Bioscaffolds embedded with regulatory modules for cell growth and tissue formation: A review

Pengju Wang, Yazhou Sun, Xiaouquan Shi, Huixing Shen, Haohao Ning, Haitao Liu

Department of Mechanical Manufacturing and Automation, School of Mechatronics Engineering, Harbin Institute of Technology, Harbin, 150001, China

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ABSTRACT

The demand for artificial organs has greatly increased because of various aging-associated diseases and the wide need for organ transplants. A recent trend in tissue engineering is the precise reconstruction of tissues by the growth of cells adhering to bioscaffolds, which are three-dimensional (3D) structures that guide tissue and organ formation. Bioscaffolds used to fabricate bionic tissues should be able to not only guide cell growth but also regulate cell behaviors. Common regulation methods include biophysical and biochemical stimulations. Biophysical stimulation cues include matrix hardness, external stress and strain, surface topology, and electromagnetic field and concentration, whereas biochemical stimulation cues include growth factors, proteins, kinases, and magnetic nanoparticles. This review discusses bioink preparation, 3D bioprinting (including extrusion-based, inkjet, and ultraviolet-assisted 3D bioprinting), and regulation of cell behaviors. In particular, it provides an overview of state-of-the-art methods and devices for regulating cell growth and tissue formation and the effects of biophysical and biochemical stimulations on cell behaviors. In addition, the fabrication of bioscaffolds embedded with regulatory modules for biomimetic tissue preparation is explained. Finally, challenges in cell growth regulation and future research directions are presented.

1. Introduction

Over the past few decades, the incidence of myocardial infarction, skin inflammation, and arteriosclerosis has increased [1–3]. Removal of diseased tissue and implantation of new tissue are reliable treatment options. To meet the growing demand for human tissues and organs because of disease or accidents, scientific research is increasingly focusing on human tissue engineering and regenerative medicine. The two common methods of preparing tissues and organs are (i) the addition of bioscaffolds, growth factors, and cells into a culture medium for in vitro organization and (ii) the implantation of bioscaffolds and growth factors into the body for organization [4,5]. Cartilage, muscle, fat, skin, liver, and other organs have been successfully fabricated this way [6]. However, there are a few challenges in natural cell growth on bioscaffold surfaces: a lack of vascularized networks for transporting nutrients and waste and a lack of regulation of cell behaviors, such as the growth, proliferation, and differentiation of cells adhering to bioscaffolds [7,8]. Therefore, studies are investigating the regulatory mechanisms underlying cell behaviors [9]. Ideal tissues can be fabricated in vitro or even in vivo, not only because cells are the basis for tissue and organ formation but also based on mechanisms underlying regulation of cell behaviors, bringing new hope for disease treatment, tissue repair and regeneration, drug delivery, and medical diagnosis. Cells are usually cultured on bioscaffolds to form tissues and organs. Cells serve as the basis for tissue organization, and research on cell behaviors under stimulation is of vital importance for tissue reconstruction.

A key issue in tissue regeneration is bioscaffold fabrication. Bioscaffolds are complex structures. Their 3D topography, surface topology, and biophysical and biochemical properties affect cell adhesion, proliferation, migration, and differentiation. With the extensive application of 3D printing technologies, the research on bioscaffold manufacturing has made significant progress. 3D bioprinting is one type of 3D printing and involves the deposition of biomaterials or cells [10,11]. With regard to bioprinting, this review discusses two modalities: 3D printing of acellular bioinks with subsequent cell seeding [12,13], and direct 3D printing of cell-laden bioinks [14,15].

In recent years, microfluidic technologies have been used in in vitro culture and tissue reconstruction. The flow channel accuracy of microfluidic chips matches the mammalian cell diameter, ensuring sufficient
oxygen, and nutrient transport to cells [16]. In addition, compared to conventional culture devices, it is easier to achieve experimental variables and accurate simulations of microenvironments for cells in 3D. Cell behaviors can be regulated through biochemical and biophysical stimulations. Biophysical stimulation cues include matrix hardness, surface topology, stress and strain, hydrostatic pressure, electromagnetic field, ultrasound, and light, while biochemical stimulation cues include growth factors, proteins, kinases, and magnetic nanoparticles (MNPs). The effect of biochemical stimulations on cells needs to be considered because of the introduction of chemical factors. Surface morphology affects cell adhesion and growth. Mechanical stimulations affect cell proliferation, migration, and phenotypes.

3D bioprinting technologies adopt the concept of discrete stacking in order to directly integrate microfluidic chips with micron-level accuracy, which provides a tool for studying cell growth and tissue formation. Compared to 3D bioprinting, conventional manufacturing methods, such as molding-pouring-demolding, can neither prepare relatively complex structures nor ensure a sterile microenvironment for cell growth [17]. Conventional manufacturing methods also often have the disadvantages of chemical reagents remaining, complex processes, or long preparation cycles [18]. Although other emerging technologies, such as soft lithography and electrospinning/electrowriting, have become essential tools of tissue engineering, there are limitations in bioscaffold fabrication because of the multiscale morphological size of 3D bioscaffolds and the significantly different sizes of macrostructure and microstructure layers. Not only can the surface topology for cell adhesion and mechanical be arbitrarily changed using 3D bioprinting, we can also achieve synergetic regulations with different stimulations.

Fig. 1. Schematic diagram of cell culture and tissue reconstruction strategies in tissue engineering. (A) Conventional tissue engineering bioscaffolds. (B) Tissue engineering bioscaffolds embedded with biochemical and biophysical regulation modules, including surface microhardness, external stress and strain, noncontact stimulation cues, and hydrostatic pressure.
Using micro-computed tomography (μCT), 3D software, or corrosion casting of reverse engineering [19,20], bioscaffolds can be fabricated through 3D printing, and sufficient regulation of cell growth by biophysical and biochemical stimulations might be achieved.

3D bioprinting technologies and regulation of cell behaviors involving cells, bioinks, cell culture bioscaffolds, and tissue reconstruction are widely used in tissue engineering. However, the mechanism underlying regulation of cell behaviors is still unclear. Although much progress has been made in tissue reconstruction research, studies mainly focus on the pure mechanism underlying regulation of cell behaviors within the scope of bioengineering and bioscaffold modeling in tissue engineering. Several studies have reported strategies based on bioscaffold fabrication and regulation of cell behaviors. However, to our knowledge, there is no systematic, comprehensive research on cell behaviors integrating bioscaffold loaded with cells and regulation modules. There is a need for implanting cell-loaded bioscaffolds into the human body and regulate cells into ideal tissues. This review discusses tissue engineering, microfluidic chips, 3D bioprinting, and multidisciplinary knowledge of mechanics, materials science, and biomedical engineering to study the methods of fabricating bioscaffolds that conform to implantation standards in regenerative medicine.

Methods include bioscaffold fabrication and regulation of cell behaviors (Fig. 1). The mechanism underlying regulation of cell behaviors can be determined by studying the effects of single and multiple stimulations on cell growth. Specifically, cell culture bioscaffolds embedded with regulatory modules may be fabricated on the basis of biophysical and biochemical stimulations and 3D bioprinting technologies. The review also provides an overview of state-of-the-art bioink preparation methods and common 3D bioprinting processes, as well as biophysical and biochemical stimulation cues for regulating cell behaviors. Especially, biophysical and biochemical stimulation cues are reviewed as examples to illustrate synergistic stimulations for cell behaviors. In addition, regulating devices and methods related to different stimulations are described. Finally, future directions and challenges in cell growth regulation and tissue reconstruction are discussed. The findings will provide new technical references for tissue preparation in tissue engineering and regenerative medicine.

2. Bioscaffold fabrication

Broadly speaking, the two indispensable aspects in bioscaffold fabrication are raw materials and processes. A bioscaffold is the application of a scaffold in bioengineering. There are two types of bioscaffolds, cell-loaded, and cell-free. Cell-loaded bioscaffolds are formed by loading biomaterials into cells, while cell-free bioscaffolds are formed by cell adhesion. Different biomaterials and manufacturing methods contribute to the diversity of bioscaffolds. We first discuss bioinks and bioscaffold fabrication. Next, relevant techniques, such as 3D bioprinting, electrospinning/electrowriting, and soft lithography, are reviewed.

2.1. Bioinks for bioscaffold fabrication

2.1.1. Definition and types of bioinks

Bioinks are inks used for 3D bioprinting. They are cell-laden biomaterials that are cell friendly, implantable, and degradable, with a desirable immune response [18]. Accurate selection of bioinks directly affects cell growth and tissue formation, and bioinks should have biophysical and biochemical properties close to the original tissue. In addition, as bioscaffold raw materials, bioinks should ensure cell survival during molding and match the molding process. In some cases, to successfully fabricate high-precision bioscaffolds, bioinks need to have specific properties, for example, shear-thinning properties (liquid characteristics and solid characteristics at different shear rates) and cross-linking properties that contribute to process diversity and structure stability. The usual cross-linking reactions include alginate and calcium ion cross-linking [21], temperature-sensitive cross-linking [22], and light cross-linking [23] (Table 1).

Many biomaterials are used for cell growth and tissue formation, such as hydrogels. Some commonly used bioinks include natural biomaterials, synthetic materials, composite biomaterials, and decellularized extracellular matrices (dECMs) (Fig. 2). Natural biomaterials include gelatin, alginate hydrogel, collagen, and silk fibroin (SF) [24]. Natural biomaterials have good biocompatibility, numerous sources, and low cost. However, they are unstable and easily degradable. Synthetic materials include polyactic acid (PLA), polyglycolic acid copolymer (PGA), polyacrylamide gel, graphene hydrogel, poly(glycerol sebacate) (PGS), and modified biomaterials, such as methacrylic acid gelatin (GelMA), metal-based biomaterials, and phosphorylated poly(sebacoyl diglyceride) [25-28]. Compared to natural biomaterials, synthetic materials, especially polymer hydrogels, are relatively stable, but they are still unsuitable for cell-loaded bioinks because of their high melting point. Because of functional requirements, such as cell organization and formability requirement of biomaterials, composite biomaterials and dECMs [29] have become attractive choices. Composite biomaterials include composite natural [30], synthetic [31], bio-, and nanomaterials [32] and also composite biological features [33]. These materials improve the strength and diversity of bioscaffolds. For example, one material in a composite biomaterial is sacrificial, providing a perfect solution for fabricating porous bioscaffolds. In particular, gelatin, sucrose [34], and sodium chloride [35,36] are used as sacrificial materials. Adding a photoinitiator makes a composite biomaterial cross-linkable. Although dECMs contain growth factors, enzymes, and biological macromolecules, which are beneficial for cell growth and tissue formation, it is difficult to fabricate bioscaffolds that meet mechanical requirements and structural strength using only dECMs. Therefore, usually, dECMs are combined with other materials to overcome their limitations. In humans, biomaterials are not limited to these aforementioned biomaterials, and polydimethylsiloxane (PDMS), titanium alloy, etc., are also used.

