Biodegradable Inks in Indirect Three-Dimensional Bioprinting for Tissue Vascularization

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Mature vasculature is important for the survival of bioengineered tissue constructs, both in vivo and in vitro; however, the fabrication of fully vascularized tissue constructs remains a great challenge in tissue engineering. Indirect three-dimensional (3D) bioprinting refers to a 3D printing technique that can rapidly fabricate scaffolds with controllable internal pores, cavities, and channels through the use of sacrificial molds. It has attracted much attention in recent years owing to its ability to create complex vascular network-like channels through thick tissue constructs while maintaining endothelial cell activity. Biodegradable materials play a crucial role in tissue engineering. Scaffolds made of biodegradable materials act as temporary templates, interact with cells, integrate with native tissues, and affect the results of tissue remodeling. Biodegradable ink selection, especially the choice of scaffold and sacrificial materials in indirect 3D bioprinting, has been the focus of several recent studies. The major objective of this review is to summarize the basic characteristics of biodegradable materials commonly used in indirect 3D bioprinting for vascularization, and to address recent advances in applying this technique to the vascularization of different tissues. Furthermore, the review describes how indirect 3D bioprinting creates blood vessels and vascularized tissue constructs by introducing the methodology and biodegradable ink selection. With the continuous improvement of biodegradable materials in the future, indirect 3D bioprinting will make further contributions to the development of this field.

Keywords: biodegradable ink, indirect 3D bioprinting, vascularization, scaffold, tissue engineering

INTRODUCTION

At present, only a few tissue-engineered products, such as skin (Ryssel et al., 2008) and cartilage (Makris et al., 2015), have achieved clinical success. For organs and tissues with more complex structure, such as the heart, liver, or spleen, there is still a long way to go (Berthiaume et al., 2011; Datta et al., 2017). An immature vascular system is one of the most important reasons for the failure of these products (Duan, 2017; Richards et al., 2017). In recent years, the rise of 3D bioprinting technology has enabled outstanding contributions to be made towards solving the problem of vasculature fabrication, thus extending the potential application of artificial tissues (Zhu et al., 2016; Alonzo et al., 2019; Hann et al., 2019). Indirect 3D bioprinting increasingly attracts research attention because of its superior capabilities with respect to vascularization (Wang Z. et al., 2019). Biodegradable materials are a crucial part of tissue engineering. They are typically designed to promote new tissue generation by serving as temporary templates and...
INDIRECT 3D BIOPRINTING FOR TISSUE VASCULARIZATION

Vascularization Challenges in 3D Bioprinting

Vascularization is an essential part of the human body. Mature vasculature provides continuous perfusion, transporting nutrients to and removing metabolic wastes from cells, thus maintaining high cell viability and normal tissue function (Leong et al., 2013; Kim et al., 2016; Hann et al., 2019). For nonvascular tissues, the diffusion range of oxygen and nutrients is generally 100–200 μm (Carmeliet and Jain, 2000; Kaully et al., 2010). This means that, for successful in vivo implantation of engineered tissue constructs any larger in scale than this limit, it is necessary to ensure sufficient vasculature throughout the construct and good integration with the host vascular system (Ko et al., 2007). However, the vascularization of thick tissues has always been a major challenge and a research hotspot in tissue engineering.

3D bioprinting technology, also referred to as additive manufacturing, uses special 3D printers and bioinks containing cellular/bioactive components to fabricate scaffolds that imitate living tissues, in a layer-by-layer deposition approach with the help of computer-aided design (CAD) (Datta et al., 2017; Abdollahi et al., 2019). Compared with other tissue fabrication methods, 3D bioprinting stands out because of its convenience of customization, precise multi-dimensional control, and ability to fabricate 3D biostructures with suitable mechanical properties. In particular, its accurate control of complex and delicate structures within constructs, combined with the adoption of bioinks, make 3D bioprinting a powerful and efficient tool to address the problem of vascularization by creating a vasculogenetic (Laschke and Menger, 2012; Balaji et al., 2013) (generating new blood vessels from endothelial cells or vascular progenitor cells) and angiogenic vascularization by creating a vasculogenic (Laschke and Menger, 2016). It relies on seeding or encapsulating endothelial cells (ECs) in the construct in vitro. However, the process of cell infiltration and growth in vitro is uncertain, and the resultant vasculature is usually inhomogeneous (Leong et al., 2013). When implanted, immature vasculature results in insufficient blood perfusion and induces core necrosis in constructs (Asakawa et al., 2010), causing the failure of thick constructs to survive (Rouwkema et al., 2008).

Nowadays, many 3D bioprinting methods have been developed to mimic vasculature, including extrusion-, laser-based systems, electrospinning, stacking of micropatterns or modules, and cell sheet techniques (Yamamura et al., 2007; Kolesky et al., 2014; Sarker M. et al., 2018; Sarker M. D. et al., 2018). New methods are constantly being explored to solve the abovementioned problems, and indirect 3D bioprinting may cast light on vascularization in tissue engineering.

Indirect 3D Bioprinting to Address the Problem of Tissue Vascularization

Instead of directly simulating and manufacturing the target constructs, indirect 3D bioprinting fabricates a sacrificial mold providing physical or chemical signals for cells (Asghari et al., 2017). Biodegradable materials also have interesting applications and new challenges in indirect 3D bioprinting for vascularization. In this review, we describe the printing methods, selection of biodegradable inks, and applications of indirect 3D bioprinting for blood vessels and vascularized tissue constructs; furthermore, we point out the existing challenges and trends for future development.
or template (negative model) and fills it with a second material. After selective dissolution, the sacrificial mold or template is removed while the construct (patrix) formed by the second material is preserved (Lee et al., 2013; Van Hoorick et al., 2015). Through this approach, pores, cavities, and channels can be precisely made throughout thick constructs (Lee et al., 2008; Schumacher et al., 2010). Owing to its specific manner of printing, other names for indirect 3D printing include lost-mold technique, indirect rapid prototyping, indirect additive manufacturing, and indirect solid free-form fabrication (Chu et al., 2001; Schumacher et al., 2010; Van Hoorick et al., 2015; Houben et al., 2016; Houben et al., 2017).

Currently, indirect 3D bioprinting has made progress in both hard and soft tissue engineering systems. Some specific tissue constructs such as functional nerve guide conduits, human knee meniscal scaffolds, sized vascular grafts, can be successfully produced in laboratories as proof of concept (Schöneberg et al., 2018; Huang et al., 2021; P et al., 2022). At the same time, with the assistance of sacrificial molds, materials that are traditionally difficult to print can be utilized to manufacture fine...

**FIGURE 1 |** Schematic illustration showing the indirect 3D bioprinting process for blood vessels and vascularized tissue constructs. (A) Three major techniques used in indirect 3D bioprinting for both sacrificial mold and patrix fabrication, including extrusion-based printing, inkjet-based printing, and DLP printing. (B) Process of blood vessel fabrication. (C) Process of vascularized tissue construct fabrication. Sacrificial mold and patrix fabrication can be further divided into three methods according to the sequence of fabrication in step 1. (D) Applications in vitro or in vivo. Blood vessel grafts constructed by indirect 3D bioprinting are currently used for studies in vitro, while vascularized tissue constructs are used for both studies in vitro and animal experiments in vivo.
structures. For example, ceramic materials play a vital role in bone tissue engineering, but its additive manufacturing is considered challenging. Chawich et al. creatively utilized a sacrificial honeycomb mold of polyactic acid (PLA), and they ultimately produced customizable stable Si-based 3D non-oxide cellular ceramic structures (El Chawich et al., 2022). Another team produced stiffness memory nanohybrid scaffolds from poly(urea-urethane) (PUU) solution which exhibits outstanding performance in long-term implantable cardiovascular devices but can not be printed directly (Wu et al., 2000). Of all the breakthroughs, the greatest concern is the potential for this technique to address problems where vascularization is a key issue.

