bioprinting.

element vaporizes a small volume of bioink inside the reservoir, creating a vapor bubble that propels a droplet through the nozzle. This process briefly exposes encapsulated cells to high temperatures (~ 300 °C) for a few microseconds (~ 2 μ s), inducing transient pore formation in the cell membrane. By contrast, piezoelectric inkjet systems generate and eject droplets *via* mechanical deformation of a piezoelectric transducer (actuated by an external voltage), thereby avoiding significant temperature increases.^{48,49} To prevent nozzle clogging, bioinks used in inkjet printing are formulated with low viscosity (on the order of 1–10 mPa s) and relatively low cell densities (typically below 10^6 cells per mL).⁵⁰

In a typical extrusion-based 3D bioprinting setup, cells suspended in a prepolymer solution are loaded into a syringe or a disposable cartridge and then extruded onto a flat substrate *via* compressed air or mechanical actuation from a rotating piston or screw.⁴³ Temperature regulation modules are often incorporated into the bioprinter to maintain the bioink at an optimal temperature during printing, which is critical for controlling viscosity and inducing *in situ* solidification.⁵¹ In addition, light sources and specialized nozzle configurations may be integrated to facilitate *in situ* cross-linking of the bioink, thereby enhancing print fidelity and the structural stability of the resulting constructs.^{52,53} A major advantage of this approach is its capacity to process highly viscous, high-density bioinks into 3D constructs of clinically relevant size, a capability not achievable with inkjet or laser-assisted bioprinting. Conversely, significant limitations include relatively low resolution (around 200 μ m), frequent nozzle clogging, and challenges in fabricating constructs that retain their intended geometry and provide a suitable

microenvironment.^{43,50} Indeed, fabricating 3D constructs with complex microarchitectures remains a significant challenge in bioprinting. To address these challenges, recent strategies have explored combining multiple crosslinking mechanisms (*e.g.*, photochemical and thermal gelation),⁵⁴ and the use of partially crosslinked hydrogels has been investigated with promising results.⁴⁹

Laser-assisted bioprinting utilizes a high-energy pulsed near-infrared laser to irradiate a donor substrate coated with the target bioink, thereby generating a localized jet of droplets. In practice, the laser pulses are focused onto a transparent support (*e.g.*, glass or quartz) coated with a thin metallic layer (commonly gold or titanium) that absorbs the laser energy and transfers it to the bioink. This absorption triggers the rapid formation of a high-pressure vapor bubble, which propels a small bioink droplet toward the receiving substrate.^{43,54} Unlike inkjet printing, laser-assisted bioprinting does not suffer from nozzle clogging and can deposit bioinks across a wide viscosity range (1–300 mPa s⁻¹) and at very high cell concentrations (on the order of 10^8 cells per mL).⁴³ However, achieving high-resolution 3D constructs requires biomaterials that rapidly photo-crosslink upon exposure to the specific laser wavelength, effectively necessitating advanced photo-reactive bioink formulations.⁵⁰

2.2 Factors affecting 3D bioprintability

Printability is an important characteristic of bioinks, as it often decides the extrudability of bioink and the formability of filaments after extrusion.⁵⁵ Thus, selecting appropriate bioinks is a critical step in bioprinting processes, particularly with respect to the characteristics of the chosen materials



(e.g., compatibility with specific printers and achievable resolution) and how these characteristics influence the printing process.⁴⁰ Generally, the printability of bioinks is affected by rheological properties, cross-linking mechanisms, and printing conditions.⁵⁶

The rheological characteristics of bioinks, characterized by viscosity, shear stress, viscoelastic shear moduli, and elastic recovery, significantly influence their printability.⁵⁷ Recent efforts to optimize bioinks have focused on improving printability and shape accuracy, often through increasing viscosity. However, the ink's bioproperties could be affected, as higher viscosity requires higher extrusion pressure, which can be detrimental to cell viability,⁵⁵ and too-viscous bioinks frequently result in poor yield and a non-homogeneous cell distribution.⁵⁸ Potent biomaterials should have an adjustable viscosity that can change with temperature and exhibit shear thinning behaviour, as required by different printing methods. Hydrogels exhibiting shear-thinning behaviour are widely regarded as optimal for 3D bioprinting, as they can flow under extrusion and simultaneously protect encapsulated cells from shear stress.⁴⁰ Elevating the polymer concentration typically enhances the material's rheological properties. Rheological modifiers such as gelatin or methylcellulose can be incorporated to impart shear-thinning behaviour.^{59–61}

The gelation strategy governs compatibility with the selected 3D bioprinting platform, and the gelation duration dictates whether supplementary support structures are required during printing.⁴⁰ Mechanisms like photoinitiated crosslinking or thermal gelation stand out as particularly effective options because they can solidify quickly while remaining gentle and cytocompatible, which helps protect encapsulated cells from stress and allows the printed structure to retain its shape right after extrusion.

The inherent biological properties of a bioink formulation play a central role in shaping cellular responses. Important characteristics to evaluate experimentally include the presence of cell adhesive ligands that encourage attachment and the hydrogel matrix's ability to degrade under standard *in vitro* culture conditions. For instance, research has shown that making alginate based bioinks more susceptible to enzymatic degradation leads to markedly improved cell attachment and proliferation throughout the bioprinted construct.⁴⁰ These qualities help the material better replicate aspects of the natural extracellular matrix and support more favourable long-term cellular behaviour.

The mechanical properties of bioinks, especially the elastic modulus that reflects stiffness, also have a substantial impact on how cells behave. Differences in modulus can guide important processes such as proliferation, migration, and differentiation by transmitting mechanotransduction signals that cells sense from their surrounding matrix.⁴⁰ Carefully tuning these biophysical features therefore becomes essential when designing constructs for biological or tissue engineering purposes.

2.3 Applications

3D bioprinting enables the generation of biological constructs that faithfully replicate native anatomical structures, offering

valuable benefits across clinical, research, and educational fields. This approach is widely applied to fabricate cell-laden scaffolds for the regeneration of tissues such as bone, myocardium, cartilage, liver, and lung.

In bone tissue research, 3D bioprinting shows a significant impact on bone treatment and healing by enabling the creation of detailed shapes and greater precision and control over structural components. Gao *et al.* studied the printing of scaffolds using bioinks composed of human mesenchymal stem cells and nanoparticles, such as bioactive glass or hydroxyapatite.⁶² Their results indicated that combining hydroxyapatite with the scaffold significantly promoted osteogenic differentiation and osteogenic extracellular matrix production, with a cell viability of $86.62 \pm 6.02\%$. Besides, the combined scaffolds exhibited a more homogeneous cell distribution, enhanced collagen synthesis, and significantly increased compressive modulus (358.91 ± 48.05 kPa) after 21 days of culture. A study by Chou *et al.* developed a biodegradable 3D-printed polylactide cage that combines an antibiotic-embedded poly(D,L)-lactide-co-glycolide nanofibrous membrane for treating comminuted fractures in rabbit models.⁶³ The results demonstrated that rabbits receiving the 3D-printed cage implant exhibited improved cortical integrity, leg length ratio, and maximal bending strength.⁶³ In this study, the 3D-printed PLA cage served as a biodegradable, shape-defined construct capable of capturing comminuted bone fragments, filling the metaphyseal defect, and assisting the intramedullary fixation in maintaining bone length and alignment, whereas the PLGA nanofibrous membrane functioned as the antibiotic-releasing component. This dual-module design differs from conventional hydrogels,⁶⁴ which mainly act as resorbable drug depots, and from PMMA cement,⁶⁵ which can provide local antibiotic delivery but is non-biodegradable and constrained by heat-related and drug-compatibility limitations. The key contribution of 3D printing is therefore the ability to fabricate a defect-conforming, mechanically supportive, biodegradable cage with controlled architecture, potential extension toward fracture pattern specific or patient matched implants. In cartilage tissue engineering, layered bioprinted scaffolds mimic the native stratified distribution of cells and the extracellular matrix, with precise control over geometry to facilitate chondrocyte differentiation. Incorporation of growth factors, such as TGF- β 1 and FGF-2, further enhances glycosaminoglycan production, thereby optimizing the functional properties of the printed cartilage.^{66,67}

In the cardiovascular realm, 3D bioprinting has enabled the fabrication of anatomically precise cardiac muscle constructs and heart valve-like structures, which can be further electrically stimulated to restore contractile function. In parallel, coaxial nozzle bioprinting has emerged as a particularly valuable strategy for vascular tissue engineering because it enables the direct fabrication of tubular or core-shell vascular channels through concentric material deposition. In this configuration, the crosslinker flowing through the core stream contacts the hydrogel precursor in the shell stream, leading to rapid *in situ* gelation and immediate formation of hollow tubular



constructs. This direct core-shell fabrication is difficult to achieve using conventional inkjet or laser-based techniques, which commonly rely on layer-by-layer assembly, scaffold-free spheroid fusion, or fugitive templates to generate perfusable channels after post-print processing. Recent studies further support the unique capability of coaxial bioprinting to produce vascular-like structures with controlled lumen geometry, multi-material organization, and improved relevance for cardiovascular models, including human coronary artery-sized endothelialized constructs.⁶⁸