2.1.2. Preparation of bioinks

Bioink preparation involves different component designs and different processes. Bioinks are biomaterials, which can be extracted and synthesized by physical or chemical processes. To prepare synthetic materials, processes usually include dissolving raw materials using chemical reagents, mixing raw material solutions, and performing dialysis and freeze-drying to filter out and extract solutes [41]. We prepared GelMA with a specific degree of methacryloyl group substitution as follows:

1. Gelatin was dissolved in phosphate-buffered saline (PBS) through magnetic stirring and temperature control.

| Table 1 | Properties of crosslinking reactions of hydrogels. |
|---------|-----------------------------------------------|
| Hydrogel | Principle | Cross-linking method | Reversible or not | Forming efficiency |
| Alginate [37] | Linking between calcium ion and alginate | Chemical cross-linking | Irreversible | High-speed |
| Gelatin [38] | Thermosensitive cross-linking | Physical cross-linking | Reversible | Low-speed |
| GelMA [39] | Thermosensitive cross-linking | Physical cross-linking | Reversible | Low-speed |
| GelMA [40] | Photo-crosslinking | Chemical cross-linking | Irreversible | High-speed |
2. Methacrylic anhydride was added dropwise to the solution, and preheated PBS was added to stop the chemical reaction.
3. The solution obtained was packed into dialysis bags and dialyzed in deionized water for ~1 week.
4. The solute obtained was wrapped in Petri dishes and stored in a freeze dryer at ~80 °C for ~3–4 days.

To prepare metal-based [27] and synthetic polymer biomaterials [28], quenching and evaporation are usually used. To prepare dECMs, ECMs are decellularized, powdered, and pregelled to form dECM-derived hydrogels [42]. For multiple mixed materials, such as poly(ethylene) glycol-alginate [43] and alginate-gelatin–gelatin hybrid hydrogels [44], different ingredients determine the different properties of bioinks, such as viscosity, forming ability, and mechanical properties. After material selection, components are designed. There are two methods of determining the ratio of various materials selected: repeated experiments and artificial intelligence [45]. You et al. [46] investigated the alginic dialdehyde-gelatin hydrogel ratio to achieve good printability and mechanical properties of the fabricated structures, while Lee et al. [45] used collagen, hyaluronic acid, and fibrin as biomaterials. Similar methods have been reported by other studies [47,48]. Many experiments on different ratios of bioinks have been conducted. On the basis of multiple regression analysis, machine learning is used to determine the elastic modulus of structures and the printability of different bioinks.

Mixed materials are prepared by two common methods, thorough mixing of all solutions with stirring and heating and full mixing of solid materials in the molten state. Lei et al. prepared mixed materials comprising PGS and salt particles [36]. They ground salt particles into smaller sizes and then mixed in melted PGS prepolymer at different ratios to obtain bioinks. Using the cross-linking characteristics of hydrogels, combined with a variety of biomaterials and their composite materials, mixtures of cells, and bioinks with different ratios are prepared to meet specific requirements.

2.1.3. Relationship between bioinks and bioscaffolds

Accurate selection of a bioink during bioscaffold fabrication is critical. Successfully applying a solution to the design of a bioscaffold requires a certain understanding of the structures and compositions of bioscaffolds as well as the selection of a suitable bioink. Using bioinks, bioscaffolds are fabricated by the mold-casting process, 3D bioprinting, electrowriting, electrosprinning, and soft lithography. Bioinks are constituent materials of bioscaffolds and directly determine the biochemical (e.g., degradability, biocompatibility) and biophysical (e.g., hydrophilicity, mechanical strength, stiffness) properties of bioscaffolds, and the properties of bioinks can be changed. Although 3D structures of bioscaffolds depend on the processes, bioinks directly contribute to the manufacturability of the structures. In addition, connectivity of the internal structures of bioscaffolds is easily achieved by sacrificial materials as part of the bioink mixture. One bottleneck in fabricating ideal bioscaffolds is the limited number of bioinks, which, in turn, limits the types and parameters of fabrication and affects cell differentiation and adhesion on the bioscaffold surface, and tissue formation [49]. For example, the forming ability of bioscaffolds is limited by the rheological properties, freezing point, melting point, and solubility of bioinks.

2.2. Bioscaffold fabrication by 3D bioprinting

In addition to meeting the biocompatibility and biodegradation requirements considered during bioink preparation, 3D morphology and surface properties of bioscaffolds also significantly affect cell behaviors. The bioscaffold-forming precision significantly affects cell adhesion, migration, differentiation, and organization. Therefore, bioscaffold fabrication has stricter requirements. The conventional bioscaffold-forming method is molding-pouring-demolding. Although this method can be used to fabricate 2D or 3D bioscaffolds, it has a few limitations, such as the inability to ensure vascular connectivity requirements, the introduction of chemical reagent residues, complex processes, and long preparation cycles. In addition, it cannot realize the direct forming and manufacturing of complex and heterogeneous bioscaffold networks and does not regulate the 3D cell distribution in bioscaffolds.

These limitations of the conventional method can be overcome by the application of 3D bioprinting technologies. Skylar-Scott et al. developed new 3D printing devices [50], while other groups investigated bioink materials [16,51,52]. Using existing equipment, He developed new forming technologies to meet the specific requirements of bioscaffolds [53]. At present, there are three common methods of bioprinting cells: extrusion technology, inkjet technology, and ultraviolet (UV)-assisted technology.

2.2.1. Extrusion-based 3D bioprinting

Extrusion-based 3D bioprinting (Fig. 3A) is widely used to print
gradient cell-loaded heterostructures. In extrusion-based 3D bio-
printing, bioinks form continuous microfilaments under an air pressure
or mechanical forces. The microfilaments contain cells and are contin-
uously stacked and formed under a specified printing path. The printing
of complex structures is of two types, coaxial [54] and multiple nozzle
[55]. Heinrich et al. [56] mixed GelMA and cells into bioinks. On the
basis of the physical properties of GelMA, the extruded bioinks were
cured by the substrate–bioink temperature difference. Ozbolat et al.
[57] used extrusion-based 3D bioprinting with sacrificial bioinks to
fabricate a microfluidic device. Gao et al. [47] used coaxial
nozzle-assisted 3D bioprinting to create a cell-laden hydrogel 3D struc-
ture with built-in microchannels for nutrient delivery. In addition,
Andrique et al. used extrusion-based 3D bioprinting with alginate
hydrogel as a bioink to investigate cell self-organization [58]. Lei et al.
[36] reported a general strategy for 3D printing thermosets. Using so-
dium chloride particles, they bioprinted various thermoset constructs
with extrusion-based 3D bioprinting.

However, extrusion-based technology has limited bioprinting reso-
lution. Future directions may focus on 3D bioprinting in suspension
baths [59], which will enable bioprinting 3D complex structures with
high-precision. In addition, a too small nozzle diameter during molding
easily causes blockage and subjects cells to severe pressure, which af-
ects cell survival. Although extrusion materials can be formed on a solid
support or directly in a liquid environment, the forming speed in a liquid
environment is relatively slow, and the problem of bioink salivation
needs to be overcome. However, extrusion-based 3D bioprinting is still
the most common bioprinting technology.

2.2.2. Inkjet 3D bioprinting

The principle of inkjet 3D bioprinting (Fig. 3B) is based on droplet
formation under an electric field. Droplets overcome surface tension and
are sprayed. Compared to extrusion-based 3D bioprinting, inkjet 3D
bioprinting can print with small-diameter nozzles and high-
concentration cell-loaded bioinks. Xie et al. fabricated GelMA micro-
spheres using inkjet 3D bioprinting [41]. Lv et al. used inkjet 3D bio-
printing to study microfluidic chip–based bioscaffolds [60]. Experimen-
tal results show that electric field forces can realize the 3D spatial
positioning of cell-loaded bioinks and maintain the integrity and
better accuracy of corresponding structures. By adding electrostatic jet
deflection devices, Liashenko et al. obtained submicrometer features
using inkjet 3D bioprinting [61]. These findings indicate that living cells
can achieve ideal 3D deposition by a change in printing parameters,
which obtains accurate tissue reconstruction through microfluidic
chip–based bioscaffolds. However, 3D bioscaffolds fabrication is not just
the accumulation and superposition of bioinks; it is also the fabrication
of high-gradient, multimaterial, multicomponent heterostructures. Ink-
jet 3D bioprinting has a few limitations, which need to be further
studied.

2.2.3. UV-assisted 3D bioprinting

UV curing is noncontact and nozzleless. It is directly applied to print
high-precision, heterogeneous, and complex structures. It avoids prob-
lems such as nozzle blockage and viscosity limitation, and is used for in
situ printing [62]. Photosensitive hydrogel materials are directly used to
fabricate cell-loaded structures or microfluidic devices used for cell
culture. UV-assisted 3D bioprinting is of two types, stereolithography
(SLA) and digital light processing (DLP) [63]. SLA uses a laser to directly
scan patterns through a point, line, and surface. DLP (Fig. 3C) is more
complex.