Generally, indirect 3D bioprinting for blood vessels and vascularized tissue constructs includes two general aspects: channel fabrication and vascularization. Channel fabrication can be further divided into three steps: sacrificial mold fabrication, patrix fabrication, and sacrificial mold removal. Manufacture of the negative mold is designed to simulate the shape and extension direction of the vascular network in a solid columnar form. In patrix fabrication, the patrix is built around the sacrificial mold to form the final scaffold or construct. Afterwards, special steps are performed to remove the sacrificial mold, leaving isometric hollow channels inside the scaffold or construct. The detailed production process and precautions of each step have been described in excellent reviews (Lee and Yeong, 2016; Houben et al., 2017; Zhu et al., 2021). Figure 1 provides a brief overview of the printing process. Here we mainly introduce the techniques frequently used and bioink selection.

The three main techniques are extrusion-based, inkjet-based, and light-assisted 3D printing (Vijayavenkataraman et al., 2018). Each technique has advantages and disadvantages, and raise different requirements for bioinks. In extrusion-based printing, the most common technique, 3D structures are built layer by layer using a nozzle to dispense continuous filament (Stumberger and Vihar, 2018; Yang et al., 2020). This technique can achieve resolutions of >100 μm, uniform cell distribution, and moderate printing speed (Placone and Engler, 2018). The technique is compatible with various materials with a viscosity range of 30 mPa s to 6 × 10^7 mPa s (Murphy and Atala, 2014). Furthermore, its cell-friendly printing environment allows it to be used for printing scaffolds with high cell density (Skadal et al., 2015). One promising method is coaxial extrusion, also referred to as coaxial extrusion. Its key features are the concentric and multi-layered nozzles that enable concentric multi-material deposition. With the use of sacrificial materials, this technique allows synchronous manufacturing of the sacrificial mold and scaffold, simplifying the production process (Kjar et al., 2021). However, for extrusion-based printing, the shear stress produced during printing is an important cause of cell death and, because the nozzle size is inversely proportional to extrusion pressure, small nozzles used to improve resolution may have a greater negative impact on cell viability (Kyle et al., 2017).

Inkjet-based printing is defined as dispensing bioink through small orifices via piezoelectric, thermal, or electrostatic actuators, and accurately positioning a very small amount of the bioink on a substrate (Calvert, 2007; Datta et al., 2017). This technique can reach high resolutions of 10–100 μm with short printing times (Poellmann et al., 2011; Gudapati et al., 2016). Different inkjet printing techniques require different material viscosities. For thermal, piezoelectric, and electrostatic inkjet printing the ink viscosity should be relatively low, because the nozzle diameter is usually small and is prone to clogging (Li et al., 2020). However, for electrohydrodynamic jet bioprinting which produces droplets through an electric field, bioink with a high viscosity of over 2000 mPa s can be printed (Workman et al., 2014). Nozzle clogging, small nozzle aperture indirectly leading to cell viability damage, and low-viscosity bioink resulting in poor shape fidelity present challenges to broadening the range of implementation (Chahal et al., 2012; Malda et al., 2013; Zhu et al., 2017).

Light-assisted printing includes digital light processing (DLP)-based and laser-based printing. DLP printing is realized by dynamically projecting an entire computer-generated optical mask into a photosensitive prepolymer solution to induce photopolymerization, while laser-based printing uses a bottom-up approach (drop-by-drop) to build constructs (Zhu et al., 2021). DLP printing is now attracting greater attention as this technique provides precise control of scaffold structures and features and high printing speed (Mandrycky et al., 2016). Nevertheless, the selection of biomaterials is limited to photosensitive polymers and additional chemical modifications are required for most biomaterials (Yue et al., 2015). Moreover, the properties of printing complex patterns directly make DLP more frequently used in direct printing. Therefore, light-assisted printing has only been used in a few cases of indirect 3D printing (Thomas et al., 2020).

Another technique called embedded extrusion bioprinting involves printing a cell-containing hydrogel in a supporting bath, which serves as a sacrificial printing environment (Rocca et al., 2018). As atypical indirect 3D bioprinting, this technique does not impose any requirements for the shape of the sacrificial mold but instead skips this step and directly prints the male mold. The advantage is that low-viscosity hydrogels can be used to construct complex draping and hollow structures with the support of the sacrificial supporting bath via free-form bioprinting, which is difficult to achieve in air. Limitations are that shape fidelity of the complex pore structure is low, resolution is currently at the millimeter level, and the removal of the sacrificial supporting bath is relatively cumbersome (Afghah et al., 2020).

Table 1 summarizes the characteristics of the common techniques covered in this review.

**Vascularization Strategies**

Vascularization is essential for engineered tissue grafts to achieve biological function and can be realized by seeding ECs into hollow scaffold channels (Bae et al., 2012; Massa et al., 2017). After migration, survival, proliferation, and differentiation of ECs, a monolayer of cells is ultimately formed on the surface of the channels, mimicking the physiological vessel wall. Usually, growth factors and other bioactive factors are perfused along with ECs to regulate cell behavior or promote cell differentiation.
The generation of new blood vessels involves complex effects of a variety of cells, factors released by platelets, extracellular matrix, and angiogenic and anti-angiogenic factors. Vascular endothelial growth factor (VEGF) and fibroblast growth factor b (FGFb) are the most effective angiogenic growth factors and are commonly used in the making of angiogenic biomaterials (Sun et al., 2011; Matsui and Tabata, 2012). Research reports successful seeding of human umbilical vein endothelial cells (HUVECs) into microchannels after mold sacrifice, and a uniform HUVEC layer formation coating the channels. It was demonstrated that the HUVEC layer could, to some extent, act as a barrier to protect scaffold cells from harmful substances (Massa et al., 2017). However, a potential problem is that the channel diameter may affect cell inoculation. Narrow channels will become blocked by ECs, hindering perfusion, while diameters too large will require more time for cells to cover the channel surface. Studies have shown that ideal channel diameter ranges from 280 to 1,270 μm (Zhang et al., 2016).

Another elegant vascularization strategy of indirect 3D bioprinting is to encapsulate ECs in the sacrificial ink to realize synchronous distribution of ECs in the process of blood vessel (sacrificial channel) fabrication. This approach reduces the manufacturing process and can realize precise control of cell distribution (Thomas et al., 2020). Owing to direct contact with cells and bioactive factors, biocompatibility and biodegradability of materials are particularly important. Studies have demonstrated that cells show increased proliferation and proliferation in hydrogels with faster degradation rates (Patterson and Hubbell, 2010). A further study suggests that biodegradable materials can promote angiogenesis through temporarily controlled delivery of siRNAs (Nelson et al., 2014). Application of biodegradable materials may better promote vascularization.

**BIODEGRADABLE INK SELECTION**

Biodegradable materials are the main scaffold materials in the field of tissue engineering, which have a broad range of applications in medicine and pharmacy because of good biosafety, reducing inflammation, ability to mimic the extracellular matrix (ECM), and enzymatic degradation in vivo (Asghari et al., 2017). Eventually these scaffolds are replaced by host tissues (Silva et al., 2020). Biodegradable materials are generally classified into synthetic materials and natural materials. Synthetic materials such as polyglycolic acid (PGA), polycaprolactone (PCL), PLA, poly(N-isopropylacrylamide), and their copolymers have already been widely used in tissue engineering. Advantages are that the key properties of these materials as scaffolds can be artificially controlled, such as degradation rate and some specific mechanical properties, including stiffness and elasticity; however, their biosafety is comparatively poor. Natural materials, such as chitin, alginate, collagen, and gelatin, are polymers produced by biological systems. Natural materials have many advantages and disadvantages. The most prominent advantage is their similarity to host tissues, including the ability to communicate with biological systems, enzymatic degradation, metabolic compatibility, and low inflammatory response (Asghari et al., 2017). However, owing to the poor strength of natural materials, scaffolds often collapse before finishing their task. In addition, they generally show slow and inhomogeneous degradation in vivo, and are inconsistent with host tissue regeneration rate.

In the field of indirect 3D printing, biodegradable polymers play vital roles in fabricating vascular or vascularized tissue: 1) As scaffold materials, they allow endothelial cells and a variety of angiogenic factors to exist, as well as providing a flexible
environment for complex angiogenesis. 2) As sacrificial materials, compared with non-degradable ones, the damage to cell activity when manufacturing or removing the sacrificial molds is much lower. 3) Since the biodegradable materials can degrade in vivo, the use of them as sacrificial material helps to skip the processing step of sacrificial mold removal in vitro, thus shortening the manufacturing time. 4) In theory, by regulating and matching the degradation rate of scaffolds and sacrificial materials, the removal of sacrificial channels and channel endothelialization can be completed in vivo before the scaffolds completely degrade, thus realizing the direct in vivo application of constructs.