For hepatic applications, 3D bioprinting has advanced beyond the simple encapsulation of hepatocytes within bulk hydrogels by enabling the spatially controlled organization of hepatic and non-parenchymal cells into liver-mimetic architectures. Bioinks containing hiPSC-derived hepatocyte-like cells, primary hepatocytes, or hepatocyte-supporting cell co-cultures can be printed into multilayered or lobule-like constructs that partially recapitulate key architectural features of native liver tissue, including defined parenchymal or non-parenchymal cell placement, hexagonal lobule-inspired geometry, and engineered microchannels or sinusoidal-flow environments. Such structural control is important because native liver function depends strongly on microscale organization, heterotypic cell-cell interaction, and mass transport. Original research has shown that rapid 3D bioprinting can pattern hiPSC-derived hepatic progenitor cells together with endothelial and mesenchymal supporting cells in microscale hexagonal units, leading to enhanced hepatic maturation, liver-specific gene expression, metabolic secretion, and cytochrome P450 inducibility.⁶⁹ Similarly, 3D bioprinted primary human liver tissues have been used to assess organ-level drug-induced liver injury and to discriminate hepatotoxic responses among clinically relevant compounds.⁷⁰ Therefore, the major value of liver bioprinting is not only in providing a permissive hydrogel microenvironment, but also in introducing reproducible, designable tissue architecture that improves physiological relevance for drug metabolism, toxicity screening, disease modelling, and future regenerative applications.

In neural engineering, scaffold free 3D printing of Schwann cells and bone marrow derived stem cells within agarose mold drives self-assembly into nerve grafts, which, upon implantation in animal models, show promise for functional neural regeneration after injury.⁷¹ Building on this foundation, bioprinting is now being applied more broadly in neural engineering to address both peripheral and central nervous system challenges. For peripheral nerve repair, advanced 3D-printed bionic scaffolds loaded with neural crest stem cell-derived Schwann cell progenitors have been developed to guide oriented axonal extension and significantly enhance myelination compared to conventional nerve conduits.⁷² Similarly, hydrogel-based constructs containing living Schwann cells printed in aligned architectures maintained high post-printing viability (>89%) and promoted directed neurite outgrowth *in vitro*.⁷³ In central nervous system applications, 3D bioprinting strategies for spinal cord injury repair have incorporated neural stem cells and glial populations within

biomimetic architectures designed to recapitulate native tissue organization and mechanical properties, with several preclinical studies reporting enhanced axonal regeneration and improved functional outcomes.⁷⁴

Skin tissue engineering represents one of the most intensely pursued applications of 3D bioprinting. This technique facilitates the direct deposition of stratified skin layers, including an epidermal layer populated by keratinocytes and a dermal compartment containing fibroblasts within an extracellular matrix. Such bioprinted constructs closely emulate the architecture of native skin, enhancing graft integration and accelerating wound closure. Notably, *in situ* 3D bioprinting whereby skin is printed directly onto the wound bed has demonstrated superior regenerative outcomes compared to conventional dressings and grafting methods.⁷⁵ Beyond cutaneous applications, 3D bioprinting has also been extended to pancreatic tissue fabrication: endocrine cell-laden constructs are printed to serve as therapeutic implants for diabetes management; however, optimizing insulin secreting cell function remains challenging due to ionic interactions (*e.g.*, calcium) during crosslinking processes.⁷⁶

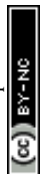
Moreover, 3D bioprinting is increasingly leveraged to develop *in vitro* cancer models that faithfully replicate the tumour microenvironment, thereby advancing both basic oncological research and drug development efforts:⁷⁷ (i) 3D tumour constructs allow the systematic spatial arrangement of malignant cells alongside stromal and endothelial cell populations, recreating the layered organization of actual tumours. These models enable detailed investigation of cell-cell and cell-matrix interactions, as well as gradients of oxygen, nutrients, and growth factors, offering critical insights into mechanisms of tumour proliferation, invasion, and metastasis. (ii) Bioprinted tumour tissues also serve as high fidelity platforms for anticancer drug screening, improving the predictive accuracy of *in vitro* assays relative to traditional two-dimensional cultures. Consequently, this approach accelerates the identification of promising therapeutic compounds, optimizes dosing regimens, and reduces variability arising from the absence of a realistic tumour microenvironment.

3. Overview of hydrogels

3.1 Definition and classification

Hydrogels are three-dimensional (3D), crosslinked polymeric networks capable of absorbing large amounts of water while maintaining structural integrity, rendering them highly advantageous for biomedical applications, including tissue engineering and 3D bioprinting.⁷⁸ Their structural and chemical diversity enables precise tuning of mechanical, rheological, and biological properties, which is critical for advanced biofabrication strategies.⁷⁹

The defining characteristic of hydrogels lies in their cross-linked network architecture, which governs their mechanical strength, swelling behavior, and responsiveness to external stimuli. The classification of hydrogels is shown in Fig. 2. Based



CLASSIFICATION OF HYDROGEL

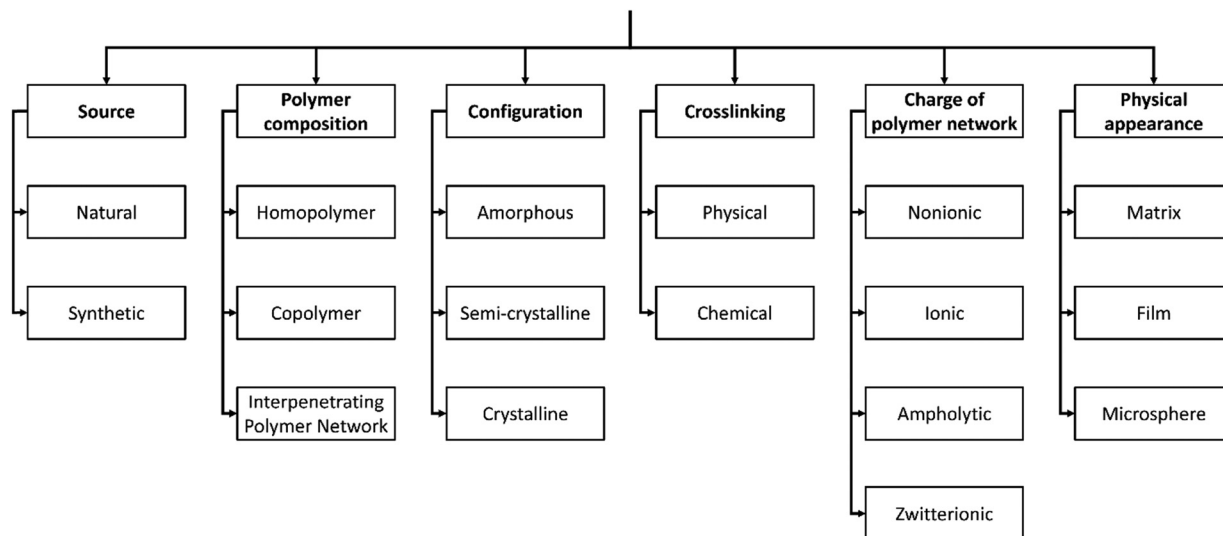


Fig. 2 Classification of hydrogels.

on crosslinking mechanisms, hydrogels are broadly categorized into physically and chemically crosslinked systems.⁷⁸ Physically crosslinked hydrogels are formed through reversible non-covalent interactions among polymer chains, including hydrogen bonding,^{80,81} ionic interactions,⁸² hydrophobic interactions,⁸³ and other secondary molecular forces.⁸⁴ These transient junctions enable dynamic and stimuli-responsive behavior but generally provide lower mechanical robustness compared to covalent networks. In contrast, chemically crosslinked hydrogels rely on permanent covalent bonds to establish stable 3D structures, resulting in improved mechanical integrity and structural resilience. The physical and chemical crosslinking mechanisms in hydrogels are illustrated in Fig. 3. Such networks can be formed through thermal polymerization,⁸⁵ photopolymerization,^{85–87} enzymatic crosslinking,⁸⁸ and other chemical approaches.⁸⁹