Compared to extrusion-based 3D bioprinting, UV curing can print
biological powders with a higher cell concentration. For example, after
the photosensitive hydrogel GelMA was mixed with fibroblasts, a 3D

Fig. 3. Schematic diagram of common 3D bioprinting technologies, bioprinting devices, and fabricated bioscaffolds. (A) Extrusion-based and (B) inkjet 3D bioprinting technologies. Reproduced with permission [60]. Copyright 2020, IOP Publishing. (C) UV-assisted 3D bioprinting. Reproduced with permission [64]. Copyright 2019, Springer Nature. (D) Multihead 3D bioprinting device of our laboratory. (E) Multinozzle 3D printing head. Reproduced with permission [50]. Copyright 2019, Springer Nature. (F) Complex 3D vascular bioscaffolds. Reproduced with permission [52]. Copyright 2019, The American Association for the Advancement of Science. UV, ultraviolet.
adhesion plays a key role in forming the shape and direction of tissues, is easily affected by the treatment of corresponding adhesion surfaces:

3. Stimulation cues for cell adhesion

and extrusion-based 3D bioprinting [68], in order to expand the freedom are combined into a single-step approach, such as inkjet 3D bioprinting [64]. Hong et al. fabricated cartilage tissues using DLP with a SF hydrogel [65]. However, the high cost, limited photoinitiator materials, and relatively low forming rate limit the wide application of UV-assisted 3D bioprinting.

2.2.4. Comparing pros and cons

In general, of the three types of 3D bioprinting, extrusion-based 3D bioprinting has the highest forming efficiency. The effects of inkjet and UV curing on cells are small, so the cell density is large. UV curing has the highest forming accuracy and the best forming stability. All three types can ensure flexibility. The materials used in extrusion-based 3D bioprinting are the most extensive, while UV-assisted 3D bioprinting is limited by printing materials. In addition, 3D bioprinting tissues and organs is not a simple voxel increase. The low cell and biomaterial throughput limits the application of UV-assisted 3D bioprinting in the fabrication of macro cell-loaded 3D structures. Melt electrowriting, a type of inkjet 3D bioprinting, usually uses polymers. This method was expanded with the advent of hydrogel materials [66].

The tissue networks printed by bioprinters are not limited to simple tissues; tissues with complex vascularized networks, such as the heart, are also being bioprinted. Our laboratory has a multifile 3D bioprinting device (Fig. 3D), which bioprints tissues and organs using the synergy of multiple nozzles. To improve bioprinting efficiency, bioprint complex structures, and increase the types of cell bioprinting, both bioink materials and printing devices [50] have been researched and developed (Fig. 3E). Lei et al. [67] built a four-axis bioprinting system with rotary receivers to fabricate tubular bioscaffolds. In addition, a highly complex network of vascularization has been successfully fabricated (Fig. 3F) [52]. For cell-free biomimetic bioscaffold fabrication, other 3D printing technologies, such as fused deposition modeling (FDM) 3D printing [34, 35], are usually involved. In addition, multiple bioprinting technologies are combined into a single-step approach, such as inkjet 3D bioprinting and extrusion-based 3D bioprinting [68], in order to expand the freedom of architecture design and the use of multiple materials and cell types.

3. Stimulation cues for cell adhesion

3.1. Stimulation with surface modification for cell adhesion

Tissues and organs are formed through cell differentiation [69]. Cell adhesion plays a key role in forming the shape and direction of tissues, laying the foundation for tissue formation [70, 71]. In vivo microenvironments are complex, and conventional animal tests cannot accurately simulate cell adhesion in the human body [11]. Therefore, cell adhesion is widely studied in vitro. In tissue engineering, the cell adhesion degree is easily affected by the treatment of corresponding adhesion surfaces: surface modifying or surface coating. Superior cell adhesion is the basic condition to maintain the stability of tissue structures and promote cell proliferation and differentiation. However, abnormal cell adhesion leads to vascular disorders and even tissue-related diseases, such as cardiovascular diseases and cancer [72, 73].

Bioprinting technologies can significantly control cell positioning. Using cell-laden bioinks, 3D bioprinting of proper process paths can achieve an ideal spatial cell distribution. Lei et al. fabricated a vascularized tissue using coaxial 3D bioprinting [74]. Briefly, cells were loaded in a sacrificial material, which was then dissolved and a structure was prepared. The cells automatically deposited and adhered to the bioscaffold surface. However, one limitation of this method is the lack of bioinks matching corresponding bioprinting processes. Microneedles are widely used in biomedicine. Cell microneedles [75, 76] are used for quantitative injection of cells at designated adhesion surfaces. Although cell-free bioprinting and subsequent cellularization provide a way for fabricating cellular microneedles, they cannot control patterned cell distribution on bioscaffolds. The other two commonly used control methods during cell adhesion are bioscaffold surface modification with topography and bioscaffold surface coating with other materials.

Surface modification created by surface topography, including the pillar, pit, and grating shape; feature size; spacing; and substrate arrangement, plays a key role in the induction of cell adhesion [63]. Generally, cell adhesion can respond to surface topological structures at the micrometer and even the nanometer scale. For example, the presence of nanotopography on microscale wrinkles of substrates promotes the adhesion of fibroblasts, endothelial cells, osteoblasts, and human mesenchymal stem cells (MSCs) [77]. Cell adhesion is the basis for cell proliferation, cell migration, cell differentiation, and tissue formation, and a comprehensive understanding of the effects of surface modification on cell adhesion is important for regenerative medicine and diagnosis. Therefore, the adhesion of cells on bioscaffold surfaces has attracted increasing attention in tissue engineering. Bioprinting of tissue engineering bioscaffolds paves the way for the creation of surface structures. After determining models and bioprinting materials, a high-precision-topology bioscaffold can be easily fabricated using 3D bioprinting. For example, Xie et al. [78] fabricated heterogeneous bioscaffolds with different fiber diameters and pore sizes using inkjet 3D bioprinting (Fig. 4A). They systematically investigated cell adhesion affected by bioscaffolds with controllable pore sizes, the difference in the adhesion characteristics of bioscaffolds with different fiber diameters, and the different adhesion behaviors of large and small cells on the same bioscaffold. Results showed that cells in bioscaffolds with small pores and large fibers have superior adhesion behavior. Compared to large cells, small cells are more inclined to adhere and bridge adjacent fiber spaces. In addition to fabrication of grid structures, 3D bioprinting can also fabricate bioscaffolds with pillars and pits. Choi et al. [79] fabricated microwrinkle circular pits on the inner surface of a bioscaffold to achieve accurate cell adhesion. 3D bioprinting provides many methods of fabricating this type of bioscaffold surfaces. Common methods include auxiliary printing of support materials or sacrificial materials, bioprinting in suspension baths, and auxiliary molding-pouring-demolding. In addition to surface topologies, post-treatment processes may also affect cell adhesion. Researchers have demonstrated a microchannel of a fabricated bioscaffold to increase its adhesion properties in an oxygen plasma chamber [80]. Compared to surface topology modification, these methods are relatively simple. Yet, precise treatments of bioscaffold surfaces are difficult to achieve because of the requirement of micron-level structures for cell adhesion, leading to inaccurate adhesion positions. Therefore, bioscaffolds with surface topography are preferred. Surface topography modification is a physical treatment that achieves cell adhesion without introducing other materials on the bioscaffold surface. So, bioscaffolds with surface topography modification, processed by 3D bioprinting, and appropriate post-processing, have good cell compatibility, in addition to direct and effective promotion of cell adhesion.

3.2. Stimulation with surface coating for cell adhesion

Unlike the direct effects of surface topography on cells, new chemical factors and material coatings on 3D-bioprinted bioscaffold surfaces also induce corresponding cell adhesion behaviors. In contrast to surface topography modification, surface coating is a conventional and easily controlled method because of its similarity to conventional experimental and clinical injection. In an in vivo setting, it may not be necessary to add additional chemical factors into bioscaffolds. However, coating materials on bioscaffold surfaces is required in order to change the hydrophilicity and hydrophobicity of bioscaffold surfaces in vitro. Surface coating is realized not only manually but also by 3D bioprinting. Although the manual method is simple, accurate surface coating is difficult. Gao et al. [81] used extrusion-based 3D bioprinting to deposit a coating material on a bioscaffold surface. The position of the coating material on the bioscaffold was controllable. Natural materials, such as poly-γ-lysine [82], and synthetic materials are widely used for surface
coating. Surface coating regulates cell adhesion. There is no need to form complex structures of coating materials. Hence, surface coating by 3D bioprinting, including but not limited to extrusion-based 3D bioprinting, is feasible. In addition, gradient cell adhesion on the same bioscaffold surface can be achieved without time-consuming and extraordinarily complex processes. For example, 3D-bioprinted gradient coating can be easily achieved by controlling the pressure and moving speed of the nozzle in extrusion-based 3D bioprinting, the droplet velocity in inkjet 3D bioprinting, and the exposure time in UV-assisted 3D bioprinting.

Do different concentrations of coating materials also affect cell adhesion? He et al. developed composite bioscaffold surfaces using diverse combinations of magnetic graphene oxide, ferroferric oxide, and graphene oxide in order to determine the relationship between cell adhesion behavior and the corresponding chemical composition of bioscaffold materials (Fig. 4B) [83]. Within a specific concentration range, the cell adhesion density increased with an increase in the concentration of magnetic graphene oxide. However, beyond a specific concentration threshold, the cell adhesion density significantly decreased. There are significant differences in the regulation of cell adhesion with different concentrations of materials used for coating cell adhesion surfaces. Therefore, an appropriate material concentration is important. Microfluidic technologies have the advantage of controlling, operating, and inspecting complex fluids at the microscale, which enables a controllable gradient distribution of concentration by designated microchannels [16]. Combining 3D bioprinting and microfluidic technologies enables fabrication of surface-coated bioscaffolds with controllable distribution of material concentration, promoting the regulation of cell adhesion and tissue regeneration in vitro. In addition, for more accurate cell distribution, other technologies, such as casting, may also be involved, partly because the fabrication of molds for surface coating can be done by 3D printing.

3.3. Biophysical stimulations for cell growth and tissue formation

In conventional tissue engineering, bioscaffolds are successfully fabricated by combining bioink preparation and 3D bioprinting. Cell-loaded or cell-loaded bioscaffolds are distributed in 3D space according to the location given by the 3D bioprinting device. Subsequently, tissues are prepared in a culture medium or bioreactors for regenerative medicine and cell therapies; however, cell proliferation, migration, and differentiation in specified growth directions cannot be guided [84]. The
tissues formed are quite different from ideal tissues. Studying the mechanism underlying regulatory mechanism of bioscaffold-based approaches for cell behaviors provides a new way to solve this problem. The common regulation methods include biophysical and biochemical stimulations. The ultimate goal of bioscaffolds is to be implanted into the human body. However, biochemical stimulations might lead to cytotoxicity for cells. Applying biophysical stimulations to cells can effectively avoid this problem. To better define the regulatory behavior of stimulations, it is important to investigate biophysical stimulations. Several factors contribute to biophysical stimulations, such as change in surface microhardness, external force and strain, and non-contact-dependent factors (light, ultrasound, electrical, and magnetic stimulations).