In the following section, we mainly enumerate the commonly used biodegradable polymers in indirect 3D bioprinting for blood vessels and vascularized tissue constructs. Table 2 summarizes the characteristics, combinations, and current applications of different scaffold and sacrificial materials.

### Scaffold Materials

Scaffold materials, whose purpose is to simulate the ECM and provide support for tissue regeneration in the human body, should have good biosafety, biocompatibility, biodegradability, and mechanical properties that are favorable for cell growth, proliferation, and migration (Lee A. Y. et al., 2014). Ideally, they

| Biomaterials | Scaffold or sacrificial material | Natural or synthetic | Biodegradable or non-degradable | Biocompatibility | Mechanical property | Combinations | Applications | References |
|--------------|---------------------------------|---------------------|-------------------------------|-----------------|-------------------|--------------|-------------|-----------|
| Gelatin      | Scaffold and sacrificial material | Natural             | Biodegradable                 | Good            | Poor              | PVA, Pluronic, agarose, HA, xanthan-gum | Earlobe-shaped channel system, liver model | Liu et al. (2008) Bentzon et al. (2009) Nichol et al. (2010) O’Bryan et al. (2017) Steunenberg and Vihar. (2018) Hu et al. (2018) Schöneberg et al. (2018) Duarte Campos et al. (2020) |
| Fibrin       | Scaffold material                | Natural             | Biodegradable                 | Good            | Poor              | Gelatin, carbohydrate glass             | Arteriole/venule                              | Lee and Mooney, (2012) Rocca et al. (2018) Piras and Smith, (2020) Liu et al. (2013) Bidgoli et al. (2019) |
| Alginate     | Scaffold and sacrificial material | Natural             | Biodegradable                 | Good            | Medium            | PVA, agarose, Pluronic F127, carbohydrate glass | Human heart- and kidney-like objects | Tocchio et al. (2015) Hernández-Córdoba et al. (2016) Hu et al. (2018) Kim et al. (2018) Charron et al. (2019) Wu et al. (2019) Thomas et al. (2020) |
| SF           | Scaffold material                | Natural             | Biodegradable                 | Good            |                  | Thermoplastic, plaster                 | Bone and cartilage engineering              | Liu et al. (2013) Bidgoli et al. (2019) |
| PVA          | Sacrificial material             | Synthetic           | Biodegradable                 | Good            |                  | Gelatin, silk, agarose, alginates, fibrin, Matrigel, PLCL, PUU | 100–1750 μm diameter channels | Tocchio et al. (2015) Hernández-Córdoba et al. (2016) Hu et al. (2018) Im et al. (2018) Kim et al. (2018) Charron et al. (2019) Wu et al. (2019) Thomas et al. (2020) |
| HA           | Sacrificial material             | Synthetic           | Biodegradable                 | Good            | Poor              | Gelatin                               | Enzymatically digestible, 380–720 μm diameter channels | Bertassoni et al. (2014) Massa et al. (2017) López-Marcial et al. (2018) Khattak et al. (2005) Miller et al. (2012) Homan et al. (2016) Daly et al. (2018) Ding and Chang, (2018) Hu et al. (2018) Xu et al. (2018) Yang et al. (2020) |
| Agarose fiber | Sacrificial material             | Natural             | Biodegradable                 | Good            |                  | Alginates, Gelatin                    | 100–1,000 μm diameter channels               | Bertassoni et al. (2014) Massa et al. (2017) López-Marcial et al. (2018) Khattak et al. (2005) Miller et al. (2012) Homan et al. (2016) Daly et al. (2018) Ding and Chang, (2018) Hu et al. (2018) Xu et al. (2018) Yang et al. (2020) |
| Carbohydrate glass | Sacrificial material            | Synthetic           | Non-degradable               | Cytotoxic when dissolved | Good | PEG, fibrin, alginate, agarose, Gelatin, sodium alginate, decellularized extracellular matrix | 150–750 μm diameter channels | Bertassoni et al. (2014) Massa et al. (2017) López-Marcial et al. (2018) Khattak et al. (2005) Miller et al. (2012) Homan et al. (2016) Daly et al. (2018) Ding and Chang, (2018) Hu et al. (2018) Xu et al. (2018) Yang et al. (2020) |
should also be able to promote angiogenesis (Do et al., 2015; Xu et al., 2018). After crosslinking, scaffold materials need a certain degree of stiffness to maintain structural integrity during removal of sacrificial materials, and to support the flow of perfusion (Skylar-Scott et al., 2019). Controllable degradation rate consistent with the growth and repair rate of host tissue in vivo is also necessary (Malda et al., 2013; Serbo and Gerecht, 2013).

To date, various natural biodegradable polymers have been used for scaffold bioprinting, such as gelatin (Chen et al., 2012), fibrin (Schöneberg et al., 2018), and alginate (Jia et al., 2014); they are collectively referred to as hydrogels. They show excellent human ECM features and allow cell encapsulation (Malda et al., 2013), but they have all been deficient in some respect. Uncontrolled degradation and poor mechanical properties are the main problems, and they are also the research focus of material modification. The simplest and most used method is blending modification. In this approach, the hydrogel ratio is controlled to prevent excessive polymer concentration or dilution, which may have an adverse effect on cell behavior or mechanical properties (Malda et al., 2013; Duarte Campos et al., 2016; Zou et al., 2020).

Gelatin is one of the most widely used scaffold materials extracted from collagen, the main component of natural human ECM. Gelatin is composed of 85–92% protein, water, and mineral salts, and is highly susceptible to several proteases (Bello et al., 2020). Its excellent biocompatibility and similarity with collagen have made it the preferred material for the assembly of scaffolds. However, as a biomaterial ink, gelatin does have several drawbacks; these include low viscosity and yield stress, as well as relatively long crosslinking time, which leads to poor shape retention and structural collapse, and is the main obstacle to creating high resolution 3D pore or microchannel structures (O’Bryan et al., 2017). These disadvantages can be improved by making gelatin composites, or applying sacrificial materials to support the scaffold hydrogel before crosslinking (Moroni et al., 2018). In addition, the degradation rate of large solid gelatin in vivo is relatively slow (Daly et al., 2018). Large numbers of gelatin residues within thick tissues remain a challenge. Solutions may include lowering the degree of metacrylation and the macromer concentration (Nichol et al., 2016; Chen et al., 2012). Furthermore, by creating microchannels within the hydrogels to enhance host interaction, the degradation of gelatin can be enhanced (Daly et al., 2018). This is possibly because of the increased invasion by host immune cells, such as macrophages, which is vital for degrading gelatin hydrogels (Kim et al., 2014).

Gelatin-methacryloyl (GelMA) is the most commonly used material among gelatin derivatives and composites. GelMA is a photopolymerizable hydrogel made of gelatin derivatized with methacrylamide side groups (Van Den Bulcke et al., 2000). It has high biosafety compared with several gelatin-based hydrogels formed by chemical crosslinking methods, such as ones fabricated using glutaraldehyde or transglutimase derived from bacteria (Benton et al., 2009). GelMA is sensitive to matrix metalloproteinases and can be degraded by cells (Nichol et al., 2010). By adjusting the rate of polymerization and ratio of methacrylic acid, GelMA can maintain relatively good shape with adjustable mechanical properties (Benton et al., 2009). Research has shown that ECs as well as endothelial colony-forming cells can undergo active angiogenesis or vasculogenesis in GelMA hydrogels either in vivo or in vitro (Chen et al., 2012; Massa et al., 2017).