Beyond crosslinking type, hydrogel networks exhibit diverse structural organizations that influence functional performance. Based on configuration, hydrogels may be amorphous, semi-crystalline, or crystalline, reflecting differences in structural order and mechanical properties.⁷⁹ More advanced architectures include gradient, anisotropic, microstructured, and nanostructured hydrogels. Gradient hydrogels display spatial variations in polymer composition or crosslinking density, often generated *via* controlled mixing of polyethylene glycol (PEG) precursors to create stiffness gradients that guide cellular migration.^{90,91} Anisotropic hydrogels exhibit direction-dependent properties achieved through aligned fibers or pores, typically introduced *via* directional freezing or mechanical stretching during gelation.⁹⁰ Microstructured hydrogels incorporate micron-scale patterns using techniques such as photolithography on PEG-DA or HEMA precursors,⁹² whereas nanostructured hydrogels constructed through nano-molding, initiated chemical vapor deposition, or supramolecular

assembly contain nanoscale features such as nanofibers or nanotubes that enhance functionality.⁸⁰

Hydrogels can also be classified according to origin, composition, charge, and morphology. Based on source, they are categorized as natural (*e.g.*, collagen, gelatin, alginate, and chitosan) or synthetic (*e.g.*, polyacrylamide, PEG, and polyvinyl alcohol (PVA)).⁹³ In terms of polymer composition, hydrogels may be homopolymers, composed of a single repeating monomer unit;⁹⁴ copolymers, consisting of multiple monomers arranged in random, block, or alternating sequences;⁹⁵ or interpenetrating polymer networks, comprising two entangled networks, including semi-interpenetrating polymer networks with one uncrosslinked polymer for enhanced elasticity and recovery behavior.⁹⁶ Network charge further distinguishes hydrogels into nonionic, ionic, ampholytic, and zwitterionic systems, the latter containing both positive and negative charges within each monomer unit.⁷⁹ Morphologically, hydrogels may exist in structures as bulk matrices, films, or microspheres, each tailored to specific biomedical applications.⁷⁹

3.2 Characterization of hydrogels

3.2.1 Degree of substitution. The degree of substitution (DoS) quantifies the average number of substituent groups introduced per monomer unit in a polymer chain and is a key parameter in hydrogel synthesis. It reflects the extent of chemical modification and is typically calculated as:⁹⁷

$$\text{DoS (\%)} = \left(1 - \frac{\text{apparent sample conc}}{\text{nominal sample conc}} \right) \times 100\%$$

Various techniques can be used to determine DoS, depending on the polymer matrix and substituent groups. The colorimetric method using reagents such as ninhydrin or 2,4,6-trinitrobenzenesulfonic acid is commonly employed to quantify



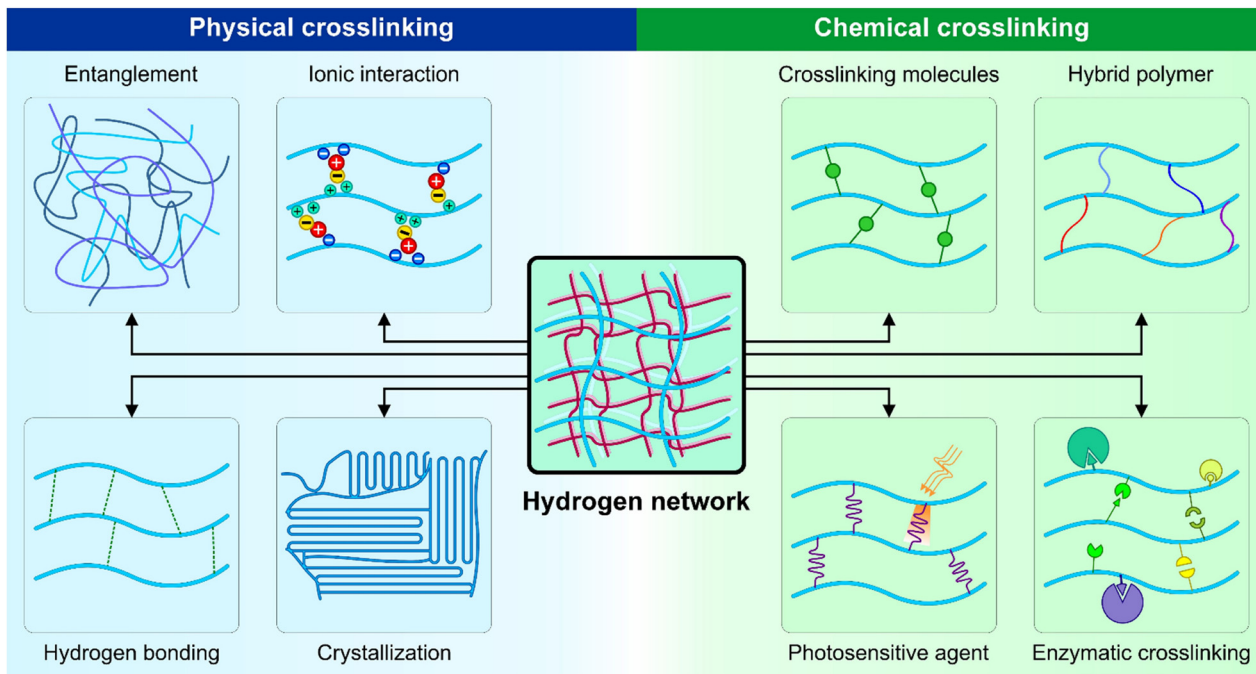


Fig. 3 Diagrammatic representation of the various physical and chemical crosslinking mechanisms in hydrogels.

unreacted amines after reaction with methacryloyl groups in GelMA hydrogels.⁹⁸ More precise measurements can be obtained using ¹H NMR spectroscopy by integrating characteristic signals of functional groups introduced during modification.⁹⁹

DoS has a significant effect on the physicochemical properties of hydrogels, particularly in 3D bioprinting. An increase in DoS enhances crosslinking density, stiffness, and structural stability, while also reducing swelling capacity.^{99–101} Highley *et al.* (2015) demonstrated that tuning DoS enables the fabrication of self-healing, shape-retaining hydrogels.¹⁰²

3.2.2 Morphology. The morphological characteristics of hydrogels, particularly porosity and pore size, are critical for biomedical applications such as tissue engineering and 3D bioprinting. A porous structure not only facilitates water uptake but also promotes cell distribution and tissue ingrowth.^{103–105} Pore size greatly affects cellular infiltration and nutrient diffusion, with optimal ranges varying by tissue type: 5–15 μm for fibroblasts, 20–125 μm for skin, and 100–350 μm for bone regeneration.¹⁰⁶

Hydrogel morphology is commonly characterized by scanning electron microscopy (SEM), following freeze-drying (freeze-drying) to preserve structural integrity. SEM enables high-resolution imaging of hydrogel structures, including pore size, distribution, and network connectivity.¹⁰³ Studies using SEM have shown that hydrogels typically exhibit highly porous surfaces with large surface areas, which significantly affect swelling ability and mechanical properties.¹⁰⁷ In addition, SEM is also used to study the effects of environmental factors such as pH and ionic strength on the swelling behaviour of hydrogels, providing important information for tailoring the application of hydrogels in various fields.¹⁰³

Notably, porosity and pore size in hydrogels can be modulated through compositional adjustments. Chen *et al.* reported that the amino group content was inversely proportional to the pore size of GelMA hydrogels.¹⁰⁸ Similarly, Celikkin *et al.* demonstrated that increasing the GelMA concentration reduced both porosity and average pore size.¹⁰⁹ These findings highlight the tunability of hydrogel microstructure, offering strategic opportunities for material optimization in 3D bioprinting.

3.2.3 Mechanical and rheological properties. In 3D bioprinting, the mechanical and rheological properties of hydrogels are critical for ensuring printability, post-printing structural durability, and the ability to support tissue growth. An ideal 3D bioink should exhibit appropriate viscosity, pronounced shear-thinning behaviour, rapid structural recovery, and an optimal crosslinking capacity to maintain its shape after deposition. However, a single universal viscosity value cannot be defined because the printable window depends strongly on the printing modality, nozzle diameter, extrusion pressure, temperature, cell density, and crosslinking mechanism. For extrusion-based bioprinting, which is among the most widely used approaches for hydrogel bioinks, printable materials have been reported across a broad viscosity range of approximately 30 mPa s to 10^7 mPa s; however, many practical hydrogel inks for direct ink writing or extrusion fall within a low-shear viscosity window of about 10^2 – 10^6 mPa s.¹¹⁰ In contrast, inkjet bioprinting generally requires much lower viscosities, commonly below 10–20 mPa s, to enable stable droplet ejection and avoid nozzle clogging.¹¹⁰

For extrusion-based printing, shear-thinning behaviour is particularly desirable because the viscosity decreases under high shear inside the nozzle, facilitating smooth extrusion



and reducing the mechanical stress imposed on encapsulated cells. After deposition, viscosity and elastic recovery should increase rapidly to preserve filament shape and prevent spreading or collapse. This behaviour is commonly described using the power-law or Herschel–Bulkley models, where a flow index (n) < 1 indicates shear-thinning behaviour; values around $n \approx 0.3$ – 0.6 are often considered indicative of pronounced shear thinning suitable for smooth extrusion. In addition, a moderate yield stress, typically in the range of 10–1000 Pa, and rapid post-printing recovery, preferably within seconds, are useful design features for maintaining shape fidelity. Therefore, bioink optimization should balance low apparent viscosity during extrusion, sufficient viscosity and elasticity after deposition, and cytocompatible crosslinking to achieve both printability and biological performance.¹¹¹

