Biomaterials for stem cell engineering and biomanufacturing

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Abstract

Recent years have witnessed the expansion of tissue failures and diseases. The upsurge of regenerative medicine converges the sight onto stem cell-biomaterial based therapy. Tissue engineering and regenerative medicine proposes the strategy of constructing spatially, mechanically, chemically and biologically designed biomaterials for stem cells to grow and differentiate. Therefore, this paper summarized the basic properties of embryonic stem cells (ESCs), induced pluripotent stem cells (iPSCs) and adult stem cells. The properties of frequently used biomaterials were also described in terms of natural and synthetic origins. Particularly, the combination of stem cells and biomaterials for tissue repair applications was reviewed in terms of nervous, cardiovascular, pancreatic, hematopoietic and musculoskeletal system. Finally, stem-cell-related biomanufacturing was envisioned and the novel biofabrication technologies were discussed, enlightening a promising route for the future advancement of large-scale stem cell-biomaterial based therapeutic manufacturing.

1. Introduction

Stem cell, a particular cell type with self-renewing ability, has the potential to differentiate into different cell lineages and has gradually become the most versatile and valuable cell source for organ transplantation, disease treatment and cosmetology. Technically, the term stem cell can be divided into three specific categories: embryonic stem cell, induced pluripotent stem cell and adult stem cell, depending on the cell's developmental potential. All of them play a crucial role in tissue regeneration. However, some therapeutic defects such as low efficiency of differentiation protocols, high risk of teratoma formation and poor immune compatibility [1] remain unsolved when the stem cells are used alone. Therefore, a new reliable approach has been applied - the introduction of advanced biomaterials.

Biomaterials are usually designed to possess appropriate biochemical and biophysical properties - including benign molecular compatibility, high porosity and suitable mechanical strength - mimicking the microenvironment of natural extracellular matrix (ECM) [2]. Such an artificial environment may either serve as a bio-adhesive surface for 2D cell culture, for example hydrogel microarrays used for supporting stem cell attachment and proliferation [3], or as 3D scaffold for a specific cell type to interact with and to control their function, thereby inducing the multi-spatial and temporal cellular processes of tissue formation and regeneration. Compared to the conventional cell type-specific biomaterial [4], the novel stem cell-interacting biomaterial is designed to be capable of meeting the need of diverse cell types due to the presence of bioactive cues in the biomaterials [5].

Almost 140 different combinations between stem cells and biomaterials are reported in recent studies, and the efficiency and properties of the cell-material interactions included morphology, vitality, cytotoxicity, apoptosis and proliferation [6]. As depicted in Fig. 1, the ultimate goal of the study is to identify materials that can regulate cell function as well as to find the appropriate biomaterial-stem cell combination for human body.

In this review we firstly outlined the various properties of stem cells and biomaterials, respectively. We then summarized the innovative application of biomaterial-stem cell interactions into clinical therapy for tissue and organ regeneration.

2. Stem cell

The primary merit of stem cells is their infinite proliferation competence. Due to their distinct proliferative properties, stem cell has also been referred as an “undifferentiated” cell type. Whether the stem cells remain self-renewal or turn into differentiated functional cells is mainly determined by the intrinsic state or the microenvironment in which
stem cells reside—niche [7]. Naturally, stem cell keeps the balance between quiescence and activation. Considering the process of division, it can be categorized into asymmetric division (a retained stem cell and a daughter cell) and symmetric division (two daughter cells). Stem cells can be specified as **totipotent stem cells** (capable of producing all cell types of body), **pluripotent stem cells** (capable of producing all cells of the embryo), **multipotent stem cells** (capable of producing cells impacted by the microenvironment) and **unipotent stem cells** (capable of producing only one or two types of cells).

In mammals, only zygotes and spores are totipotent, and they will develop into embryonic lineages and extraembryonic lineages. Extraembryonic cells form the placenta, whereas the inner cell mass (ICM) becomes pluripotent, possessing the capacity to give rise to all cell types of the body. When the ICM is isolated and cultured in vitro, we can formulate **embryonic stem cells** (ESCs) that are able to generate every cell type within the human body [8].