Fibrin scaffolds have a wide range of applications in tissue engineering, especially in bone tissue engineering. Fibrin is a natural biopolymer produced by thrombin cleavage of fibrinogen, and serves as a temporary scaffold for tissue healing in physiologic processes (Noori et al., 2017). It also plays an important role in specific receptor-mediated interactions with cells because of its ability to bind different types of proteins and growth factors, including FGF and VEGF that promote angiogenesis (Breen et al., 2009; Litvinov and Weisel, 2017). Fibrin hydrogel can be easily remodeled by ECs, which is favorable for fast angiogenesis. Nevertheless, similar to gelatin, fibrin rapidly degrades, and has poor mechanical stability, durability, and shape fidelity (Calderon et al., 2017). To overcome these problems, fibrin composites and mimics have been developed in 3D bioprinting. For example, by combining fibrin and gelatin, the stiffness of the hydrogel increased, and lower water loss on compression was observed (Schöneberg et al., 2018). Another attempt at a fibrin composite is called ELP-RGD, composed of ELP (elastin-like protein) hydrogel along with a cell adhesion RGD amino acid sequence derived from fibrinectin. ELP hydrogel contains elastin (a kind of fibrin) -like repeat units alternating with biologically active domains (Madl et al., 2017). As a scaffold material, ELP-RGD has adjustable stiffness, is readily hydrolyzable with protease, and is able to promote matrix remodeling as well as cell proliferation (Chung et al., 2012). It has been demonstrated to be suitable for on-chip platforms with vascular-like networks (Duarte Campos et al., 2020).

Alginate is a natural polysaccharide extracted from alginic acid. The long polysaccharide chains provide it with pliability and gelling adeptness, and it undergoes hydrolytic cleavage under acidic conditions or enzymatic degradation by lyase (Pawar and Edgar, 2012; Rastogi and Kandasubramanian, 2019). Despite its mechanical instability and poor cell attachment, alginate is widely used as a hydrogel because of its low cost, good biosafety, and its ability to be rapidly but reversibly crosslinked by Ca²⁺ under mild conditions (Jia et al., 2016; Rastogi and Kandasubramanian, 2019). However, one important limitation of alginate application in vivo is the low degradation rate and unpredictable degradation process owing to the lack of alginate degrading enzyme in the human body (Reakasame and Boccaccini, 2018). Moreover, alginate shows relatively poor cell adhesion and infiltration (Balakrishnan et al., 2014). Measures like modification or mixing in additives, such as nanomaterials, peptides, and growth factors, are useful to regulate rheological properties, promote cell adhesion, or guide cell differentiation (Lee and Mooney, 2012; Piras and Smith, 2020). For example, by oxidative modification, more reactive sites are provided to the structure, accelerating the alginate’s biodegradability (Liang et al., 2011). Ino et al. combined sodium alginate with sacrificial molds of sugar structures, and by soaking the structure in a CaCl₂ solution they achieved
simultaneous dissolution of the mold and formation of calcium alginate hydrogel, thus simplifying and hastening the manufacturing process (Ino et al., 2020).

Silk fibroin (SF), a silk-derived protein-based material approved by the FDA (Perrone et al., 2014), has been used to make clinical sutures for many years. In recent decades, because of new processing techniques and further understanding of its properties, SF has attracted great interest in bone and cartilage engineering (Liu et al., 2013). Compared with the abovementioned materials, native silk fibers have excellent mechanical properties including good strength and toughness (Kundu et al., 2013; M et al., 2017). Other advantages of SF include good biosafety, good biocompatibility, controllable biodegradability and bone induction, and low immunogenicity (Wenk et al., 2009; Kundu et al., 2013). The host immune system mediates the process, suggesting that SF is also biodegradable (Wang et al., 2008). It is worth noting that SF remains strong during degradation, which is its unique advantage in tissue engineering (Kundu et al., 2013). Studies were carried out to produce vascularized scaffolds via SF. In combination with indirect 3D bioprinting, a layered SF-bioactive glass composite scaffold with excellent compressive strength, flexibility, and 10–50 μm micropores has been fabricated (Bidgoli et al., 2019). The scaffold comprises hierarchically micro and sub-micro pores, which are important features for promoting cell migration, differentiation, bone formation, and angiogenesis (Qi et al., 2018); results showed that the scaffold enhanced cell adhesion and cell proliferation (Bidgoli et al., 2019). However, drawbacks of SF such as lack of biological activity, general poor performance under humid conditions, and the difficulty of transportation and long-term storage may limit its further application. These limitations are potential future research directions.

Other scaffold materials commonly used in indirect 3D bioprinting but that are not yet, or rarely, used for vasculature fabrication include polyethylene glycol (PEG), hyaluronic acid (HA), PLA, PCL, poly (ε-caprolactone) (PLCL), and poly(lactic-co-glycolic) acid (PLGA) (Park et al., 2014; Houben et al., 2016; Alijohani et al., 2018; Im et al., 2018). Inorganic substances are more frequently used for osteochondral tissue fabrication because of their excellent mechanical properties. Future studies on these materials can be performed in the field of indirect 3D bioprinting for tissue vascularization.

**Sacrificial Materials**

Ideal sacrificial materials should have good fluidity for free molding, rapid solidification to save printing time, a low expansion rate, and appropriate mechanical strength to achieve good shape fidelity. During the process of printing, there should be no adverse reactions with the scaffold material that result in deformation of the scaffold structure. If working in combination with scaffold materials containing cells or other bioactive factors, sacrificial materials should be chosen to ensure non-toxic and non-stimulatory conditions. Also, the conditions for their state transformation and removal should be mild and easy to achieve, preserving the shape and properties of the scaffold. When exposed to living cells *in vivo* or bioactive components, cytotoxicity, biocompatibility, and whether the removal conditions are conducive are usually considered first.

Currently the commonly used sacrificial materials that come closest to meeting the conditions described earlier are carbohydrate glass (Miller et al., 2012), Pluronics (Afghah et al., 2020), and PVA (Zou et al., 2020). Carbohydrate glass and Pluronics are nonbiodegradable materials. Carbohydrate glass is a simple glass consisting of a mixture of carbohydrates, including glucose, sucrose, and dextran, and was one of the first materials applied to indirect 3D bioprinting as a sacrificial biomaterial ink (Miller et al., 2012). The synthetic glass shows sufficient mechanical stiffness to maintain its shape in air, as well as rapid dissolution to accelerate the process. Results demonstrated that it can be compatible with a variety of natural or synthetic hydrogel materials, such as agarose, alginate, fibrin, and Matrigel, adapting well to their different properties and means of crosslinking (Miller et al., 2012). Currently, vessels with diameters ranging from 150 μm to 1 mm and smooth in-plane junctions can be achieved with this sacrificial ink (Pollet et al., 2019). Pluronics are a class of amphiphilic tri-block copolymers popular in drug and clinical applications. Because they have the characteristics of solubilizer, emulsifier, and stabilizer, they are often used as excipients in pharmaceutical preparations (Jarak et al., 2020). The most frequently used Pluronic in tissue engineering is Pluronic F127, which can rapidly dissolve in aqueous media or biological fluids. Its sol-gel transition at room temperature and convenient removal attracts attention as a sacrificial ink. Channels with a diameter as small as 150 μm can now be printed with Pluronic F127 (Homan et al., 2016). However, both carbohydrate glass and Pluronic F127 show cytotoxicity when dissolved, which is one of the most prominent shortcomings (Miller et al., 2012; Deng et al., 2020). Besides, Pluronic F127 liquefies at low temperatures, making it difficult to use with some scaffold materials that require these temperatures during casting, such as collagen and matrix gelatin (Ding and Chang, 2018; Hu et al., 2018).

As mentioned above, nonbiodegradable materials usually exhibit certain cytotoxicity and their removal is relatively cumbersome. As a result, researchers have investigated biodegradable materials, which show higher biosafety and are expected to achieve direct application in vivo.