There are many methods to evaluate the mechanical properties of hydrogels including compression and tension tests to determine the strength and elasticity of the material; dynamic mechanical analysis to test the ability of hydrogels to withstand vibration and deformation; and rheological tests to evaluate the rheological properties and viscosity of hydrogels, which are especially important in 3D printing applications.^{112,113}

Rheology describes how hydrogels deform and flow under external forces, reflecting their viscoelastic nature possessing both solid-like and liquid-like behaviour. Key parameters include storage modulus (G'), loss modulus (G''), and loss factor ($\tan \delta$), which are crucial for evaluating printability and structural stability.¹¹⁴ G' indicates elastic energy storage and mechanical strength, with higher values supporting shape retention after printing. G'' reflects viscous energy dissipation and flowability, facilitating extrusion but potentially compromising shape fidelity. The $\tan \delta$ (G''/G') ratio distinguishes material behaviour: $\tan \delta < 1$ indicates elastic-dominant, while $\tan \delta > 1$ suggests viscous-dominant behaviour.^{114,115} The linear viscoelastic region (LVR) defines the strain range where G' and G'' remain stable; printing within this region ensures mechanical integrity and shape recovery.¹¹⁵

Cross-linking conditions and polymer composition affect the gelation kinetics and mechanical strength of the hydrogel. Jeong *et al.* reported a temperature-triggered sol–gel transition in chitosan-graft-(PEG-PAF), determined by the change in the correlation between G' and G'' .¹¹⁶ Moura *et al.* demonstrated enhanced elastic moduli in genipin-crosslinked chitosan, indicating denser network formation.¹¹⁷ A physically cross-linked PAA/sodium alginate hydrogel incorporating amorphous calcium carbonate exhibited self-healing behavior and exceptional flexibility, enabling conformation to complex surfaces.¹¹⁸

4. Optimization of hydrogel synthesis for 3D bioinks

4.1 Application of DoE in hydrogel synthesis optimization

Traditionally, the development and optimization of biomaterials have relied on one factor-at-a-time approach, which, while widely used¹¹⁹ is time-consuming, costly, and inadequate for

capturing complex factor interactions,¹²⁰ resulting in delays in clinical translation.¹¹⁹ Over the past decade, these traditional methods have been gradually replaced by statistical testing methods such as DoE. DoE enables the simultaneous variation of multiple parameters, thereby increasing experimental efficiency, identifying key contributors to material performance, and supporting predictive model development to streamline optimization.^{121,122}

DoE is a statistical method that enables systematic planning and conducting experiments to efficiently extract maximal information from each trial. By elucidating the relationships between independent and dependent variables, DoE facilitates the identification and optimization of critical parameters using a reduced number of experiments.^{123,124} To enhance the reliability and objectivity of experimental outcomes, DoE incorporates foundational principles such as replication, randomization, and blocking.¹²⁵

In DoE, response surface methodology (RSM) employs polynomial regression models to investigate the nonlinear effects of multiple factors and determine optimal conditions for the desired response variables.^{126–128} Commonly used RSM designs include central composite design (CCD) and Box–Behnken design (BBD), among others.^{129,130} Considering specific experimental applications will help clarify the high applicability and optimization efficiency of DoE in hydrogel material research. In practice, DoE has been widely applied to identify and optimize the relationships between input parameters such as polymer concentration, photoinitiator content, environmental conditions or synthesis process parameters with important properties of hydrogels including storage modulus, swelling ratio, rheological behavior, printability, and mechanical strength.^{26,131–134} By employing appropriate design models, researchers have not only minimized the number of required experiments but also developed reliable predictive models, thereby guiding the formulation of bioinks tailored to specific requirements in biomedical and tissue engineering applications. Table 1 presents representative studies that have applied various DoE models to optimize hydrogel synthesis, highlighting the role of DoE as a powerful tool for the rational design and development of bioink materials for 3D bioprinting.

The application of response surface methodology in hydrogel optimization has been shown to be effective in improving important properties such as mechanical strength, swelling ability, and rheology. Experimental design models such as CCD, BBD, *etc.* not only reduce the number of experiments required but also provide reliable data to determine the optimal conditions.

4.2 Application of ML in the optimization of hydrogel printability

ML utilizes data-driven statistical relationships to generate predictive models, contrasting with traditional rule-based programming.¹³⁵ In 3D bioprinting, ML is increasingly integrated to reduce development time and enhance print fidelity.³² Although currently in its nascent stages, this integration demonstrates significant potential.^{136,137} Key applications



Table 1 Summary of DoE applications in optimizing synthesis conditions

Model	Hydrogel	Factor	Response	Results	Ref.
CCD	GelMA/HAMA	Concentrations of GelMA, HAMA, and photoinitiator LAP	Storage modulus and diffusion coefficient	The optimal composition was 8.0% (w/v) GelMA, 2.0% (w/v) HAMA, and 0.1% (w/v) LAP, which achieved a maximum storage modulus of 34.98 kPa and an optimal diffusion coefficient of $1.172 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$, suitable for biomedical applications.	131
	Self-assembling peptide gel (SPG-178)	Peptide (SPG-178) concentration, NaCl concentration, Milieu type	Rheological properties of hydrogels	Helps hydrogels achieve the highest stiffness, suitable for applications in hard tissue and other biomedical fields.	132
	Gum tragacanth-acrylic	pH and concentration of the photoinitiator	Swelling	Helps determine the maximum swelling degree of 5307% when $\text{pH} \approx 7$ and at a photoinitiator concentration of about $21\text{--}23 \times 10^{-6} \text{ mol L}^{-1}$	26
BBD	Gel GelMA/HA-Tyr/MC)	GelMA concentration (w/v) HA-Tyr concentration (w/v) Methylcellulose concentration (w/v)	Printability	Helps 3D frames have a stable structure during and after printing, providing greater precision and flexibility	133
Doehlert	GelMA	[MA]/[gelatin] ratio (mL g^{-1}), addition rate of MA (mL min^{-1}), stirring speed (rpm)	Lysine substitution rate, swelling, $\log G'$, compressive modulus	<ul style="list-style-type: none"> - The interaction between temperature and the [MA]/[gelatin] ratio had a positive effect on swelling and a negative effect on $\log(G')$ - The interaction between the [MA]/[gelatin] ratio and MA flow rate had a negative effect on lysine substitution rate and compression at 15% 	134

include optimizing material properties, tuning printing parameters, enabling *in situ* monitoring, and refining scaffold architectures (Fig. 4).³²

Traditional ML relies on pre-extracted numerical (*e.g.*, statistical, frequency-domain) or image-based (*e.g.*, edges, textures) features for predictive tasks. Achieving optimal performance necessitates selecting highly relevant and discriminative features.¹³⁸ Through iterative training to minimize errors, these models optimize their parameters, enabling them to generalize and accurately predict outcomes for novel inputs.

Depending on the data structure and specific predictive task, these ML models generally utilize three primary approaches: supervised,¹³⁹ unsupervised,¹⁴⁰ and reinforcement learning.¹⁴¹ Supervised learning maps labelled input-output

pairs to predict defined outcomes,¹³⁹ whereas unsupervised learning extracts intrinsic patterns from unlabelled datasets.¹⁴⁰ Operating between these paradigms, reinforcement learning optimizes actions through evaluative feedback signals rather than relying on explicit target outputs.¹⁴¹

Applying these approaches to 3D hydrogel bioprinting, ML not only predicts pre-print material properties but also optimizes process stability and printability. Specifically, it deciphers complex relationships between diverse input parameters (*e.g.*, polymer concentration, printing settings, UV exposure) and critical functional outcomes, such as filament quality, structural integrity, and gelation.¹⁴² Table 2 provides a comprehensive summary of the primary algorithmic categories utilized in hydrogel optimization, highlighting their individual