3.3.1. Surface microhardness

Surface microhardness contributes to the migration behavior of cells. A change in microhardness is achieved by controlling 3D bioprinting processes and adding external supports [85], electric and magnetic fields [86], and gamma ray radiation [87] to the substrates formed. Matrix hardness for cells is good for tissue regeneration, but too low hardness is not conducive to structural stability and cell adhesion on hard materials is difficult. To some extent, the migration behavior of vascular cells to stiffer regions is observed on uniform substrates, promoting the regulatable direction, and distribution of newly formed tissues on bioscaffolds. For example, Song et al. fabricated fibrous networks with different stiffness values and found that meniscal fibrochondrocytes migrate to stiff fibers through the networks after day 6 (Fig. 5A) [88]. In addition, they found better repair effects of damaged meniscus tissues in stiff fibrous networks.

To regulate stiffness-responsive cell behaviors, accurate fabrication of bioscaffolds with surface microhardness is necessary. Xue et al. used DLP 3D bioprinting to fabricate a hydrogel bioscaffold with regionally varied stiffness [89]. The surface microhardness of the bioscaffold was tuned by different exposure times on print layers (Fig. 5B–i), and fibroblasts were seeded onto the bioscaffold to investigate cell behaviors. Results showed a higher cell population (Fig. 5B–ii) and fibrous-like tissue layers in the area of a longer exposure in culture. Similarly, in inkjet 3D bioprinting, curing jetted ink droplets at different UV exposure times facilitates the formation of bioscaffold with different hardness values. In addition to photo-cross-linking, ionic cross-linking is also involved. Idaszek et al. [90] used extrusion-based 3D bioprinting to develop a microfluidic print head and formed inks with different calcium chloride–alginate mixing ratios. They achieved different surface microhardness values by regulating the cross-linking reaction between calcium ions and alginate. Studies have also reported indirect regulation of bioscaffold stiffness. Bastola et al. [91] added a magnetoreheological material in matrices to fabricate structures with tunable stiffness using extrusion-based 3D printing. They encapsulated magnetoreheological materials in the structures layer by layer, and regulated the bioscaffold hardness by applying electric fields (Fig. 5C).

Ramy et al. presented a similar method of changing matrix hardness [87]. They subjected the prepared composite films to gamma ray irradiation at various dosages. By using hybrid materials embedded with materials responsive to electrical stimulation, magnetic stimulation, or light, microhardness can be changed using 3D bioprinting and external fields. Another representative method of regulating surface microhardness is to add external supports [85]. Cell substrates bonded with underlying patterned supports can be fabricated by 3D bioprinting, followed by corresponding assemblies. Therefore, cell migration can be observed and quantified by varying the geometry underlying corresponding cell substrates. Usually, connectors integrated in these two modules may be used to achieve high-precision assembly and microscale regulation.

3.3.1.1. Discussion of pros and cons. Overall, the reported methods have a few advantages and disadvantages. Both controlling 3D bioprinting processes and applying an electric field, magnetic field, and gamma ray radiation have specific requirements for bioinks and materials. In addition, the time and cost of complex processes for adding external supports require extra designing and preparation. Compared to other treatments, realizing different hardness distributions in cell matrices during forming is the most effective and precise method. On the basis of cross-linking reactions (ionic or photosensitive cross-linking), ideal hardness can be achieved by extrusion-based, inkjet, or UV-assisted 3D bioprinting. For example, Giachini et al. [92] used extrusion-based 3D bioprinting to design and print different structures with a controllable hardness gradient (Fig. 5D). They considered the material distribution and composition and cross-linking processes in the structural design. Adding external supports has low requirements for 3D bioprinting to obtain the ideal matrix hardness for cells. The fabrication of these bioscaffolds with surface microhardness involves two Lego-like modules, including a bioscaffold and its corresponding external supports, so there is no need to consider the regulatory factor when bioprinting bioscaffolds and it is widely used in the study of in vitro cell culture. Non-contact-dependent methods, including applying an electric field, magnetic field, and gamma ray radiation, regulate cell behaviors in noncontact and remote ways, which are conducive to in vivo tissue formation. In addition, controllable distribution of materials related to external fields into bioscaffolds is easily achieved by 3D bioprinting. However, most reports about regulation of cell migration by surface hardness are still focused on phenomenon. An in-depth study of the regulatory mechanism, such as the redox state [93], needs to be considered.

3.3.2. External force and strain

External force and strain affect the secretion of growth factors, proteins, and gene expression by acting on cells through force conduction. They also affect the direction of cell growth, proliferation, migration, and differentiation. External forces include shear force, hydrostatic pressure, and cyclic stress, generated by a controlled flow rate of the culture medium [94–96]. Bioscaffolds with self-designed regulation, including microchannels and resilience materials, provide the mechanical strain stimulation [97–99]. Shear force acting on cells is realized by controlling fluids. Trachtenberg et al. [100] fabricated bioscaffolds with different pore sizes using 3D bioprinting (Fig. 6A–i), forming a shear stress gradient within the bioscaffolds after perfusion of the culture medium. Molladavoodi et al. [101] found that a low shear stress created by the flow velocity of the culture medium promotes wound healing (Fig. 6A–ii). Microfluidic methods that generate cyclic stress are similar to those that generate shear stress; the only difference is the pump that drives the flow velocity of the culture medium. For example, researchers generated a cyclic fluid stress with different reciprocating frequencies by a self-designed mini-oscillator [102]. They found that a specific frequency of reciprocating force promotes fibroblast arrangement and polarization (Fig. 6B–i) but that another frequency promotes fibroblast differentiation (Fig. 6B–ii). Optimization of the reciprocating frequency is important for cell differentiation and tissue regeneration. In addition, there are reports on hydrostatic pressure. Park et al. [103] used extrusion-based 3D bioprinting to fabricate composite bioscaffolds with different mass fractions of beta-tricalcium phosphate (β-TCP) (Fig. 6C–i). After MSC seeding, they generated a more marked ECM by applying hydrostatic pressure (Fig. 6C–ii).

In contrast with the combination of shear and stress for direct cell regulation, strain indirectly acts on cells by changing the shape of cell matrices. Usually, methods of applying strain include external and self-generating control. The expansion and contraction due to a change in the air pressure in microchannels provides external control, while the resilience and gravity of bioscaffolds provide self-generating control. Kim et al. [104] used UV-assisted 3D bioprinting to fabricate a microfluidic chip that enabled the real-time control of strain in capillary channels surrounded by source and sink channels (Fig. 6D–i). The strain
Fig. 5. Regulation surface microhardness for cell behaviors. (A) Migration behavior of meniscal fibrochondrocytes on a soft/stiff fibrous network. Reproduced with permission [88]. Copyright 2019, Wiley-VCH. (B) Fabrication of bioscaffolds with regulatable surface microhardness and its effect on cell behaviors: (i) DLP 3D bioprinting of bioscaffolds with regulatable surface microhardness and (ii) cell migration and proliferation on bioscaffold layers with different surface hardness values. Reproduced with permission [89]. Copyright 2019, American Chemical Society. (C) Extrusion-based 3D printing of structures to modulate surface micro-hardness using electric fields. Reproduced with permission [91]. Copyright 2017, Elsevier. (D) Design strategy of hardness gradient with controllable material distribution and composition for extrusion-based 3D bioprinting. Reproduced with permission [92]. Copyright 2020, Giachini et al. DLP, digital light processing; MR, magnetorheological materials.
was generated by stretching the flanking channels. Engeland et al. [105] applied a cyclic circumferential strain on vascular and smooth muscle cells and found both types of cells arranged and elongated in the vertical direction of strain (Fig. 6D–ii). Hsieh et al. used self-generating control to design a model to generate gradient strain for cells [4] by the actions of the resilience and gravity of a deformed bioscaffold (Fig. 6E). To achieve this type of regulation, bioscaffolds and regulatory modules can be fabricated separately using 3D bioprinting, followed by corresponding assemblies.

3.3.2.1. Discussion of pros and cons. 3D Bioprinting has outstanding applications in fabricating complex bioscaffolds embedded with the regulatory modules of external force and strain. Taking external force regulation as an example, precise bioscaffolds embedded with regulatory modules are easily fabricated by 3D bioprinting. Because of the direct control of fluids acting on cells, there is usually no need to integrate the regulatory modules into bioscaffolds, but the modules are needed for bioscaffold strain-mediated regulation. 3D bioprinting of high-precision structures facilitates not only integration between bioscaffolds and regulatory modules but also the fabrication of micro-channel bioscaffolds. UV-assisted 3D bioprinting is a feasible method of fabricating these structures. Especially, for fabricating microchannel bioscaffolds, inkjet and extrusion-based 3D bioprinting of sacrificial materials provide supports during forming. These sacrificial materials can be dissolved postprocessing. Numerous regulation methods can be generated by combining the interdisciplinary knowledge of mechanics and physics. In addition, the regulation of parameters by these methods helps clarify the mechanism underlying cell behaviors. However, long-term studies need to consider the degradation of bioscaffolds and avoid the problems of pollution and operational errors generated during bioscaffold operation, which may be harmful to cells. Mechanical regulation of self-generating control does not need to consider the introduction of harmful reagents, so self-generating control has unique advantages in in vivo regulation, although there are few reports compared to external regulation. Overall, bioscaffold design has been widely studied, and it is expected that researchers will conduct in-depth studies of cell behaviors by stress and strain stimulation.