PVA, a commonly used biodegradable sacrificial material, has satisfactory biocompatibility and similar functions to natural tissues, including high water content, high elasticity, and low interfacial tension with biological fluids (Teodorescu et al., 2019). It shows resistance to protein absorption, which is important for bone formation (Kim et al., 2018). The ease of printability allows it to be used for repeatable fabrication of complex vascular patterns (Hu et al., 2018). In terms of mechanical properties, PVA has good strength and stiffness at 25°C (Charron et al., 2019) and fits well with different biodegradable natural polymers, such as gelatin and silk (Mohanty et al., 2016). Tocchio et al. used PVA successfully to make sacrificial templates with characteristic sizes of 100–500 μm in multi-branch structures, which helped to further simulate the complex environment of cell growth, and
showed great potential for production of large-sized vascularized scaffolds that would meet clinical needs (Tocchio et al., 2015). Compared to other sacrificial materials, the stable chemical properties, convenience of preservation, and low cost of PVA are huge advantages for its use in industrial production. However, owing to the lack of bioactive components, PVA tends to be resistant to protein absorption and cell adhesion, which limits its further application outside bone tissue engineering (Schmedlen et al., 2002). Furthermore, if PVA filaments are too large in diameter (>500 μm reported), they are likely to deform as they cannot support their own weight (Hernández-Córdova et al., 2016). More in-depth research is needed in the future.

HA-based enzymatically degradable photoink was developed by different research teams (Zhu et al., 2017; Thomas et al., 2020). HA is a linear polysaccharide. It is an essential component of the ECM and has vital effects on many cellular responses, including cellular signaling, wound repair, morphogenesis, and angiogenesis (Burdick and Prestwich, 2011). In the abovementioned two experiments, HA was chemically modified to achieve photopolymerization and mixed with gelatin to form the photoink. The hydrogels could be digested with hyaluronidase to achieve fast (within hours) and collaborative (not limited by graft size) degradation. Channels ranging from 50 to 720 μm have been successfully fabricated and achieved vascularization. However, the enzymolysis process may reduce EC activity, and because the enzyme is encapsulated in the bioink, degradation occurs at the same time as printing, which exerts a certain amount of time pressure (Zhu et al., 2017; Thomas et al., 2020).

Other research teams have also produced alternate solutions; for example, agarose was used as a sacrificial ink (Massa et al., 2017). It shows good rheological properties and printability similar to Pluronics when combined with alginate (López-Marcial et al., 2018). Currently, agarose has been shown capable of fabricating 100–1,000 μm diameter pipes and forming a smooth channel surface for cell inoculation (Massa et al., 2017). Another example is gelatin, that has been used as a sacrificial material and is removed by warming (Skylar-Scott et al., 2019). Channels over 400 μm in diameter could be printed with high fidelity. Another team used alginate and CaCl2 to create ultrafine fibers with a size range of 150–200 μm, and clear, interconnected microchannel structures were observed through the hydrogel as a result (Hammer et al., 2014). It is worth mentioning that some of the removal processes of these biodegradable sacrificial materials are physical processes. The utilization of their biodegradable properties needs to be further explored, as they show great potential to facilitate simplified fabrication processes and direct in vivo application of tissue constructs.

**Cells and Biological Factors**

When 3D printing techniques print with cells, the printing ink is also referred to as bioink. It includes cells, biomaterials that serve as a cell-delivery medium, and biological factors (Groll et al., 2018). The existence of cells and biological factors require higher biosafety of the materials to maintain good bioactivity of the bioinks as discussed before. Also, to prevent cells from excessive shear forces, low viscosity fluid is required. Viscosity and rheological properties greatly influence the printability of the bioinks, and they are mainly determined by the molecular weight and concentration of polymer in solution (Schwab et al., 2020). Gels with shear-thinning properties or solutions containing hydrogel precursors are preferred. In current studies of indirect 3D bioprinting for vascular systems, HUVECs, smooth muscle cells (SMCs), and fibroblasts have been used, and the cell viability is generally over 80% with high cell density (Table 3). Still, more efforts should be put on developing optimized bioinks in future investigations.

**PRECLINICAL AND CLINICAL APPLICATIONS OF INDIRECT 3D PRINTING FOR DIFFERENT TISSUE VASCULARIZATION AND BIODEGRADABLE INK SELECTION**

Nowadays, only a few tissues with less stringent vascular structures, such as cartilage and the cornea, have achieved good clinical outcomes (Rouwkema et al., 2010; Seifu and Mequenin, 2012; Pimentel et al., 2018). Advanced techniques and suitable biomaterials are essential for the development of functional engineered tissues. Indirect 3D bioprinting and the related biodegradable inks provide new opportunities for commercial development. The process has been applied to manufacturing various tissues and has achieved corresponding results. In general, our appraisal can be divided into vascular grafts and vascularized tissue; the latter includes highly vascularized tissue, vascularized osteochondral tissue, and vascularized skin, as shown in Table 3.

**Vascular Grafts**

Most vascular regions treated during surgery are larger than 1 mm in diameter, which, theoretically, both direct and indirect 3D bioprinting can achieve. Biomaterial selection requirements for tissue-engineered vascular grafts are low immunogenicity, good mechanical properties, and similarity to native tissue characteristics (Liu et al., 2018). Compared with non-degradable polymers, biodegradable natural polymers (such as decellularized tissue scaffolds and fibrin) have relatively poor mechanical properties, but they have lower antigenicity and can better simulate natural tissue structures (Aper et al., 2016). Meanwhile, biodegradable synthetic polymers (such as PGA and PCL) can have adjustable mechanical properties and degradation rate, and they are considered the ideal biomaterials for tissue-engineered vascular grafts (Liu et al., 2018). Afghah et al. used embedded extrusion bioprinting with a composite Pluronic-nanoclay support-bath and biocompatible alginate to create a branched vascular structure with diameters of several millimeters. This vascular mold showed good mechanical properties and preservation of shape fidelity after removal from the support-bath, but its biological functions have not yet been verified (Afghah et al., 2020). While Schöneberg et al. used the indirect bioprinting technique to fabricate biofunctional multi-
| Category                     | Sub-category          | Sacrificial material | Scaffold material | Cells and cell density | Cell viability | Progress                       | Limitations                  | References          |
|------------------------------|-----------------------|----------------------|-------------------|------------------------|---------------|-------------------------------|-----------------------------|----------------------|
| Vascular grafts              | Arteriole/venule      | Gelatin              | Fibrin and collagen/fibrin blends | HUVECs (~10^5 cells/ml); SMCs (~10^5 cells/ml); normal human dermal fibroblasts (−) | ~83%/91% (1/d/4/d, SMCs) | In vitro model success         | Unable to meet human transplantation standards | Schöneberg et al. (2018) |
| Highly vascularized tissue   | Branched vascular structure | Pluronic-nanoclay   | Alginate           | —                      | —             | In vitro non-cell model success | No biological function          | Afghah et al. (2020)   |
|                              | Heart-like structure  | Pluronic F127        | Alginate           | —                      | —             | Simplified models for conceptual validation | No good method to fabricate complex structures | Zou et al. (2020)     |
|                              | Valentine-shaped heart | PVA                  | Alginate and agarose | HUVECs (~10^6 cells/mL); H9c2 rat myoblasts (~10^6 cells/mL) | ~95%/90% (1/d/14/d) | A hollow structure containing a network of microfluid channels | Difficult to imitate the ultrastructure of capillaries; low degree of simulation | Zou et al. (2020)     |
|                              | Simplified cardiac scaffolds | PVA                  | PUU                | Primary human cardiac myocytes (~10^6 cells/scaffold) | 94% (1/d) | A perfusable scaffold with mechanical properties similar to cardiac tissue, and good biocompatibility with cardiac myocytes | A geometrically simplified in vitro scaffold mainly for material performance test | Hernández-Córdova et al. (2016) |
|                              | Cardiac spheroids     | Gelatin              | Collagen I and Matrigel | Cardiomyocytes with primary cardiac fibroblasts (~10^6 cells/ml in total); HUVECs (~10^6 cells/ml) | Enhanced cell viability throughout the bulk tissue compared to nonvascular tissue | A perfusable cardiac tissue that fuses and beats synchronously over a 7-day period with high cellular density | Lack of sufficient microvascular network formation; a modest contractility (~1% strain) only | Skylar-Scott et al. (2019) |
|                              | Gut-like tissue fragments | PVA                  | Matrigel, gelatin, and fibrin | Caco-2 intestinal epithelial cells; HUVECs (~10^6 cells/ml); HUVECs (~10^6 cells/ml); HepG2/C3A cells (~10^6 cells/ml) | Good cell coculture results | An in vitro gut model capable of sustaining cells long term | A simplified model mainly for conceptual validation | Hu et al. (2018)   |
|                              | Liver tissue model    | Agarose fiber        | GelMA              | HepG2, HUVECs (~10^6 cells/m1); HepG2/C3A cells (~10^6 cells/m1) | >80% (2d) | A vascularized liver tissue model for mimicking in vivo conditions and testing drug diffusion and toxicity | Difficult to imitate the ultrastructure of capillaries | Massa et al. (2017) |
|                              | Liver tissue fragments | PVA and PLA          | Gelatin            | Liver hepatocellular carcinoma (HepG2) cells (~10^3–10^5 cells/m1) | Good HepG2 cell proliferation to a high cell density | A perfusable thick engineered construct with cellular densities of native tissues | A simplified model for conceptual validation; difficult to create channels with diameter <1 mm | Pimentel et al. (2018) |
|                              | Renal proximal tubule models | Pluronic F127        | Gelatin            | Proximal tubule epithelial cells (~10^2 cells/ml); glomerular microvascular epithelial cells (−) | Healthy cell phenotype was observed | A 3D vascularized proximal tubule model that can be independently addressed to investigate renal reabsorption | The reabsorptive properties may be improved by reducing the proximal tubule lumen diameter and the separation distance between the | Lin et al. (2019) |