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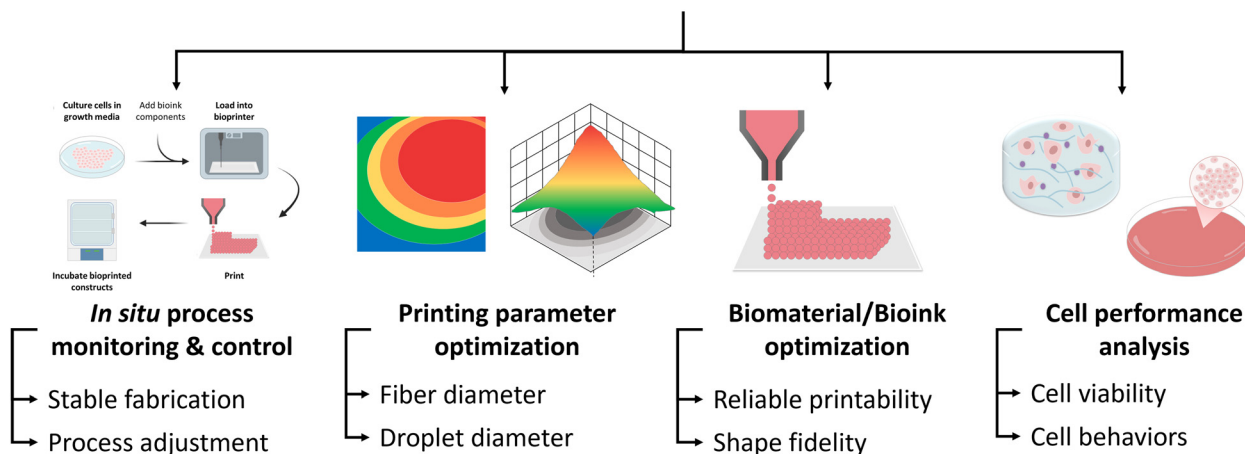


Fig. 4 Applications of machine learning in 3D bioprinting.



Table 2 Support vector learning algorithms employed for the optimization of hydrogels

Algorithm	Task	Inputs (I)/outputs (O)	Input feature/sample size	Advantages (A)/disadvantages (D)	Ref.
Linear regression	Regression		Input feature: single variable Sample size: small	A: a systematic framework for finding optimal solution paths D: applicable solely to datasets exhibiting linear correlations	146
Multiple linear regression	Prediction of outcomes based on multiple variables	I: printing parameters O: print structure	Input feature: ~3 to 10 variables Sample size: small to medium (20–50 samples) ^{156,157}	A: fundamental prediction and identification of critical parameters D: this methodology exhibits constraints to improve 3D hydrogel printing because the linearity assumption inadequately represents the nonlinear interactions among factors. The model is also susceptible to noisy data.	143
Decision tree	Classification/regression	I: printing parameters O: classification of target variables or prediction of material properties	Input feature: ~5 to 20 variables Sample size: medium (50–100 samples) ^{156,158,159}	A: interpretability and the capacity for handling nonlinear interactions D: susceptible to overfitting and sensitive to small alterations	144
Random forest	Classification/regression	I: printing parameters O: prediction of material properties	Input feature: 50 to 100+ variables Sample size: medium to large (>150 samples) ^{158,159}	A: handling nonlinear interactions D: black-box model with limited interpretability and obscured algorithmic structure	145
LASSO	Feature extraction, regression	I: printing parameters O: important features	Input feature: 10 to 50 initial variables Sample size: small to medium (20–50 samples) ¹⁶⁰	A: feature selection emphasises the identification of variables that have the most major effect on the target variable. D: may overlook interactions due to the assumption of linearity	161
Bayesian optimization	Optimal model design	I: process parameters and feedback O: optimal parameters and prediction plane	Input feature: ~2 to 5 core variables Sample size: very small (<50 iterations) ^{160,162}	A: performs well with limited data D: exhibits limited generalizability and presents difficulties in scaling	147
Support vector machines	Classification/regression		Input feature: ~5 to 30 variables Sample size: medium (50–100 samples) ^{156,160}	A: applicable for both classification and regression applications, and capable of integrating Boolean functions B: the system is susceptible to overfitting due to an overwhelming quantity of features, which is accompanied by restricted interpretability of the computational framework.	148

advantages, drawbacks, and applications to clarify the function of various ML methods.

ML utilizes data-driven models to optimize 3D bioprinting, significantly enhancing print fidelity and accelerating development.^{32,136,137,143} By extracting discriminative features,¹³⁸ ML employs supervised,¹³⁹ unsupervised,¹⁴⁰ and reinforcement learning¹⁴¹ paradigms. In hydrogel bioprinting, these models decipher complex, nonlinear relationships between multi-variable inputs (*e.g.*, polymer concentration, UV exposure) and structural outcomes.¹⁴² Specific algorithms address distinct challenges: decision trees and random forests manage nonlinearities and noise,^{144,145} multiple linear regression identifies key linear factors,^{143,146} Bayesian optimization maximizes limited datasets,¹⁴⁷ and support vector machines balance training efficiency with predictive accuracy.^{148,149}

Beyond single algorithms, hybrid ML frameworks significantly enhance predictive performance. For instance,

integrating the least absolute shrinkage and selection operator (LASSO) for feature selection with random forests or gradient boosting yields highly accurate models.¹⁵⁰ For global process optimization, Bayesian optimization excels by efficiently identifying near-optimal parameters (*e.g.*, pressure, speed) from limited experimental trials to maximize “printability scores”.^{147,151,152} Table 3 summarizes these diverse ML applications in optimizing hydrogel synthesis and bioprinting workflows. Specific ML models could be applied to address distinct phases of bioprinting. Bayesian optimization rapidly identifies optimal material compositions (*e.g.*, GelMA/HAMA ratios) and printing parameters to maximize printability with minimal experimental expenditure.¹⁴⁷ For natural bioink formulation, integer linear programming combined with multiple regression ensures requisite mechanical properties without chemical modifications.¹⁴³ Furthermore, integrating random forests with Boruta (feature selection) and SHAP accurately classifies



Table 3 Several ML models are used for synthesis and process optimization in 3D bioprinting of hydrogels

Model	Optimal target	Input	Output	Performance	Ref.
Bayesian optimisation	Evaluate quantitative printability to achieve and accelerate the extrusion printing optimization process	Printing parameters: extrusion pressure, printing speed, temperature and nozzle diameter Composition and concentration of materials (GelMA, HAMA at varying ratios)	Printability score (based on filament formation and layer stacking)	Optimal parameters found with ~19–47 experiments vs. ~6000–10 000 possible combinations (<i>i.e.</i> , major experiment reduction)	147
Inductive logic programming	Design of bioink from natural materials (collagen, fibrin, hyaluronic acid) for stable 3D printing without chemical modification	Collagen concentration (AC): strongly influences elasticity (G')	Storage modulus (G') for shape fidelity	ML models have successfully identified critical mechanical property windows enabling reliable extrusion and shape fidelity, with experimental confirmation affirming a strong agreement between predicted and measured printability.	143
Multiple regression	Ensure appropriate mechanical properties (elasticity, yield stress) to maintain printed shape	Fibrin concentration: partly determines the yield stress	Yield stress (τ_y) for extrudability		
Random forest	Identify the key rheological measures that govern the printability of the hydrogel upon the incorporation of rheological additives	Hyaluronic acid concentration (HA): imparts rheological properties (viscosity) A total of 65 initial rheological features derived from frequency sweep, amplitude sweep, and shear-stress-dependent viscosity tests (including G' , G'' , viscosity, yield stress, damping factor, and flow parameters) across 180 formulations	Printability score: determined based on G' - τ_y combination The 3D printability of each hydrogel was predicted by classifying formulations as “printable” or “not-printable” based on a printability score threshold of ≥ 0.33 .	The model identified 13 critical rheological features strongly associated with printability. Feature reduction improved the $F1$ -score for the printable class from 0.91 to 0.94; for the non-printable class, $F1 = 0.99$; OoB score = 0.96, indicating high generalizability and minimal overfitting.	150
Boruta feature selection					
SHAP					
Linear regression (LR)	Predict and optimize the gel fraction of conductive	3 feature groups:	Gel fraction (%)	Best model: SVR (MAPE = 3.13%, $R^2 = 0.79$)	153
Support vector regression	GelMA-PEDOT:SPSS hydrogels to improve curing efficiency and reduce experimental cost	G1: bioink formulation (GelMA, LAP, PEDOT:SPSS concentration) + UV parameters (intensity, exposure time) G2: absorption coefficient (measured directly using an <i>in situ</i> UV sensor during the curing process)		Random forest regression (MAPE = 3.42%, $R^2 = 0.76$); deep neural network (MAPE = 3.81%, $R^2 = 0.74$)	
Decision tree regression				Absorption coefficient alone: poor performance ($R^2 \leq 0.27$)	
Random forest regression				Combined features improved prediction (best deep neural network: MAPE = 6.31%, $R^2 = 0.54$)	
Deep neural network					
Hierarchical machine learning (HML):	Identify printing parameters that achieve <10% error in linewidth and corner radius relative to CAD (high-fidelity criterion)	Bottom layer: printing parameters include concentration, nozzle diameter, extrusion speed, and print-head speed. Middle layer: physical variables are calculated from predictors.	Printing fidelity (dimension error in linewidth and corner radius)	$R^2 = 0.643$ on the test set (HML with middle layer + LASSO leave-one-out CV)	161
Predictor layer → derived physical variable layer → LASSO regression for feature selection and regularization				Compared to LASSO using only predictors ($R^2 = -0.439$), the middle layer significantly improves the accuracy of the model.	