3.3.3. Non-contact-dependent factors

Non-contact-dependent factors affect cell migration, proliferation, and differentiation, in addition to intracellular microenvironments. Non-contact-dependent factors include light, ultrasound, electrical, and magnetic stimulations [106–109]. These stimulation cues regulate cell behaviors by acting directly on cells or changing cell matrices, such as surface microhardness. Direct actions on cells activate intracellular signaling pathways and enhance intracellular delivery, while indirect stimulations regulate cells by controlling cell microenvironments. Zhu et al. [110] used UV-assisted 3D bioprinting to fabricate a composite GelMA and polyethylene (glycol) diacrylate bioscaffold. They applied red laser light to stimulate neural stem cells seeded on the bioscaffold (Fig. 7A–i). After culturing for a few days, they observed neuronal differentiation of neural stem cells (Fig. 7A–ii). Ramya et al. fabricated methyl methacrylate, iron-zinc, and HA bioscaffolds [87] and subjected them to different doses of gamma rays, which changed the surface roughness, hydrophobicity, and conductivity and enhanced the viability of NIH-3T3 fibroblasts. Osborn et al. [111] used 3D bioprinting to fabricate porous PLA bioscaffolds for ultrasound stimulation (Fig. 7B–i) and seeded human MSCs on the bioscaffolds. Compared to controls, cell morphologies showed a significant increase in proliferation under low-intensity pulsed ultrasound stimulation (Fig. 7B–ii). An et al. reported an indirect stimulation method of using ultrasound stimulation [112]. They cultured cells on bioscaffold surfaces and modified the surfaces for cell adhesion through low-intensity pulsed ultrasound stimulation. After cell culture, microfilaments, pseudopodia, and ECM mineralization nodules of cells increased, and the related protein expression for osteoblast differentiation was upregulated.

Stimulation based on bioscaffolds and material coating has also been reported. Liu et al. team [113] used extrusion-based 3D bioprinting to fabricate functionalized poly(propylene fumarate) (PPF) bioscaffolds coated with a nanocomposite of single-stranded DNA (ssDNA) and carbon nanotubes (CNTs) (Fig. 7C–i). Pre-osteoblast adhesion and proliferation were enhanced by electrical stimulation after cell seeding (Fig. 7C–ii). Compared to external stimulation cues, Chen et al. [114] formed a built-in stimulation of a nanoscale electric field on bioscaffold surfaces by changing the piezoelectricity of materials using polarization. Continuous electrical stimulation was provided without additional external fields. In addition, cell behaviors can also be regulated through nanovibrational stimulation using piezoelectric materials or piezo actuators [115]. Bioscaffold surfaces for cell adhesion are deformed by an electric field force. A vibration stimulation can be created by applying a time-varying electric field. Lin et al. [116] applied a magnetic field to study cell behaviors with surface-functionalized iron oxide nanoparticles (IONPs). They used extrusion-based 3D bioprinting to fabricate bioscaffolds embedded with IONPs (Fig. 7D–i). A magnetic field can improve the proliferation and mineralization of osteoblasts (Fig. 7D–ii).

3.3.3.1. Discussion of pros and cons. Various stimulations enable remote regulation of cell behaviors without cell contact, decreasing the pollution of cell culture chambers caused by conventional regulation. Some factors are required for imitating physiological microenvironments. For example, electrical stimulation stimulates myocardial cell growth. There is a closed relationship between electric and magnetic fields in physics, leading to many similarities in their stimulations in the design methods, principles, and regulation modes of cell behaviors. Ultrasound stimulations have been applied to study cell behaviors both in vitro and in vivo [117]. In addition, UV light irradiation kills bacteria. Compared to conventional methods, a bioscaffold can serve as a source of stimulations including but not limited to electrical, magnetic, ultrasound, and light stimulations [118].

External fields usually act on cells through corresponding media materials, and these cell stimulations may be mediated by bioscaffolds embedded with these media materials. To target regulation of cell behaviors, piezoelectric and light-responsive materials and MNPs, need to be reasonably distributed in the 3D space of composite bioscaffolds by 3D bioprinting. In contrast to UV-assisted 3D bioprinting, the distribution of these materials in bioscaffolds is more diversifed by inkjet and extrusion-based 3D bioprinting with multinozzle printheads. So far, reports on the remote regulation of cell behaviors have focused on
Fig. 7. Regulation of non-contact-dependent factors for cell behaviors. (A) Effects of light stimulation on cell growth: (i) bioscaffold fabrication by UV-assisted 3D bioprinting and schematic illustration of light stimulation and (ii) enhanced neuronal differentiation of neural stem cells under low-level light stimulation. Reproduced with permission [110]. Copyright 2017, IOP Publishing. (B) Effects of ultrasound stimulation on cell proliferation: (i) 3D-bioprinted PLA bioscaffolds visualized with SEM and (ii) enhanced osteogenic differentiation of human MSCs under low-intensity pulsed ultrasound stimulation. Reproduced with permission [111]. Copyright 2018, John Wiley and Sons. (C) Effects of electrical stimulation on cell proliferation: (i) 3D-bioprinted PPF bioscaffolds before and after functionalization with nanocomplex material and (ii) improved cell adhesion and proliferation of pre-osteoblasts under electrical stimulation. Reproduced with permission [113]. Copyright 2020, Elsevier. (D) Effects of magnetic stimulation on cell behaviors: (i) 3D-bioprinted (left) chitosan bioscaffolds and (right) chitosan bioscaffolds embedded with IONPs and (ii) improved proliferation and mineralization of osteoblasts under magnetic stimulation. Reproduced with permission [116]. Copyright 2020, Elsevier. UV, ultraviolet; PLA, polylactic acid; SEM, scanning electron microscopy; MSCs, mesenchymal stem cells; PPF, poly(propylene fumarate); IONPs, iron oxide nanoparticles.
bioscaffolds for cell culture because of the combination of external field-responsive materials and 3D bioprinting providing the possibility for precise cell regulation. However, there are few reports on direct cell stimulation, except for ultrasound stimulation. Future studies need to shift their focus from the regulation of bioscaffolds to the direct regulation of cells.

3.4. Biochemical stimulations for cell growth and tissue formation

Tissue formation is similar to seedling growth. Nutrient solutions contain nutrients that induce the cell activity of trees and provide active substances required for tree growth. Nutrient solutions for trees are similar to biochemical components for cells, and the infusion control devices of trees are like the biochemical control bioscaffolds of cells. Biochemical components induce cell differentiation and reprogramming. Biochemical stimulations mainly involve three aspects: types of biochemical factors, regulatory bioscaffolds, and concentration regulation.

3.4.1. Biochemical factors

Biochemical factors coming from internal bodies mainly include cell growth factors, cellular proteins, kinases, and others, such as heparin, cellulose, and vitamins [119]. Chemical stimulants are from outside the body, such as MNPs [120]. In vivo biochemical stimulation cues, which have good cell compatibility and do not need to consider immune effects, are usually the best choice for regulating cell behaviors. For example, Song et al. investigated the effects of transmembrane 4 L six family member 5 (TM4SF5) on cell metabolism [121]. Compared to untreated cells, after TM4SF5 treatment, cells in 3D collagen showed more aggressive growth, in addition to aggressive pseudopodia, invasive foci, invadopodia, and endothelial-like networks. Poukkula et al. analyzed the migration behavior of border cells using receptor tyrosine kinases and platelet-derived growth factor/vascular endothelial growth factor–related receptors (PVRs) [122]. They found that receptor tyrosine kinases affect the presence and size of border cells, while PVRs induce front extensions and cell migration. However, in vivo stimulation cues sometimes cannot completely meet experimental requirements. To regulate cell behaviors in different ways, synthetic reagents, including composite materials, are used. For example, antioxidants [123], poly (vinyl alcohol)–chitosan [124], and cannabinoid [125] are used to regulate cell proliferation. However, their degradation characteristics and biocompatibility need to be considered using stimulants from outside the body. Biochemical factor delivery and concentration regulation are new aspects of biochemical regulation of cell behaviors. The delivery of biochemical factors and the distribution of chemical factor concentration on demand are realized through the design of bioscaffolds. The function and implementation principle of bioscaffolds are similar to those of drug delivery, which accurately determines the injection position and corresponding controllable gradient concentration of biochemical reagents without creating bulk flows.

3.4.2. Regulatory bioscaffolds

Inductive bioscaffold materials, bioscaffolds loaded with biochemical factors, and biochemical factors generated by degradation of bioscaffolds provide stimulations. Zhao et al. [126] used extrusion-based 3D bioprinting to fabricate bioscaffolds with the regenerative material of bioactive glass (Fig. 8A–i). Cell culture experiments showed that stimulation with bioactive glass bioscaffolds enhances osteoblast migration and extramembranous osteogenesis (Fig. 8A–ii). A more common method of regulating cell behaviors is to use bioscaffolds loaded with biochemical factors. Liu et al. [127] bioprinted biomimetic gelatin/alginate hydrogel bioscaffolds loaded with niosilicates (Fig. 8B–i). They exploited the bioinert property of alginate to avoid rapid degradation of the bioscaffolds and ensure effective regulation. They found that the addition of niosilicates significantly promotes the formation of mineralized matrices in the culture medium (Fig. 8B–ii).

Biochemical factors can be not only loaded by bioscaffolds, but also generated by degradation. To generate biochemical factors, Liu et al. [128] 3D-bioprinted strontium-containing HA/PCL bioscaffolds (Fig. 8C–i). Strontium and calcium ions were released from the bioscaffolds in a sustained manner. MSC proliferation and new tissue formation on the composite bioscaffolds increased compared to PCL bioscaffolds (Fig. 8C–ii). In addition, material synthesis processes are involved before bioscaffold fabrication [27]. Bioscaffolds are also used as a delivery vehicle for biochemical reagents. Paolini et al. [129] used extrusion-based 3D bioprinting to fabricate PLA bioscaffolds and coated them with poly-amidoamine hydrogel. They observed improved cell proliferation on the coated bioscaffolds as a function of time compared to PLA bioscaffolds. However, biochemical factors have an uncontrollable release rate. Dang et al. [130] improved this by fabricating macroscale and microscale porous bioscaffolds using 3D bioprinting, and the burst release decreased. In addition, microneedles for delivery of reagents have also been investigated [75,131]. To release drugs for cell regulation, Derakhshandeh et al. prepared resin-based miniaturized needle arrays by 3D printing in support material (Fig. 8D–i) [132]. After perfusing cefazolin salt through microneedles, granulation tissues, new blood vessels, and hair significantly improved in the microneedle-assisted group (Fig. 8D–ii). Unfavorable flows created by bioscaffold delivery can be improved compared to conventional delivery methods [80].