(Continued on following page)
layered blood vessel models in vitro, which direct printing cannot yet achieve (Schöneberg et al., 2018). They used three different degradable hydrogels with three different cell types to simulate and reconstruct the adventitia (fibroblast matrix), medial layer (elastic SMC), and intima (endothelium), and successfully replicate the three-layered natural vascular channel structure with a wall thickness of up to 425 μm, and diameters of around 1 mm (Figure 2). These bioinks provide a friendly living environment for cells. Currently, engineered vascular grafts do not meet human transplantation standards, and most of them are used for in vitro experiments, such as drug prescreening or preliminary concept verification.

Highly Vascularized Tissue

Currently, most studies are in the conceptual validation phase and, to date, there are no good methods or compatible bioinks to fabricate a 3D functional, highly vascularized network in thick tissue or organ constructs, which limits the development of tissue engineering (Duan, 2017). For highly vascularized tissue, such as liver, kidney, and heart, more precise microvessels are required, ranging from a few microns to millimeters. However, the current techniques have difficulty achieving this accuracy and resolution. Zou et al. created a valentine-shaped synthetic heart with a simplified aorta by using indirect 3D bioprinting and biodegradable alginate (Zou et al., 2020). They demonstrated no collapse of the scaffold structure and 90% cell viability. Nevertheless, the constructed microchannels are still at the level of hundreds of microns, and it remains difficult to emulate the ultrastructure of capillaries. An in vitro liver model (GelMA loaded with HepG2/C3A) was developed to test drug toxicity (Massa et al., 2017), and 3D human heart- and kidney-like objects, composed of dozens of alginate layers, were produced by embedded extrusion bioprinting (Rocca et al., 2018); however, these models were about 2 cm³ in size, with very simple vascular structure, and were used only for concept validation. Lin et al. used indirect 3D bioprinting to fabricate adjacent open cavities (representing proximal renal tubules and parallel blood vessels) embedded in the permeable ECM. After endothelialization and epithelization, the selective reabsorption and vectorial transmission of solute were realized through external circulation devices, which favor the further study of tissue-engineered kidneys (Lin et al., 2019).

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**TABLE 3** (Continued) Summary of indirect 3D bioprinting applications and bioink selection for different tissue vascularization covered in this review.

| Category                        | Sub-category | Sacrificial material | Scaffold material | Cells and cell density | Cell viability | Progress | Limitations | References               |
|--------------------------------|--------------|----------------------|-------------------|------------------------|----------------|----------|-------------|-------------------------|
| Kidney-like structure          | Pluronic F127| Alginate             | —                 | —                      | Simplified models for conceptual validation | proximal tubule and vascular conduits | No good method to fabricate 3D highly vascularized network in thick tissue or organ remodelling during endochondral bone repair | Rocca et al. (2018) |
| Vascularized osteochondral tissue | Pluronic F127| GeiMA                | Bone marrow derived mesenchymal stem cells (10⁷ cells/ml) | Cells remained viable after 24 h | A promising approach for guiding vascularization and implant remodeling during endochondral bone repair | No obvious enhanced overall-level bone formation | Daly et al. (2018) |
| Vascularized skin              | PVA          | PLCL                 | Human dermal fibroblasts (10⁷ cells/ml) | Considerable collagen and new blood vessels were observed at 4 weeks | A customized scaffold successful in vitro animal experiments and may act as a dermis substitute | A simplified model without hierarchical structure | Im et al. (2018) |
| Thermoresponsive 'stiffness memory' elastomeric nanohybrid scaffolds | PVA          | PUU-POSS             | 3T3-J2 mouse embryonic dermal fibroblasts (10⁴ cells/scaffold) | Good ingrowth of tissue and new blood vessels were observed at 4 weeks | A unique smart elastomer scaffold that can guide the growth of myofibroblasts, collagen fibers, and blood vessels at real 3D scales | Slow ingrowth of host blood capillaries; local inflammatory response | Wu et al. (2019) |
There are also some special needs for biomaterials. Highly vascularized tissues usually consist of large amounts of active cells, which require the biomaterials to be sufficiently friendly to cells and cell migration, growth, and proliferation, and should facilitate substance exchange (Xie et al., 2020). Usually, softer biomaterials are needed; indeed, one important requirement for cardiac scaffolds is that it should not provide resistance to muscle contraction during systole while providing mechanical support to resist the tensile stress during diastole (Hernández-Córdova et al., 2016). This requires the elastic modulus of the biomaterial to match that of the myocardial reported interval (7.9–1,200 kPa) (Courtney et al., 2006; Chan et al., 2013; Hernández-Córdova et al., 2016). Alginate, gelatin, fibrin, and collagen can be used for cardiac scaffolds (Alonzo et al., 2019). In an indirect 3D bioprinting case, PUU was developed as a scaffold material that had suitable elastic modulus. They also used PVA as sacrificial material, and finally created a cardiac scaffold containing 300–500 µm channels. The biomaterial showed good biocompatibility with cardiomyocytes (Hernández-Córdova et al., 2016). Meanwhile, to verify the feasibility of multicellular tissue printing, Skylar-Scott et al. reported an embedded indirect 3D bioprinting method to create perfusible vascular channels in ECM solution with organ building blocks (OBBs) composed of thousands of patient-specific-induced pluripotent stem cell–derived organoids, as shown in Figure 3. As an example, they then fabricated a cardiac tissue with physiological functions over a 7-day period with high cellular density, showing the huge prospects of indirect 3D bioprinting (Skylar-Scott et al., 2019).

**Vascularized Osteochondral Tissue**

The 3D printing technique is relatively mature in the field of osteochondral tissue engineering and has been used in the fabrication of long bones, mandibles, cheekbones, human finger bones, and other structures (Hollister et al., 2005; Lee et al., 2005; Wang et al., 2009). Biomaterials for osteochondral tissue engineering emphasize mechanical properties (Daly et al., 2017). For the design and production of vascularized bone tissue engineering scaffolds, pore-forming agents are usually added to form pores of specific sizes (>300 µm) to promote angiogenesis (Karageorgiou and Kaplan, 2005; Wang J.-Q. et al., 2019; Gonzalez-Fernandez et al., 2019). Uniform channels of controlled size that simulate the natural morphology of bone tissue, including Volkmann’s and Haversian canals for better osteogenesis, are more and more created in molds via indirect 3D bioprinting (Houben et al., 2016; Houben et al., 2017).
However, few studies have focused on the analysis of angiogenesis and new bioink development.

For endochondral bone repair, current techniques enable in vivo angiogenesis around cartilage models (Thompson et al., 2016), but the internal regions of the models remain nonvascular (Mesallati et al., 2015). Daly et al. constructed a microchanneled cartilage template using sacrificial Pluronic ink. After in vivo cultivation, they found that, compared to solid templates, channeled templates showed better vascularization, more degradation of cartilage precursor hydrogel in the core region, less ectopic bone formation, and better integration with the host tissue, which are all clinically important (Daly et al., 2016).
2018). However, more evidence is needed to demonstrate a difference in total bone formation between the channeled and solid templates for endochondral bone repair.