hydrogel printability (out-of-bag score = 0.96) by isolating the 13 most critical rheological features.¹⁵⁰

Extending ML to crosslinking optimization, support vector regression effectively captures the nonlinear dynamics of GelMA-PEDOT:PSS gelation (MAPE = 3.13%, $R^2 = 0.79$).¹⁵³ Furthermore, hierarchical machine learning (HML) utilizing intermediate physical variables vastly outperforms traditional LASSO ($R^2 = 0.643$ vs. <0).¹⁵⁰ As summarized in Tables 2 and 3,

ML offers a scalable toolkit ranging from basic regression to complex ensembles. Moving forward, synergizing these versatile ML algorithms with DoE will be essential to accelerate bioprinting optimization, reduce costs, and ensure structural reproducibility.^{119,154,155}

To overcome the resource and cost barriers associated with limited experimental data, optimization strategies utilizing machine learning (ML) algorithms tailored for small datasets



are proposed. Unlike deep learning models that necessitate vast datasets, Bayesian optimization (BO) enables the identification of optimal printing parameters within merely 19 to 47 experimental iterations.¹⁴⁷ Furthermore, coupling ML with DoE facilitates the generation of highly representative and structured datasets, thereby mitigating the need for large-scale random data collection.¹⁶³ Notably, the integration of CFD simulations allows for the prediction of material behavior prior to printing, which significantly reduces the requisite number of physical trials while maintaining high accuracy in capturing complex, nonlinear relationships.³⁵

4.3 Applications of CFD in the optimization of hydrogel printability

In the bioprinting field, CFD has become a helpful tool for evaluating key printing parameters, including nozzle velocity, shear stress, printability, and cell viability. Traditionally, optimizing parameters requires multiple experimental iterations, which can significantly increase time and cost to achieve accurate results. Applying CFD simulations can considerably reduce the number of physical experiments required, thereby improving research efficiency.³⁵ For instance, CFD can be used to simulate the material's flow behaviour and deformation prior to testing to save cost, since the bioinks required for printability testing are often expensive. OpenFOAM software has been applied to examine the relationships among hydrogel flow rate, various nozzle designs, and operating pressure.³⁶ Similarly, other CFD platforms such as ANSYS Fluent, COMSOL Multiphysics, and FLOW 3D have been utilized to investigate additional design parameters and nozzle configurations. The application of CFD in 3D bioprinting extends beyond nozzle geometry to simulate how bioink characteristics and nozzle design jointly influence cell viability. For instance, COMSOL was used in one study to compare the effect of shear stress on cell viability between cylindrical and conical nozzles.³⁹

CFD in hydrogel bioprinting is most impactful when directly linked to experimental bioink behaviour and printing outcomes. Modern CFD studies focus on (i) nozzle geometry optimization to control local shear and extensional flows that

determine filament formation and print fidelity; (ii) prediction of shear stresses and residence times that correlate with cell damage or viability; and (iii) process design for high-viscosity dispensing systems (*e.g.*, screw dispensers) and multi-material printheads. These targeted CFD applications have been shown to guide nozzle design parameters (diameter, taper angle, and length), select safe operating windows for extrusion, and reduce the number of empirical trials required for new bioinks.^{39,164,165}

CFD methodologies integrate a preprocessor, a flow solver, and a post-processor phase, each facilitated by various software platforms and computational techniques. The general computational workflow is illustrated in Fig. 5, where fluid problems are first formulated based on fluid mechanics principles, translated into the Navier–Stokes equations, discretized using numerical methods, and then solved computationally on structured or unstructured grids to obtain simulation results that are subsequently validated against experimental data. To illustrate the practical implementation of CFD in hydrogel bioprinting, representative simulation platforms and their typical applications are discussed. These tools, such as IPS IBOFlow, COMSOL Multiphysics 4.2, OpenFOAM, and FLOW 3D, have been used to model bioink rheology, nozzle flow behavior, shear-stress distribution, and droplet formation under realistic printing conditions. When employed alongside discrete numerical methods, CFD enables the in-depth assessment of distinct components within diverse bioprinting technologies.³⁸

In extrusion-based 3D bioprinting, filament morphology plays a critical role, as it constitutes the foundational elements of printed constructs and directly influences resolution, surface characteristics, and the mechanical stability of the final output. Many studies have been conducted using CFD as a useful tool in the optimization of nozzle design, characterization of bioink rheological behaviour, including flow velocity, pressure gradients, and shear stress, and the evaluation of resultant print attributes such as shape fidelity, biocompatibility, and cell survival rates.³⁸ CFD simulations have also been effectively employed to predict and optimize printability. By modeling the extrusion process, CFD enables analysis of how flow rate,

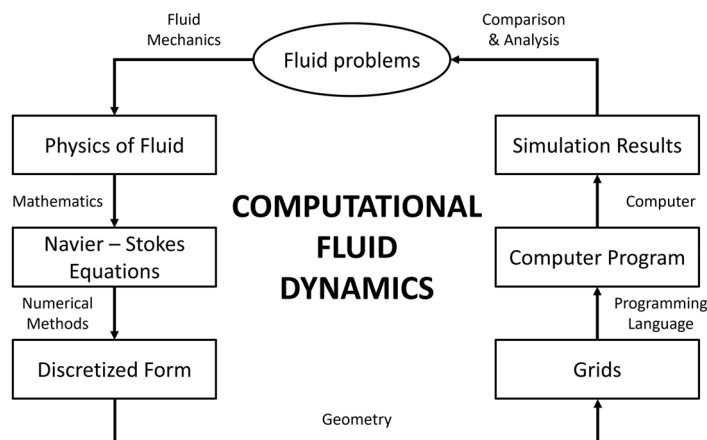


Fig. 5 Computational fluid dynamics process.



nozzle geometry, applied pressure, temperature, and ink rheology influence filament formation and diameter.¹⁶⁴ These simulations, as reviewed by Fareez *et al.* (2024), enable computation of relationships between flow parameters, nozzle design, and printing outcomes, and in several studies CFD predictions have been validated experimentally, supporting the use of CFD to guide optimal settings for enhanced print fidelity.³⁸ A major application of CFD is evaluating bioprinting's impact on cell viability. Wall shear stress within the nozzle is a critical factor influencing cell viability during extrusion-based bioprinting.^{38,166} CFD simulations of different nozzle geometries (cylindrical, conical, and tapered) map shear stress fields and correlate them with experimental viability data, revealing a strong inverse relationship between peak shear stress and post-printing cell survival.³⁵ Conical nozzles produce high but localized shear stress near the tip, while cylindrical nozzles generate lower peak stress over a longer distance, resulting in greater cell damage due to prolonged exposure, insights that guide nozzle redesign to minimize shear while preserving flow.³⁵ CFD further supports microfluidic integration, where organ-on-a-chip platforms enable precise cell–bioink mixing and multi-material dispensing at micro-scales. By simulating velocity, pressure, and shear rate fields in microchannels, CFD optimizes device design for controlled flow and reduced shear.¹⁶⁷ CFD has been applied to simulate and analyze flow characteristics, including pressure distributions, shear stresses, and oxygen concentration profiles, in perfusable microchannels and engineered tissue constructs. In perfusion models with embedded microchannel networks, CFD enables evaluation of flow velocity, oxygen transport, and shear environments, informing design parameters that improve tissue viability and functional perfusion.^{168,169}

Several recent studies exemplify how CFD has been integrated with experiments to address bioprinting-specific questions. Blanco *et al.* review nozzle geometry optimizations that identify favourable internal taper angles and length/diameter ranges to minimize damaging shear while preserving filament continuity; these geometry guidelines have been validated in multiple extrusion experiments.¹⁶⁴ Chand *et al.* and Malekpour *et al.* quantified wall and bulk shear stresses in model bioinks and correlated CFD predicted stress fields with measured cell viability and print quality; these works demonstrate that threshold shear metrics from CFD can be used to screen candidate formulations before wet-lab printing.^{35,39} For high viscosity bioinks, Lee *et al.* combined rheological characterization with CFD of a screw based dispenser to predict extrusion pressures and local shear histories, then validated survival and filament fidelity experimentally, which directly reduced the actual iterations required when developing viscous hydrogel inks.¹⁶⁵ Several recent optimization studies demonstrate that surrogate models trained on combined experimental and CFD features improve prediction of printability and cell viability compared with models trained on experimental inputs alone.^{170,171} Including CFD features therefore enhances model interpretability by linking physical stresses to outcomes and reduces the need for exhaustive experiments.