3.4.3. Concentration regulation

The concentration of biochemical factors also significantly affects cell behaviors. A reasonable concentration of biochemical factors is required for cell proliferation, cell migration, cell differentiation, and tissue formation [133]. Gradient concentration regulation is beneficial for cell research. Microchannels can be similarly converged and branched to form a concentration gradient on the basis of the principle of series and parallel networks of circuits in electrotechnique. Nie et al. [134] fabricated Lego-like modular microfluidic devices using 3D bioprinting to create a concentration gradient for the regulation of cell behaviors. The concentration gradient was controlled by the location of microchannels and the flow velocity of the culture medium or other biochemical substances (Fig. 8E–i), and formulations with different concentrations were created by assembled microfluidic bioscaffolds. Cell culture experiments showed that the spherical size of fibroblasts increases and osteoblasts proliferate and form mutual contact under an appropriate concentration of the culture medium (Fig. 8E–ii). To create a diversified concentration gradient, auxiliary devices, such as a rocker platform, may be involved [133].

3.4.3.1. Discussion of pros and cons. Overall, in contrast to biophysical stimulations, biochemical stimulations provide unique stimulation cues for cell growth and tissue formation. The regulation of biochemical factors by bioscaffolds involves the development and delivery and the concentration control of reagents. Reagents come from direct or indirect chemical factors, such as in vivo acellularization and in vitro–synthesized composite materials. Because of good cellular compatibility and immunity, primary biochemical factors, and their derivatives in vivo will probably be the main regulatory factors. Yet, the mechanism underlying biochemical regulation of cell behaviors is relatively clear. Therefore, the focus is on the way these biochemical stimulations work and the concentration control of biochemical factors.

For reagent delivery, there are three application aspects of bioscaffolds. First, bioscaffold with materials functions of biochemical stimulations or embedded with reagents provide direct regulation of cell behaviors. 3D Bioprinting technologies have unsurpassed advantages in controlling material distribution and fabrication of materials loaded with reagents. 3D bioprinting achieves precise targeting and sustained release of stimulations. Delivery by microneedles is convenient for regulation of cell behaviors because of the advantage of minimal
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Fig. 8. Regulation of biochemical stimulations for cell behaviors. (A) 3D-bioprinted bioscaffolds of inductive materials for regulation of cell behaviors: (i) fabrication of bioactive glass bioscaffolds using extrusion-based 3D bioprinting and (ii) enhanced osteoblast migration (top) and extramembranous osteogenesis (bottom) after stimulation with bioactive glass bioscaffolds. RAW is a type of cells. Reproduced with permission [126]. Copyright 2018, John Wiley and Sons. (B) 3D-bioprinted bioscaffolds loaded with biochemical factors for regulation of tissue behaviors: (i) fabrication of gelatin/alginate hydrogel bioscaffolds loaded with nanosilicates using extrusion-based 3D bioprinting and (ii) enhanced formation of mineralized matrices in the culture medium after stimulation with bioscaffolds loaded with nanosilicates. Reproduced with permission [127]. Copyright 2020, Elsevier. (C) 3D-bioprinted bioscaffolds loaded with generable biochemical factors for regulation of tissue behaviors: (i) fabrication of strontium-containing HA/PCL bioscaffolds using extrusion-based 3D bioprinting and (ii) enhanced proliferation of MSCs and formation of new bone tissues. Reproduced with permission [128]. Copyright 2019, Elsevier. (D) 3D-bioprinted microneedle arrays to release drugs for regulation of tissue behaviors: (i) 3D-bioprinted microneedle structures for regulation of tissue behaviors by drug delivery and (ii) enhanced effect on tissue repair by microneedles; granulation tissue is denoted with black arrowheads, and hair is denoted with white arrows. Reproduced with permission [132]. Copyright 2020, John Wiley and Sons. (E) 3D-bioprinted modular microfluidic bioscaffolds to create a gradient concentration for regulation of cell behaviors: (i) several formulations with different concentrations of the culture medium created by modular microfluidic bioscaffolds and (ii) increased spherical size of fibroblasts (left) and enhanced proliferation and mutual contact of osteoblasts (right) under perfusion of an appropriate concentration of the culture medium. Reproduced with permission [134]. Copyright 2018, IOP Publishing. BGSE, bioactive glass bioscaffold extract; NM, normal medium; HA, hydroxyapatite; PCL, polycaprolactone; MSCs, mesenchymal stem cells.

invasion during drug delivery. The types and injection locations of biochemical reagents can be easily adjusted by changing the microneedle position. Second, indirect delivery can be achieved by releasing biochemical stimulation cues during the degradation of bioscaffolds. Occasionally, sustainable release of biochemical stimulation cues is required. In some cases, the method of embedding growth factors into bioscaffolds is not feasible because of the limitations of 3D bioprinting and the requirements of targeted regulation. An alternative is to generate interesting and meaningful regulatory cues through the degradation of bioscaffolds. Some biochemical factors, such as calcium, iron, zinc, and magnesium ions, are beneficial for cell growth and tissue formation. If these factors are considered during the design and preparation of bioscaffold materials, cells can be stably regulated for a long time. Third, bioscaffolds can act as transportation tools for delivering biochemical reagents to cells. Cells are commonly disrupted by conventional targeted injections of reagents. These defects can be eliminated by following bioscaffold-based approaches through guiding the smooth flow and precise delivery of biochemical reagents. The only requirements are an appropriate geometry design and 3D bioprinting. In addition, microchannel bioscaffolds are involved in concentration control. Cross sections of microchannels and connections between microchannels affect the final concentration of prepared reagents. Multiple-component control puts high demands on the structures of these microchannels. Modular fabrication facilitates various types of preparation and decreased the complexity of the overall structures and manufacturing difficulties. As discussed previously, 3D-printed connector modules can help achieve the assembly between modules. Therefore, this method is operable and has application prospects.

Realization of these aforementioned regulation methods requires not only the structural accuracy of bioscaffolds but also the function of biochemical factors. Biochemical factors may be modified by 3D bioprinting, such as high-intensity UV exposure and chemical reagents (e.g., calcium chloride) added during cross-linking of extrusion-based, inkjet, and UV-assisted 3D bioprinting. These aspects must be considered to avoid inaccurate regulation of cell behaviors. In addition, the degradation rate of bioscaffold materials also needs to be investigated to match the release of biochemical stimulation cues and corresponding tissue formation. For interesting research on regulatory bioscaffolds and the achievement of concentration gradients, multidisciplines, such as physics and biology, should be integrated to deeply imitate the in vivo cell microenvironments.

3.5. Synergistic stimulations for cell growth and tissue formation

Synergistic stimulation refers to the regulation for cell behaviors by combining at least two stimulants, including biophysical and biochemical stimulation cues [135]. The cellular microenvironments in the body are complex, and they are simultaneously stimulated by various stimulants. Existing regulatory methods have been reported in recent years, and we have a general understanding of the mechanism underlying regulation of cell behaviors under single stimulation. However, the mechanism underlying regulation of cell behaviors under multiple stimulations is still unclear. To provide suitable in vitro or in vivo stimulating microenvironments to promote tissue and organ formation, it is important to research the corresponding mechanism underlying regulation of synergistic stimulations. Common synergistic stimulations include multiple biophysical, multiple biochemical, and chemical and mechanical stimulations. As mentioned before, biophysical stimulation cues include surface topology, surface microhardness, external force and strain, and non-contact-dependent factors (light, ultrasound, electricity, and magnetic stimulations). Biochemical stimulation cues include biochemical factors, such as cells growth factors, cellular proteins, kinases, and synthetic materials (and their concentrations). Synergistic stimulations are regulated by combining regulatory bioscaffolds with research methods mentioned before. Because the regulation of multiple biochemical factors by bioscaffolds is similarly consistent with single biochemical-factor regulation, synergistic stimulations mainly focus on two aspects, multiple biophysical stimulations, and stimulation by biophysical and biochemical stimulation cues.

3.5.1. Multiple biophysical stimulations

Multiple biophysical stimulations comprehensively regulate cell behaviors by combining at least two stimulation cues from surface topology, surface microhardness, external force and strain, and non-contact-dependent factors. Methods of providing a single stimulation to regulate cell growth and tissue formation have been extensively discussed previously. This section mainly discusses synergistic regulation of cell behaviors by 3D-bioprinted bioscaffolds on the basis of six points: (i) surface topology and external force or strain, (ii) surface topology and external fields, (iii) surface microhardness and surface topology, (iv) surface microhardness and external force or strain, (v) surface microhardness and external fields, and (vi) external fields and external force or strain.

The topological structure of bioscaffolds is usually involved in multiple biophysical stimulations. Matsugaki et al. [136] used extrusion-based 3D bioprinting to fabricate collagen bioscaffolds with different topologies, including oriented scaffolds and scaffolds with random orientation (Fig. 9A–i). Stimulation of shear stress was created by fluid flow. Primary osteocytes showed obvious enhanced directional distribution under the synergetic stimulation by surface topology and shear stress (Fig. 9A–ii). Synergetic stimulation by other external forces and surface topology can also be applied by adjusting the fluid state. In addition, 3D-bioprinted microchannel bioscaffolds pave the way for controllable strain regulation. A corresponding synergetic regulation can be achieved by fabricating microchannel bioscaffolds with surface topology and perfusing fluids. Aliaabouzar et al. [137] investigated the synergistic effects of ultrasound stimulation and surface topology. They used UV-assisted 3D bioprinting to fabricate nonporous hydrogel bioscaffolds and porous bioscaffolds with square or hexagonal geometry (Fig. 9B–i). Cell-loaded bioscaffolds were placed in an ultrasound

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stimulation. Synergetic stimulation promoted the spread and growth of MSCs (Fig. 9B–ii). Such regulation can also be achieved by embedding non-contact-dependent-factor-responsive materials into bioprinting materials. Xu et al. [138] used extrusion-based 3D bioprinting to fabricate pattern nanocellulose hydrogel bioscaffolds with different lattice sizes on the basis of the synergetic regulation of surface microhardness and surface topology (Fig. 9C). Bioscaffold stiffness was modified by postprinting chemical cross-linking. Within a certain range, fibroblast cell proliferation significantly improved with an increase in surface microhardness and lattice size.