**Vascularized Skin**
Creating man-made skin grafts for wounds and burn healing is the primary purpose of skin tissue engineering (Adams and Ramsey, 2005). Tissue-engineered skin grafts should be non-toxic, have low inflammatory response, allow water vapor transmission, and act as a barrier. They are also expected to quickly adhere to the wound surface, have controllable degradation, and promote angiogenesis (MacNeil, 2007). Biodegradable, non-toxic, non-immunogenic, and non-inflammatory biomaterials with low risk of disease transmission and easy access are ideal skin substitutes (Vig et al., 2017). To date, indirect 3D bioprinting has not been widely applied in vascularized skin tissue fabrication. Wu et al. created a thermoresponsive stiffness memory elastomer via indirect 3D bioprinting and found that the scaffold promoted fibroblast proliferation in vitro and angiogenesis in vivo (Wu et al., 2019). Another team found that by combining bioactive peptide hydrogels with scaffolds having finger-shaped pores created via indirect 3D bioprinting, more vessels and more collagen I and III formed in the scaffolds (1m et al., 2018). Most studies that focused on tissue design have been based on simplified skin models, while few models consider controlled porosity, biodegradable material selection, and cell distribution (Lee V. et al., 2014; Van et al., 2018). It is important to mention that for the reconstruction of simple epidermis or thin dermis, vascularization is not necessary.

**CONCLUSION**
As described in this paper, tissue vascularization has always been a critical issue in tissue engineering and is key to the application and survival of engineered tissue constructs in vivo. Many accomplishments have demonstrated the feasibility of indirect 3D bioprinting for manufacturing blood vessels and vascularized tissue constructs, and most experiments show that indirect 3D bioprinting has advantages in channel structure design and construction. On the other hand, the selection of biodegradable inks is an important aspect of indirect 3D bioprinting for successful 3D tissue construct fabrication and in vitro/vivo application. Theoretically, indirect 3D bioprinting allows a wider range of materials to be used, since the use of sacrificial materials lowers the mechanical performance requirements for scaffold materials. However, this introduces new requirements for the sacrificial materials. At present, owing to the lack of satisfactory biodegradable materials, further in vivo applications are limited. A large number of studies are investigating advanced sacrificial/scaffold bioinks, which assist in the assembly of biodegradable, biosafe, bioactive, and more bionic structures at higher resolution. Efforts are now transitioning from theoretical verification to tissue and organ model construction. We expect that with the future continuous development of biodegradable materials, the use of indirect 3D bioprinting will continue to increase and will contribute to the field of tissue engineering.

**AUTHOR CONTRIBUTIONS**
JL and YY conceived and coordinated this project. YZ and YL wrote this article. LH and YS collected and summarized literatures. PL edited pictures in this paper. PG revised this paper. All authors contributed to manuscript revision, read and approved the submitted version.

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**REFERENCES**
Abdollahi, S., Boktor, J., and Hibino, N. (2019). Bioprinting of Freestanding Vascular Grafts and the Regulatory Considerations for Additively Manufactured Vascular Prostheses. *Translational Res.* 211, 123–138. doi:10.1016/j.trsl.2019.05.005

Adams, D. C., and Ramsey, M. L. (2005). Grafts in Dermatologic Surgery: Review and Update on Full- and Split-Thickness Skin Grafts, Free Cartilage Grafts, and Composite Grafts. *Dermatol. Surg.* 31 (8 Pt 2), 1055–1067. doi:10.1111/j.1524-4725.2005.31831

Afgâh, F., Altunbek, M., Dikyol, C., and Koc, B. (2020). Preparation and Characterization of Nanoclay-Hydrogel Composite Support-bath for Bioprinting of Complex Structures. *Sci. Rep.* 10 (1), 5257. doi:10.1038/s41598-020-61606-x

Alijani, W., Ullah, M. W., Zhang, X., and Yang, G. (2018). Bioprinting and its Applications in Tissue Engineering and Regenerative Medicine. *Int. J. Biol. Macromolecules* 107 (Pt A), 261–275. doi:10.1016/j.ijbiomac.2017.08.171

Alonzo, M., AniKumar, S., Roman, B., Tasnim, N., and Joddar, B. (2019). 3D Bioprinting of Cardiac Tissue and Cardiac Stem Cell Therapy. *Translational Res.* 211, 64–83. doi:10.1016/j.trsl.2019.04.004

Aper, T., Wilhelmi, M., Gebhard, C., Hoefker, K., Benecke, N., Hilfiker, A., et al. (2016). Novel Method for the Generation of Tissue-Engineered Vascular Grafts Based on a Highly Compacted Fibrin Matrix. *Acta Biomater.* 29, 21–32. doi:10.1016/j.actbio.2015.10.012

Asakawa, N., Shimizu, T., Tsuda, Y., Sekiya, S., Sasagawa, T., Yamato, M., et al. (2010). Pre-vascularization of In Vitro Three-Dimensional Tissues Created by Cell Sheet Engineering. *Biomaterials* 31 (14), 3903–3909. doi:10.1016/j.biomaterials.2010.01.105

Asghari, F., Samiei, M., Adibkia, K., Akbarzadeh, A., and Davaran, S. (2017). Biodegradable and Biocompatible Polymers for Tissue Engineering Application: a Review. *Artif. Cell Nanomedicine, Biotechnol.* 45 (2), 185–192. doi:10.3109/21691401.2016.1146731

Auger, F. A., Gibot, L., and Lacroix, D. (2013). The Pivotal Role of Vascularization in Tissue Engineering. *Annu. Rev. Biomed. Eng.* 15, 177–200. doi:10.1146/annurev-bioeng-071812-152428
Liu, C. Z., Xia, Z. D., Han, Z. W., Hulley, P. A., Triffitt, J. T., and Czernuszkaja, J. T. (2008). Novel 3D Collagen Scaffolds Fabricated by Indirect Printing Technique for Tissue Engineering. J. Biomed. Mater. Res. 85B (2), 519–528. doi:10.1002/jbm.b.30975

Liu, M. J. J., Chou, S. M., Chua, C. K., Tay, B. C. M., and Ng, B. K. (2013). The Development of Silk Fibroin Scaffolds Using an Indirect Rapid Prototyping Approach: Morphological Analysis and Cell Growth Monitoring by Spectral-Domain Optical Coherence Tomography. Med. Eng. Phys. 35 (2), 253–262. doi:10.1016/j.medengphy.2011.09.029

Liu, R. H., Ong, C. S., Fukunishi, T., Ong, K., and Hibino, N. (2018). Review of Vascular Graft Studies in Large Animal Models. Tissue Eng. Part B Rev. 24 (2), 133–143. doi:10.1089/ten.TEB.2017.0350

López-Marcial, G. R., Zeng, A. Y., Osuna, C., Dennis, J., García, J. M., and O’Connell, G. D. (2018). Agarose-Based Hydrosols as Suitable Bioprinting Materials for Tissue Engineering. ACS Biomater. Sci. Eng. 4 (10), 3610–3616. doi:10.1021/acsbiomaterials.8b00903

M. C., Reardon, P. J., Konwar, R., Knowles, J. C., and Mandal, B. B. (2017). Mimicking Hierarchical Complexity of the Osteochondral Interface Using Electrospun Silk-Bioactive Glass Composites. ACS Appl. Mater. Inter. 9 (9), 8000–8013. doi:10.1021/acsami.6b15590

MacNeil, S. (2007). Progress and Opportunities for Tissue-Engineered Skin. Nature 445 (7130), 874–880. doi:10.1038/nature05664

Madl, C. M., LeSavage, B. L., Dewi, R. E., Dinh, C. B., Stowers, R. S., Khariton, M., et al. (2017). Maintenance of Neural Progenitor Cell Stemness in 3D Hydrosols Requires Matrix Remodelling. Nat. Mater. 16 (12), 1233–1242. doi:10.1038/nmat4920