5. An integrated DoE–ML–CFD strategy

Optimizing hydrogel bioinks for 3D bioprinting involves navigating a multifaceted design space encompassing material composition, rheological properties, printing parameters, and resulting biological performance (*e.g.*, cell viability and structural fidelity).¹⁷² Traditionally, DoE, ML, and CFD have been used separately to investigate these factors. However, integrating them into a unified DoE–ML–CFD framework enables systematic, efficient improvement of hydrogel performance by combining statistical rigor using DoE, physics-based simulation with CFD, and data-driven prediction through ML, thereby accelerating the discovery of robust bioink formulations.^{173,174}

As illustrated in Fig. 6, the refinement cycle requires technically consistent data hand-offs among the DoE, ML, and CFD modules. In the DoE-to-ML transition, experimental outputs such as viscosity, storage or loss moduli, gelation time, extrusion pressure, filament diameter, pore fidelity, and cell viability must be converted into standardized, machine-readable datasets. This step is not trivial because hydrogel datasets are often small, heterogeneous, and sensitive to metadata such as polymer batch, temperature, nozzle diameter, printing pressure, crosslinking method, and measurement shear-rate range. Without consistent units, replicate handling, feature encoding, and uncertainty reporting, ML models may capture experimental artifacts rather than true composition–process–property relationships.

The experimental-to-CFD hand-off is equally important. Rheological data obtained from oscillatory or steady-shear measurements must be translated into constitutive parameters, such as power-law, Herschel–Bulkley, or viscoelastic model coefficients, that can be used as CFD inputs. Errors at this stage can amplify through the entire cycle because an oversimplified viscosity model, an inappropriate shear-rate window, or unrealistic boundary conditions may lead to inaccurate predictions of wall shear stress, pressure drop, residence time, and flow distribution inside the nozzle. Conversely, CFD outputs are usually high-dimensional fields rather than directly usable experimental variables. Therefore, they must be reduced into physically meaningful descriptors, including maximum wall shear stress, average shear exposure, pressure requirement, velocity profile, residence time, and predicted filament expansion, before being integrated into ML models or used to refine the DoE matrix.

In this adaptive cycle, discrepancies among experimental observations, ML predictions, and CFD simulations become decision points for refinement. For example, if ML predicts a formulation to be printable but CFD indicates excessive shear stress that may compromise cell viability, that formulation can be penalized or excluded from the next experimental round. If CFD predicts stable extrusion but the printed filament collapses experimentally, the rheological model, crosslinking kinetics, or boundary conditions must be recalibrated. Similarly, if experimental extrusion pressure or filament morphology deviates from CFD predictions, updated rheological



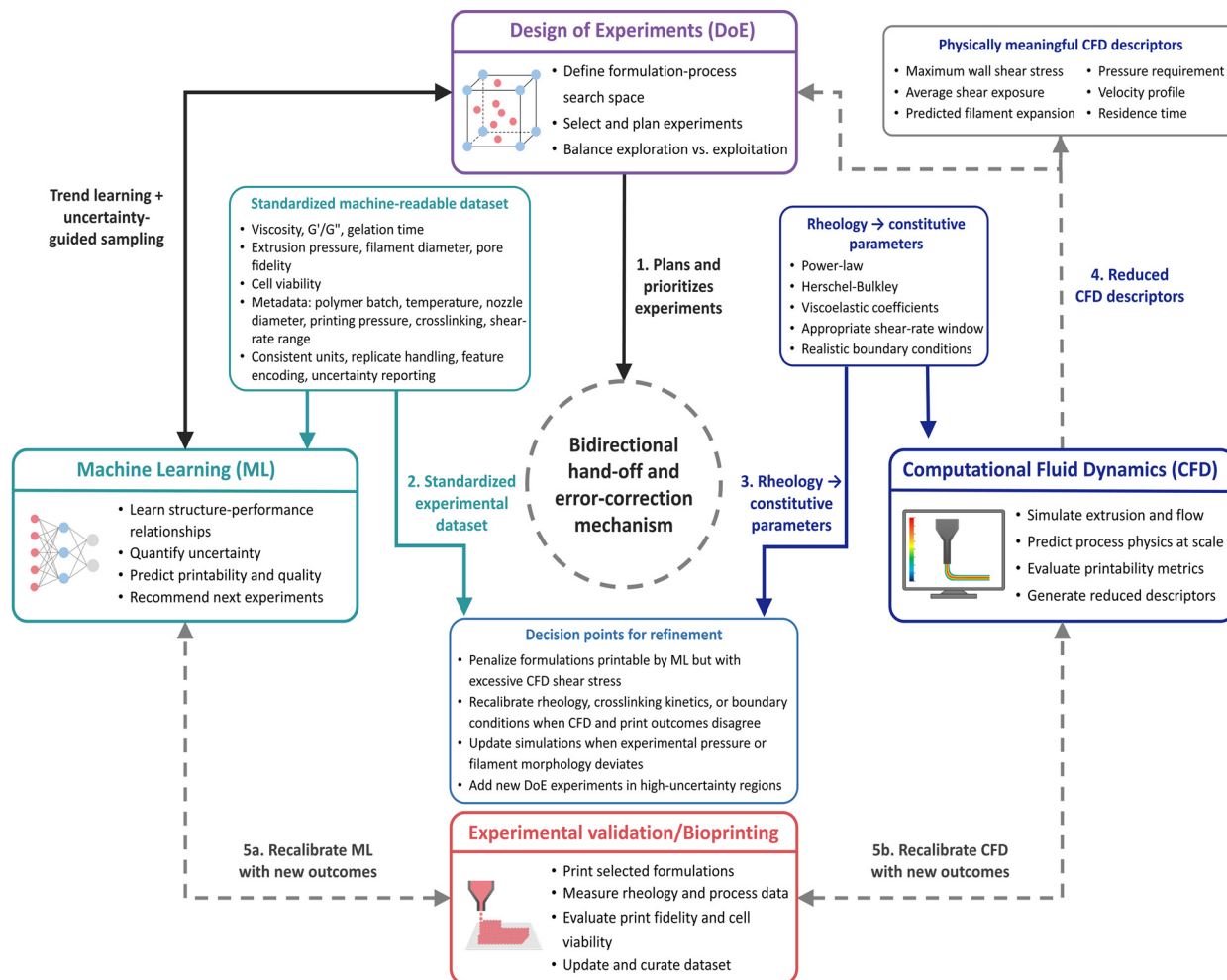


Fig. 6 Integrated DoE-ML-CFD framework for hydrogel bioprinting optimization.

measurements can be fed back into the simulation module. If ML uncertainty is high in a specific formulation region, DoE can prioritize new experiments in that region rather than randomly expanding the dataset. Thus, the dashed arrows in Fig. 6 represent an operational error-correction mechanism: DoE structures the experimental search space, ML identifies nonlinear trends and uncertainty, CFD enforces physical plausibility, and experimental validation recalibrates both models. This bidirectional hand-off transforms the framework into an adaptive optimization cycle that can progressively reduce experimental burden while improving prediction reliability.

Recognizing that multivariable interactions in hydrogel systems are difficult to resolve through conventional trial-and-error approaches, several studies have integrated DoE with ML to construct predictive and statistically structured optimization strategies. For instance, Madadian Bozorg *et al.* employed DoE to vary hydrogel printing parameters and used ML to optimize 3D printing parameters of soft material extrusion, enabling a quality-by-design characterization that systematically explores the effects of key process variables and reduces reliance on traditional trial-and-error approaches.¹⁵⁵ Similarly, Ali

Khalvandi *et al.* employed a Box-Behnken DoE to systematically generate porous PVA/gelatin hydrogel formulations and subsequently trained a supervised deep neural network to predict compressive mechanical responses.¹⁶³ This approach demonstrated that structured experimental design combined with data-driven modeling can capture nonlinear composition-property relationships and reduce reliance on extensive empirical testing. Such integration enables identification of influential formulation parameters (*e.g.*, the role of crosslinker concentration in modulating viscosity and mechanical properties), thereby supporting rational refinement of bioink formulations toward improved functional performance.¹⁷⁵ Additional studies have combined DoE with supervised ML to predict the compressive behavior of PVA gelatin hydrogels and to guide bioink design as well as structural tuning, including scaffold porosity, in 3D bioprinting workflows.^{163,175}