For synergistic stimulation by surface microhardness and external force or strain, many of the design and manufacturing processes are the same as those for synergistic stimulation by surface microhardness and surface topology, except for the additional consideration of external force or strain. Dan et al. [139] investigated the coordinate effects of cyclic stretch and substrate stiffness on endothelial cells and found that stimulation improves the integrity of cell monolayers and promotes monolayer recovery. Synergetic stimulation by surface microhardness and external fields has also been researched. Heo et al. [140] use UV-assisted 3D bioprinting to fabricate conductive hydrogel bioscaffolds (Fig. 9D–i) and determined surface microhardness by the degree of cross-linking. After seeding cells and applying electric fields, cells on bioscaffolds with different surface microhardness values were synergetically regulated. Imaninezhad et al. [141] investigated the same–ii). Similarly, bioprinting inks loaded with conductive materials are used for coordinated regulation of electrical stimulation and external strain. Wang et al. [142] fabricated conductive composite multiple-channel bioscaffolds using inkjet 3D bioprinting with graphene-containing bioinks (Fig. 9E–i). Such regulation can be achieved after additional stimulation by external strain, as discussed before. Pavesi et al. [143] investigated this regulation of cell behaviors and found significantly change in the expression of a few genes, such as connexin43 (Fig. 9E–ii).

3.5.1.1. Discussion of pros and cons. Synergetic stimulation by biological stimulation cues directly regulates the shape of new tissue and affects its functions. Cell adhesion and cell growth directions are initially determined and guided by the bioscaffold surface topology. The cell distribution on bioscaffolds can be regulated by inducing cell migration through surface microhardness. Exerting stress and strain is a way to regulate cell proliferation, cell differentiation, and tissue formation. Because of parameterized implements and wireless real-time controllable operations, non-contact-dependent factors are widely involved in cell regulation, whether direct or indirect, based on transformation into other stimulation cues. 3D topology structures of bioscaffolds can be fabricated by 3D bioprinting. Surface microhardness can be determined by 3D bioprinting processes, such as cross-linking, or external loads, such as external-field-induced surface modification. Cell regulation by an external force is usually achieved by fluid flow, external strain is exerted by microchannel bioscaffolds with the help of fluids, and noncontact regulation can be achieved by applying an external field. It may be difficult to achieve synergetic stimulation by multiple biophysical stimulation cues using only one 3D bioprinting process or technology. Step-by-step preparation by combining multiple processes (extrusion, inkjet and UV) is a feasible approach to fabricating bioscaffolds embedded with regulatory modules. After integrating multiple biophysical stimulations, nearly the entire processes of cell growth and tissue formation can be regulated. Especially, synergetic stimulation by three or more biophysical stimulation cues for regulation of cell behaviors [144] injects new vitality into tissue engineering, although there are few reports on this. In the foreseeable future, multiple biophysical stimulations are expected to play a crucial role in personalized functional tissue reconstruction.

3.5.2. Stimulations by biophysical and biochemical cues

Cell behaviors can be regulated by combining at least two stimulation cues from biophysical and biochemical stimulations. Biochemical stimulations can be selected intuitively. On the basis of whether an external field is involved, research on coordinated regulations based on methodologies mainly focuses on two aspects. One aspect excluding biochemical stimulation includes surface topology, surface microhardness, and external stress or strain. The other excluding biochemical stimulation includes ultrasound, magnetic, electrical, and light stimulations. In addition, a few uncommon but important synergetic stimulations, such as indirect stimulus by conversing stimulation cues and cell co-culture, are discussed.

Song et al. [145] investigated surface topology and biochemical stimulation cues for cell behaviors. Bioscaffolds with macro-/microstructures were fabricated by 3D bioprinting, and a biochemical stimulation cue was introduced by chemical etching of HA. The authors observed a large area of bone regeneration under synergetic stimulation. Studies have also researched surface microhardness and biochemical stimulation cues. A simple method is to design a bioscaffold the same way as with surface microhardness regulation and then to introduce biochemical factors. Another important approach was developed by Shi et al. [146], who cross-linked alginate hydrogels with different concentrations during 3D bioprinting (Fig. 10A–i). Distler et al. [147] fabricated bioscaffolds with tunable stiffness based on ionical and enzymatical dual cross-linking. Synergetic stimulation increased NIH-3T3 and ATDC-5 cell attachment (Fig. 10A–ii). In addition, cells are subject to fluid shear in dynamic culture. Chen et al. [148] analyzed the stimulation by external strain and nerve growth factors. They fabricated cell-loaded bioscaffolds using extrusion-based 3D bioprinting and applied them to a dynamic tensile culture system (Fig. 10B–i). Synergetic stimulation promoted nerve cell proliferation and the formation of aligned, spindled, and stretched shapes (Fig. 10B–ii). Co-stimulation by external stress and biochemical stimulation can likewise be achieved by using a fluid as a biochemical stimulus or by incorporating biochemical stimulation cues into bioscaffolds.

Noncontact factors are also usually involved. Non-invasive ultrasound and lipid-coated microbubbles (MBs) were analyzed by Aliaibouzar et al. [149] (Fig. 10C–i). Cells were seeded on the 3D-bioprinted bioscaffolds. Coordinated treatment of ultrasound and MBs
Fig. 10. Regulation of synergistic stimulation by biophysical and biochemical stimulation cues for cell behaviors. (A) Synergistic stimulation by surface microhardness and biochemical stimulation cues and its effect on cell behaviors: (i) schematic diagram of surface microhardness changed by cross-linking. Reproduced with permission [146]. Copyright 2017, John Wiley and Sons; (ii) increased NIH-3T3 and ATDC5 cell attachment under synergetic stimulation. Reproduced with permission [147]. Copyright 2020, American Chemical Society. (B) Synergistic stimulation by external strain and biochemical stimulation cues and its effect on cell behaviors: (i) cell-loaded bioscaffolds applied to a dynamic tensile culture system and (ii) enhanced nerve cell proliferation and formation of aligned, spindled, and stretched shapes under synergetic stimulation. Reproduced with permission [148]. Copyright 2020, Chen et al. (C) Synergistic stimulation by ultrasound and biochemical stimulation cues and its effect on cell behaviors: (i) schematic diagram of the synergistic stimulation exposure setup and (ii) enhanced cell proliferation under synergetic stimulation. Reproduced with permission [149]. Copyright 2016, Aliaobouzar et al. (D) Synergistic stimulation by electric field and biochemical stimulation cues and its effect on cell behaviors: (i) conductive CNT-incorporated bioscaffolds fabricated by UV-assisted 3D bioprinting and (ii) enhanced neural stem cell proliferation under synergetic stimulation. Reproduced with permission [151]. Copyright 2018, IOP Publishing. (E) Synergistic stimulation by photothermal effect and biochemical stimulation cues and its effect on tissue behaviors: (i) 3D-bioprinted composite bioscaffolds composed of 30CS and ROS and (ii) inhibited tumor tissue formation in vivo under synergetic stimulation. Reproduced with permission [153]. Copyright 2018, Ma et al. ADA, alginate; GEL 7.5%, 7.5% w/v gelatin; mTG, microbial transglutaminase; CNT, carbon nanotube; UV, ultraviolet; 30CS, Fe$_3$O$_4$-CaSiO$_3$; ROS, reactive oxygen species.

significantly promoted cell proliferation (Fig. 10C–ii). Chen et al. [150] investigated the effects of combined quetiapine and magnetic stimulation on cells. Synergetic stimulation increased brain-derived neurotrophic factor and protein kinase levels.

In addition to direct introduction, biochemical stimulation cues can also be integrated into bioscaffolds. Lee et al. [151] investigated electric field and biochemical stimulation cues. CNTs were added to a poly (ethylene glycol) diacrylate (PEGDA) hydrogel, and corresponding bioscaffolds were fabricated using UV-assisted 3D bioprinting (Fig. 10D–i). Synergetic stimulation promoted neural stem cell proliferation (Fig. 10D–ii) and improved neuronal differentiation and maturity. Studies have also investigated co-stimulation of light-responsive bioscaffolds. Imato et al. [152] fabricated bioscaffold materials by incorporating spiropyran into a photoresponsive polymer chain. Biochemical stimulation was adjusted by changing the spiropyran–polymer ratio. Proper UV irradiation and biochemical stimulation significantly inhibited cell adhesion. On the basis of light-responsive materials, this co-stimulation can also be exerted by introducing additional biochemical factors.

Ma et al. [153] investigated indirect stimulation by conversion of stimulation cues. They fabricated iron–calcium silicate composite bioscaffolds using 3D bioprinting (Fig. 10E–i) and analyzed the photothermal effect of iron particles and the catalytic effect of iron ions. Synergistic stimulation inhibited tumor cell growth (Fig. 10E–ii). Indirect stimulation can also be achieved by cell co-culture through cell secretion of biochemical stimulation cues [154]. Choi et al. [79] investigated the effect of fluid stress and cell co-culture. Smooth muscle cells were adhered to the inner surface of bioscaffold microchannels, and endothelial cells were perfused into the microchannels under different flow rates. Co-cultivation promoted endothelial cell-mediated vascularization.

3.5.2.1. Discussion of pros and cons. Compared to multiple biophysical stimulations, synergistic stimulation of biophysical and biochemical stimulation cues achieves more accurate simulation of microenvironments in vivo and diversified regulation of cell behaviors. Biophysical stimulations directly guide cell adhesion, migration, and oriented population and cell morphology. Biochemical stimulations significantly regulate protein secretion and gene expression and enhance cell and tissue functionalization. In addition to direct stimulation cues, there are two ways of regulation, indirect stimulus by conversion of stimulation cues and cell co-culture. The common conversion of stimulation cues involves piezoelectric, photovoltaic, and photothermal effects. Conversion provides a means to integrate the positive points of each stimulation cue. In addition, the body’s microenvironments are complex, and interactions between co-cultured cells lead to new ideas for regulating corresponding cell behaviors.

Recent studies have reported the effects of microsegment materials of bioscaffolds on cell behaviors [155]. This is synergistic regulation of cell behaviors by biophysical and biochemical stimulations. The chemical properties of microsegment materials provide biochemical stimulation cues, while topological structures provide biophysical stimulation cues. This is an interesting research direction because of the possibility of the intervals between materials promoting cell migration and growth. In addition, composite bioscaffolds are involved in this regulation.