Makris, E. A., Gomoll, A. H., Malizos, K. N., Hu, J. C., and Athanasiou, K. A. (2015). Repair and Tissue Engineering Techniques for Articular Cartilage. Nat. Rev. Rheumatol. 11 (1), 21–34. doi:10.1038/nrrheum.2014.157

Malda, J., Visser, J., Melchels, F. P., Jüntgen, T., Hennink, W. E., Dert, W. J. A., et al. (2013). 25th Anniversary Article: Engineering Hydrogels for Biofabrication. Adv. Mater. 25 (36), 5011–5028. doi:10.1002/adma.201302042

Mandrycky, C., Wang, Z., Kim, K., and Kim, D.-H. (2016). 3D Bioprinting for Engineering Complex Tissues. Biotechnol. Adv. 34 (4), 422–434. doi:10.1016/j.biotechadv.2015.12.011

Massa, S., Sakr, M. A., Seo, J., Bandaru, P., Arneri, A., Bersini, S., et al. (2017). Bioprinted 3D Vascularized Tissue Model for Drug Toxicity Analysis. Biomicrofluidics 11 (4), 044109. doi:10.1063/1.4994708

Matsui, M., and Tabata, Y. (2012). Enhanced Angiogenesis by Multiple Release of Gelatin Hydrogels. Biomaterials 33 (11), 3279–3305. doi:10.1016/j.biomaterials.2012.01.007

Perrone, G. S., Leisk, G. K., Lo, T. J., Moreau, J. I., Haas, D. S., Papenburg, B. J., et al. (2014). The Use of Silk-Based Devices for Fracture Fixation. Nat. Commun. 5, 3885. doi:10.1038/ncomms4385

Pimentel-C., R., Ko, S. K., Caviglia, C., Wolff, A., Emnéus, J., Keller, S. S., et al. (2017). Three-dimensional Printing with Sacrificial Materials for Soft Matter Manufacturing. MRS Bull. 42 (8), 571–577. doi:10.1557/mrs.2017.167

Patterson, J., and Hubbell, J. A. (2010). Enhanced Proteolytic Degradation of Molecularly Engineered PEG Hydrogels in Response to MMP-1 and MMP-2. Biomaterials 31 (30), 7836–7845. doi:10.1016/j.biomaterials.2010.06.061

Pawar, S. N., and Edgar, K. J. (2012). Alginate Derivatization: a Review of Chemistry, Properties and Applications. Biomaterials 33 (11), 3279–3305. doi:10.1016/j.biomaterials.2012.01.007

Peronne, G. S., Leisk, G. K., Lo, T. J., Moreau, J. I., Haas, D. S., Papenburg, B. J., et al. (2014). The Use of Silk-Based Devices for Fracture Fixation. Nat. Commun. 5, 3885. doi:10.1038/ncomms4385

Pimentel-C., R., Ko, S. K., Caviglia, C., Wolff, A., Emnéus, J., Keller, S. S., et al. (2017). Three-dimensional Fabrication of Thick and Densely Populated Soft Constructs with Complex and Actively Perfused Channel Network. Acta Biomater. 65, 174–184. doi:10.1016/j.actbio.2017.10.047

Pitas, C. C., and Smith, D. K. (2020). Multicomponent Polysaccharide Alginate-Based Bioinks. J. Mater. Chem. B 8 (36), 8717–8788. doi:10.1039/d0tb01059g

Placenza, J. K., and Engler, A. J. (2018). Recent Advances in Extrusion-Based 3D Printing for Biomedical Applications. Adv. Healthc. Mater. 7 (8), 1701161. doi:10.1002/adhm.201701161

Poellmann, M. J., Barton, K. L., Mishra, S., and Johnson, A. J. W. (2011). Patterned Hydrogel Substrates for Cell Culture with Electrohydrodynamic Jet Printing. Macromol. Biosci. 11 (9), 1164–1168. doi:10.1002/mabi.201000004

Pollet, A. M. A. O., Homburg, E. F. G. A., Cardinaels, R., and den Toonder, J. M. J. (2019). 3D Sugar Printing of Networks Mimicking the Vasculature. Micromachines 11 (1), 43. doi:10.3390/mi11010043

Qi, X., Wang, H., Zhang, Y., Pang, L., Xiao, W., Jia, W., et al. (2018). Mesoporous Bioactive Glass-Coated 3D Printed Borosilicate Bioactive Glass Scaffolds for Improving Repair of Bone Defects. Int. J. Biol. Sci. 14 (4), 471–484. doi:10.7150/ijbs.23872

Rastogi, P., and Kandassaburamian, B. (2019). Review of Alginate-Based Hydrogel Bioprinting for Application in Tissue Engineering. Biofabrication 11 (4), 042001. doi:10.1088/1758-5090/ab331e

Reakasame, S., and Boccaccini, A. R. (2018). Oxidized Alginate-Based Hydrogels for Tissue Engineering Applications: A Review. Biomacromolecules 19 (1), 3–21. doi:10.1021/acs.biomac.7b01331

Reis, L. A., Chiu, L. L. Y., Feric, N., Fu, L., and Radisic, M. (2016). Biomaterials in Myocardial Tissue Engineering. J. Tissue Eng. Regen. Med. 10 (1), 11–28. doi:10.1002/tber.1944

Richards, D., Jia, J., Yost, M., Markwald, R., and Mei, Y. (2017). 3D Bioprinting for Vascularized Tissue Fabrication. Ann. Biomed. Eng. 45 (1), 132–147. doi:10.1007/s10439-016-1653-z
Xie, R., Zheng, W., Guan, L., Ai, Y., and Liang, Q. (2020). Engineering of Hydrogel Materials with Perfusable Microchannels for Building Vascularized Tissues. Small 16 (15), 1902838. doi:10.1002/smll.201902838

Xu, Y., Hu, Y., Liu, C., Yao, H., Liu, B., and Mi, S. (2018). A Novel Strategy for Creating Tissue-Engineered Biomimetic Blood Vessels Using 3D Bioprinting Technology. Materials 11 (9), 1581. doi:10.3390/ma11091581

Yamamura, N., Sudo, R., Ikeda, M., and Tanishita, K. (2007). Effects of the Mechanical Properties of Collagen Gel on the In Vitro Formation of Microvessel Networks by Endothelial Cells. Tissue Eng. 13 (7), 1443–1453. doi:10.1089/ten.2006.0333

Yang, S., Tang, H., Feng, C., Shi, J., and Yang, J. (2020). The Research on Multi-Material 3D Vascularized Network Integrated Printing Technology. Micromachines 11 (3), 237. doi:10.3390/omi11030237

Yeong, W.-Y., Chua, C.-K., Leong, K.-F., Chandrasekaran, M., and Lee, M.-W. (2007). Comparison of Drying Methods in the Fabrication of Collagen Scaffold via Indirect Rapid Prototyping. J. Biomed. Mater. Res. 82B (1), 260–266. doi:10.1002/jbm.b.30729

Zhang, G., Varkey, M., Wang, Z., Xie, B., Hou, R., and Atala, A. (2020). ECM Concentration and Cell-mediated Traction Forces Play a Role in Vascular Network Assembly in 3D Bioprinted Tissue. Biotechnol. Bioeng. 117 (4), 1148–1158. doi:10.1002/bit.27250

Zhu, J., Wang, Y., Zhong, L., Pan, F., and Wang, J. (2021). Advances in Tissue Engineering of Vasculature through Three-dimensional Bioprinting. Dev. Dyn. 250 (12), 1717–1738. doi:10.1002/dvdy.385

Zhu, W., Ma, X., Gou, M., Mei, D., Zhang, K., and Chen, S. (2016). 3D Printing of Functional Biomaterials for Tissue Engineering. Curr. Opin. Biotechnol. 40, 103–112. doi:10.1016/j.copbio.2016.03.014

Zhou, N., Grottkau, B. E., He, Z., Shu, L., Yang, L., Ma, M., et al. (2020). Biofabrication of valve-shaped Heart with a Composite Hydrogel and Sacrificial Material. Mater. Sci. Eng. C 108, 110205. doi:10.1016/j.msec.2019.110205

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