The convergence of ML and CFD is establishing a novel paradigm in 3D bioprinting, facilitating a fundamental shift from empirical, trial-and-error methodologies to quantitative, predictive design strategies.^{38,39} In this synergistic framework, CFD functions as a generator of foundational physical data,



simulating parameters that are experimentally elusive, while ML leverages this dataset to construct high-throughput predictive models for printability and biological functionality.^{176,177} A quintessential example of this approach is the work by Zhang *et al.* who successfully established an integrated framework wherein shear stress data derived from CFD simulations were utilized to train a multi-layer perceptron (MLP). This model facilitated the accurate prediction of as-extruded cell viability without necessitating extensive physical experimentation.¹⁷³ However, it is critical to acknowledge that the efficacy of this hybrid model is intrinsically linked to the fidelity of CFD input assumptions; inaccuracies in characterizing the viscoelastic properties of bioinks can propagate errors throughout the ML prediction chain.^{39,178} Despite these challenges, due to its capacity to significantly abbreviate R&D cycles and optimize resource allocation, the integration of ML and CFD is expected to play an increasingly important role in the fabrication of complex tissue constructs and precision medicine.^{179,180}

DoE and CFD bridge experimental efficiency with physical insight: DoE systematically screens influential parameters such as nozzle geometry and material properties, while CFD simulates flow fields and shear stress distributions within the extrusion channel, enabling mechanistic understanding of factors that affect filament formation and cell viability.^{172,174} This ensures that simulations are guided by and validated against experimental data.¹¹⁹ Sun *et al.* demonstrated a synergistic approach combining rheological characterization with CFD simulations to model hydrogel 3D printing behaviour and relate flow properties to printing performance.¹⁸¹ Chand *et al.* assessed the effects of nozzle geometry and inlet pressure using CFD simulations, quantifying wall shear stress distributions and their potential impact on cell viability.³⁵ These findings provide mechanistic insights that can inform structured experimental design for extrusion optimization. Concurrently, in the domain of hardware optimization, Reina-Romo *et al.* and Rubio *et al.* pioneered “*in silico*” nozzle design strategies combined with DoE, employing CFD to screen nozzle geometries for minimizing shear-induced cell damage.^{164,174} Crucially, CFD does not function as a terminal validation step but as a recalibration mechanism within the loop. Experimental measurements of extrusion pressure or filament morphology can be used to refine rheological inputs and boundary conditions in subsequent simulations. Conversely, CFD-identified stress thresholds can guide the selection of new experimental regions within the DoE matrix. This bidirectional exchange establishes an adaptive cycle in which DoE structures exploration, ML predicts outcomes, and CFD enforces physical plausibility, with each module continuously informing the others rather than operating sequentially.

The selection of optimization strategies in hydrogel-based bioprinting depends on the physicochemical and rheological properties of each material system.^{182,183} Fig. 6 illustrates a flexible workflow integrating DoE, ML, and CFD, in which different methods are applied and combined according to the requirements of the hydrogel system. Natural hydrogels, such as collagen, gelatin, and alginate, are widely reported to be

biocompatible but mechanically weaker and structurally less defined than synthetic systems, resulting in significant variability and nonlinear structure–property relationships.^{182,183} Accordingly, DoE is typically employed to define the formulation–process search space, with key variables including polymer concentration, crosslinker ratio, pH, temperature, and ionic strength, which directly influence viscosity, gelation kinetics, and print fidelity.¹⁸⁴ ML models are then used to capture nonlinear relationships between these variables and performance metrics; for example, elastic modulus and yield stress have been identified as key predictors of printability in ML-assisted bioink design.^{142,143} In contrast, synthetic hydrogels, including PEG- and PVA-based systems, exhibit more controlled and reproducible compositions with tuneable physicochemical properties, enabling more systematic parameterization for optimization.^{142,182} Within this framework, DoE remains useful for exploring multivariable formulation spaces, particularly for variables such as polymer concentration, initiator content, curing conditions, pH, and temperature, which govern swelling behaviour, porosity, and mechanical strength.¹⁸⁴ ML approaches, including regression-based and tree-based models, have been applied to predict mechanical responses and structure–property relationships in polymer and hydrogel systems under nonlinear or photo-crosslinking conditions.¹⁴² In addition, the relatively well-defined rheological behaviour of synthetic hydrogels allows their description using constitutive models, for example, power-law or Herschel–Bulkley, making them particularly suitable for CFD-based analysis of extrusion dynamics.¹⁸¹ In such simulations, key parameters include viscosity models, inlet pressure, nozzle geometry, and shear-rate-dependent behaviour, which together determine shear stress distribution, flow stability, and filament formation during printing.¹⁸¹ For interpenetrating polymer network (IPN) hydrogels, the coexistence of multiple interlaced networks introduces strong coupling between formulation and process variables, leading to highly nonlinear and multiscale behavior.¹⁸⁵ Reviews of IPN systems have shown that while additional networks can enhance mechanical strength and functionality, they also significantly increase the complexity of the design space.^{185,186} Consequently, an integrated DoE–ML–CFD strategy is more appropriate for IPN hydrogels, in which DoE defines the multivariable experimental space, ML captures nonlinear interactions and feature importance, and CFD provides physics-based validation of flow and shear conditions.¹⁷⁵ Following optimization, model predictions and simulation outputs are evaluated against experimental observations.¹⁸¹ If discrepancies arise, the workflow is iteratively redirected to the corresponding module, enabling targeted refinement of the experimental design, retraining of ML models, or recalibration of CFD boundary conditions and constitutive assumptions. This adaptive logic is consistent with recent ML-enabled bioprinting studies that combine experimental datasets with *in silico* predictions to reduce trial-and-error and improve predictive efficiency.¹⁸¹ In this sense, Fig. 6 captures a modular and iterative optimization framework that integrates experimental design, data-driven modeling, and



physics-based simulation into a unified strategy for reproducible hydrogel bioink development.

To make such adaptive cycles reproducible across laboratories, future studies should move toward a standardized bioink metadata schema for FAIR data generation (Fig. 7). At a minimum, this schema should report formulation variables, including polymer type, molecular weight, concentration, degree of substitution, crosslinker or photoinitiator content, pH, ionic strength, pH, and temperature ranges; processing variables, including mixing time, sterilization method, temperature during printing, nozzle diameter, extrusion pressure, printing speed, layer height, and crosslinking dose or duration; and characterization outputs, including viscosity as a function

of shear rate, G' , G'' , $\tan \delta$, yield stress, recovery behaviour, filament diameter, pore fidelity, shape fidelity, and cell viability. For DoE-based studies, the design type, coded and actual factor levels, replicate structure, randomization, response variables, and statistical significance criteria should be explicitly provided. For ML-based studies, dataset size, feature definitions, train/test split, cross-validation strategy, performance metrics such as R^2 , RMSE, MAE, accuracy, $F1$ -score, or AUC, and uncertainty estimation should be reported. For CFD-assisted studies, the constitutive model, fitted rheological parameters, mesh settings, boundary conditions, convergence criteria, and extracted descriptors such as maximum wall shear stress, pressure drop, residence time, and velocity profile



Fig. 7 Proposed bioink metadata schema for FAIR DoE–ML–CFD workflow.



should also be included. Such harmonized reporting would make bioink datasets more findable, accessible, interoperable, and reusable, allowing future DoE–ML–CFD workflows to move from isolated case-specific optimization toward cumulative, transferable, and data-driven bioink design.

6. Conclusion

Hydrogel-based bioinks have emerged as promising materials for 3D bioprinting, offering tuneable rheological behaviour, structural fidelity, and biological functionality. While DoE, ML, and CFD have each independently advanced bioink development, their systematic integration into a unified framework represents a critical step toward rational and predictive biofabrication.

The proposed integrated DoE–ML–CFD strategy establishes a structured and adaptive optimization architecture in which statistically designed experiments generate high-quality datasets, ML models capture nonlinear composition–process–property relationships, and CFD simulations provide physics-based validation of extrusion dynamics and shear stress distributions. Through iterative bidirectional refinement, discrepancies between predictions and experimental observations can be progressively minimized, thereby enhancing model robustness and reducing reliance on empirical trial-and-error approaches.

Despite its potential, several challenges remain, including discrepancies between computational predictions and experimental measurements, limited availability of standardized and high-quality bioprinting datasets, and the inherent complexity of cell–material interactions and crosslinking kinetics in biological environments. Addressing these challenges will require improved rheological characterization, standardized reporting practices, and the development of hybrid physics-informed learning models.

Overall, the integration of DoE, ML, and CFD within a closed-loop framework provides a promising pathway toward reproducible, data-driven, and mechanistically informed bioink design, accelerating the translation of 3D bioprinting technologies from experimental research to clinical and industrial applications.

Author contributions

The concepts were proposed by Minh Hien Nguyen and Thi Tan Pham. Minh Hien Nguyen, Gia Huy Duong, Le Thao Vy Huynh, Hoang Cac Tien Le, Thi Yen Nhi Nguyen, Vinh-Dat Vuong and Thi Tan Pham contributed to collecting, summarizing, and evaluating research and news. All authors contributed to drafting and revising the final manuscript.

Conflicts of interest

There are no conflicts to declare.

Data availability

No primary research results, software, or code have been included, and no new data were generated or analysed as part of this review. All information discussed in this article is available in the cited literature.

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