3D Bioprinting technologies provide unique manufacturability for fabricating these bioscaffolds, whether embedded with biophysical or biochemical stimulation modules. Especially, extrusion-based 3D bioprinting is adapted for various materials. Inkjet and UV-assisted 3D bioprinting provide high resolution. Coordinated stimulations created by 3D-bioprinted bioscaffolds pave the way for engineering morphological and functional tissues, which is the ultimate direction of tissue engineering research. These approaches are of great importance and significance in clinical application.

3.5.3. Application prospects of synergistic stimulations

Although tissue engineering research has made significant progress, it is still limited to the simple fabrication of tissues and organs from a shape point of view. Functional tissue fabrication is still in its early stages. Cells in vivo are in complex microenvironments, and many stimulating factors simultaneously regulate the growth process. Synergetic stimulations can simulate cellular niches. In addition, there are many types of stimulation cues, contributing to numerous different synergetic regulation methods. Therefore, regulation of cell behaviors is of significant importance to ensure functional engineered tissues. However, research still focuses on single stimulation schemes. There are few reports on three or more stimulations. There are difficulties in researching the preparation process of biochemical reagents, the forming process of surface structures, the designs of external stress and strain bioscaffolds, and the build methods of non-contact-dependent biophysical stimulations.

3D bioprinting promotes high-precision fabrication of tissue engineering bioscaffolds, which provides a powerful tool for coordinated regulation of multiple stimulations. Application of the 3D-bioprinted bioscaffold technique to synergistically regulate cell growth and tissue formation is promising. First, compared to conventional approaches, bioscaffold-based regulation provides better spatial structure and growth microenvironments for cell propagation. In addition, the properties of engineering tissues, like fabrication, biomechanics, and biochemistries, regulated by 3D bioprinting approaches are better compared to conventional approaches. For example, blood vessels have a hierarchical organization, and blood cells can easily climb and grow on 3D-bioprinted bioscaffolds. In addition, the internal spatial structure and its functionalization, including the initial laminar structure of blood vessels and the tunica intima, media, and externa, can be controlled by 3D bioprinting processes and subsequent regulation methods. Second, some prerequisites are met. Especially, ultrasound and magnetic stimulation and biochemical stimulation cues have been widely applied in medicine. Bioreactors can provide regulation of cell behaviors. These stimulation cues and fluid-driven external stress can be easily achieved by nonprofessionals. Other complex stimulation cues, such as bioscaffold topology, require communication, and cooperation between engineering personnel and medical practitioners. Third, more and more regulatory measures for medical implant bioscaffolds, such as
4. Outlook and conclusion

The advantages of different cell regulation methods promote the individualized and biomimetic regulation of cell behaviors by combining the knowledge of computer science, mechanics, materials science, 3D bioprinting, and biomedical engineering. In addition, the advantages and disadvantages of these methods facilitate the development of regulation methods for cell behaviors in vitro (Table 2).

4.1. Outlook

The complex structure of in vivo ECMs provides a good microenvironment for cell growth. Conventional manufacturing methods cannot accurately simulate cell microenvironments. This problem can be effectively solved by the development of new materials and processes of 3D bioprinting. Currently, the regulation of cell behaviors is not limited to biochemical reagents. Medium concentration and biophysical stimulation cues, such as external force, strain, and electric field, are emerging as regulation methods for cell behaviors. However, there are challenges in bioprinting and regulation processes, especially the fabrication of high-precision cell-loaded bioscaffolds and accurate regulation of cell growth. In addition, image technologies, such as µCT, may be used in bioscaffold design and tissue analysis.

4.1.1. 3D bioprinting technologies

Challenges in 3D bioprinting include bioink materials, bioprinting speed and resolution, and large-scale tissue preparation. Because of cellular requirements of biocompatibility, nutrients, and immunity, bioinks are limited to natural materials or acellular ECMs. Low-speed fabrication cannot meet the requirements of mass production. So far, high resolution bioprinting limited to 2D or simple 3D structures cannot achieve high-precision bioprinting of multilayered or complex large-scale tissues. In addition, the compatibility between the degradation characteristics of bioscaffolds and cell growth needs to be considered. To obtain biomimetic bioscaffolds, composite biomaterials and 3D bioprinting processes, including process parameters and forming accuracy, need to be further studied and optimized. On the basis of the principles of bioprinting, such as extrusion-based, inkjet, and UV-assisted 3D bioprinting, personalized bioprinting bioscaffolds should be fabricated to prepare target tissue structures in the future.

4.1.2. Cell regulation by bioscaffolds

With regard to regulation, regulation methods, and bioscaffold fabrication are the main difficulties. Regulation methods include biochemical and physical stimulations of cells. The fabrication of regulatory bioscaffolds involves interdisciplinary technologies, including computer-aided design and manufacturing (CAD/CAM), materials science, control science, chemistry, and biomedical engineering. Many factors regulate cell growth in vivo. Current regulations for cell growth focus on a single regulatory factor. Attention should be paid to synergistic regulation of cell behaviors by various stimulation cues. In addition, the match between the degradation performance of bioscaffolds and the tissue regeneration rate should be studied in the future.

4.1.3. Integration of bioscaffolds loaded with cells and regulation modules

Structures integrated with bioscaffold modules and regulatory modules can be applied to the research on the mechanism underlying regulation of cell growth and tissue formation. There are two ways to integrate modules:

- Bioscaffold modules and control modules are prepared separately and then integrated on the basis of Boolean operation. Compared to the challenges in preparing corresponding modules, the outstanding challenges are integrated modules, assembly accuracy, and

Table 2

| Methods                        | Mechanisms                                      | Advantages                                      | Disadvantages                                      |
|--------------------------------|-------------------------------------------------|-------------------------------------------------|---------------------------------------------------|
| Surface topography modifying   | Optimizing the surface topography of microchannels at nano-scale | - Inducing cell adhesion [156]                  | Complex preparation processes [156–160]             |
|                                |                                                  | - No other materials were introduced [156]       | High resolution and accuracy requirements [161]    |
|                                |                                                  | - High efficiency [157]                         | Considered biocompatibility and degradation characteristics of materials [164] |
| Surface coating                | Introducing other favorable materials on the surface of microchannels | - Inducing cell adhesion [162]                  | Complete biocompatibility and degradation characteristics of materials [164] |
|                                |                                                  | - Accurate control [163]                        | Complete biocomposition processes [168]            |
| External force applying        | Changing the morphology of cells and stimulating secretion [165] | - Improved proliferation, migration, differentiation and alignment of cells [166,167] | Complex bioprinting processes [168]               |
| Strain applying                | Influencing the direction of cell growth and stimulating secretion [166] | - Easily-controlling, improved alignment and differentiation of cells [170,172] | Complex bioprinting processes [168]               |
| Medium                         | Modulating the speed of cell differentiation and reprogramming [173] | - Controllable concentration gradient [168]    | Thoughtful degradation characteristics [168]       |
| Concentration changing         |                                                  |                                                  | Complex preparation and regulation processes [169] |
| Surface microhardness changing | Inducing cell migration [174]                   | - Controllable patterning [175] and no chemical components introduced [176] | Limited researches [176]                          |
| Biochemical cues adding        | Inducing cell differentiation and reprogramming [177] | - Synergistic regulation of multiple chemical factors [176] | Introducing other materials [177]                  |
| Light irradiating              | Inducing cell proliferation and migration [178] | - No contact on cells [179]                     | Considerable biocompatibility [178]                |
| Ultrasound exerting            | Inducing cell proliferation and differentiation [181,182] | - Killing bacteria [179]                        | Limited materials [179]                            |
| Electricity stimulating        | Activating intracellular signaling pathways, and influencing microenvironments [184] | No contact on cells and no modify on scaffold surfaces [185] | Required high-precision control [180]              |
| Magnetic stimulating           | Enhanced intracellular delivery and changed cell stiffness [187] | - Precise cellular regulation by the combination of specific materials/structures and electricity stimulating [185, 186] | Low-precision control [180]                        |

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introduction of pollution. 3D bioprinting technologies, conventional mold preparation and demolding technologies, Lego-like modular building, and assembly technologies may be combined to fabricate and reassemble bioscaffolds.  

- Integrated 3D models with bioscaffold modules and control modules are directly fabricated. The structures are prepared by 3D bioprinting technologies, which have higher requirements for their preparation processes because of the increased complexity of print models.

4.1.4. Regulation of cell behaviors in vitro and in vivo

Due to material properties of bioscaffolds and rheological properties of the culture medium, there are differences between stimulations delivered to cells and externally applied stimulations. Therefore, it is necessary to evaluate the relationship between the actual stimulations acting on cells and corresponding cell behaviors. In addition, cell growth factor secretion, protein expression, related gene phenotypes, and immunity should be evaluated and quantified under single and synergetic stimulations. There are no clear test methods or standards for implanted bioscaffolds. Current studies are based on animal experiments, but animals and humans respond differently to bioscaffolds. On the basis of clarifying the mechanism underlying regulation of cell behaviors and the characteristics of bioscaffold degradation, bioscaffolds integrated with regulatory modules are expected to achieve remote regulation of in vivo cell growth. For large-scale application of bioscaffolds in a clinical setting, the evaluation criteria of implantable bioscaffolds should be investigated and the tissues prepared should be evaluated for clinical application.

4.2. Conclusion

To accurately simulate the growth microenvironments of cells and study the mechanism underlying cell growth and tissue formation, 3D bioprinting and regulation technologies have been applied to the fabrication of bioscaffolds embedded with regulatory modules. Tissue reconstruction has undergone three stages of development: (i) free cell growth on bioscaffolds, (ii) cell growth under the action of a single regulatory factor, and (iii) cell growth under the action of multiple regulatory factors. First, conventional bioscaffold fabrication is based on fabrication of bioscaffolds by 3D bioprinting and cell adhesion to grow into tissues without additional induction. However, the tissues prepared are usually quite different from requirements. To reconstruct satisfac-

CRediT authorship contribution statement

Pengju Wang: Writing - original draft. Yazhou Sun: Supervision. Xiaquan Shi: Giving Advice about presentation of the manuscript. Huixing Shen: Searching for the references. Haohao Ning: Searching for the references. Haitao Liu: Giving Advice about organization and presentation of the manuscript.

Declaration of competing interest

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

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