While pluripotent stem cells expand and differentiate, part of them are maintained locally and have the ability to either self-renew or generate specialized cell types for a confined tissue, making them multipotent stem cells. Examples include neural stem cells, hematopoietic stem cell and mammary stem cell etc. [9–11], all of which play important roles in both organogenesis and tissue regeneration. In general, most **adult stem cells** (ASCs) are multipotent and have limited potency and finite periods of regeneration. ASCs are derived from patient or their parent without ethical issues and are widely used for therapy such as leukemia and radiotherapy [12,13]. Unlike the pluripotent and multipotent stem cells, unipotent stem cells have the lowest differentiation potential along only one lineage, however, the fact that adult unipotent germline stem cells can give rise to reproducible germline-derived pluripotent stem cells [14], addresses more potential to the unipotent stem cells.

At the beginning of human developmental studies, researchers used cells from teratocarcinomas, a cancer line derived from germ cells [15]. The problems, including out-of-control differentiation into multiple cell types, called for a more feasible way to find tractable model for studying human cells and disease in vitro. In following years, scientists demonstrated that by using different transcription factors (e.g. Oct 3/4, Sox 2, Klf4 and c-Myc), differentiated somatic cells could be reprogrammed to become induced pluripotent stem cell (iPSC) [16]. These iPSCs are reset to the pluripotent state primed for cloning and medical treatment similar to that of the ESCs. Therefore, the complete pluripotency of those reprogrammed cells broadened the source of stem cells and promised the future of cell therapy.

A brief summary of ESC, ASC, and iPSC are described in Table 1.

3. **Biomaterial for stem cell culture**

Biomaterials serve as non-viable materials in medicine to repair malfunctioned tissues and organs. The past five decades have witnessed tremendous growth in biomaterial science and engineering as a result of vast investment in developing new products. Biomaterials comprise of two species: natural and synthetic materials. As discussed earlier, a specific environment is required for stem cell survival. Therefore, to mimic the in vivo microenvironment, biomaterials open up a new avenue for regulating stem cell fate via cell-matrix interactions. Biomaterial scaffolds can provide cell adhesion sites and maintain the merits of stem cells. In contrast to traditional 2D culture, the novel 3D biomaterial scaffolds construct a more satisfactory microenvironment for stem cells by including both chemical and physical signals across the ECM. Upon well-designed configuration, scaffolds can directly regulate cell signaling and trigger lineage-specific differentiation of stem cells by chemical cues or cell-matrix interactions [24].

With the growing interest in utilizing biomaterial-based approaches, the properties of the biomaterials were found to affect stem cell lineage specification. Hence, surface, mechanical, electrical, electrostrictional, morphological and chemical properties must be precisely considered when designing a new scaffold [25]. After elaborate selections, the cell adhesion, cell transportation, cell differentiation and matrix
organization can be modulated to direct stem cell differentiation. Table 2 summarized typical biomaterials for stem cell culture and the detailed properties of each category will be unfolded in the following part.

3.1. Natural biomaterials

With the goal of mimicking the 3D ECM to regulate stem cell behavior, certain natural biomaterials have been adopted to support stem cell proliferation and differentiation, including collagen, gelatin, hyaluronic acid hydrogels, fibrin, glycosaminoglycans (GAGs), alginate, matrigel, silk and hydroxyapatite (HA), etc. These materials exhibit specific advantages, including similar mechanical and adhesive properties as the natural ECM, while batch variability, short degradation period, difficulty in purification and quality control make up the main disadvantages of these materials.

Collagen, present in all connective tissue, acts as main component of ECM with superior biocompatibility, due to the fact that collagen-derived acellular ECM would not cause serious adverse immune responses. Of note, the biodegradability of collagen makes it a better choice in skin tissue restoration because of its high rate of degradation. While it is the most abundant protein in animals, collagen is difficult to obtain for research and clinical treatment, leading to the production of recombinant collagen for unlimited supply. Moreover, collagen-based biomaterials are now used for cartilage regeneration treating osteochondral defects and cornea defects.

Extracted from cartilage, hyaluronic acid hydrogels (also known as hyaluronan) serve as a native component and play an essential role in cartilage homeostasis and biomechanical integrity, including morphogenesis, proliferation, cellular signaling and wound repair. Hyaluronic acid hydrogels has been used in tissue repair and regeneration as well as adriamycin-induced cytotoxicity prevention by forming a bioartificial stem cell niche. Hyaluronic acid hydrogels were also photo-crosslinked for the application in chondrogenesis where almost all of the encapsulated mesenchymal stem cells (MSCs) survived.

Gelatin, derived from collagen, is now widely employed as a scaffold material for cartilage tissue engineering due to its biocompatibility, biodegradability and ability to form hydrogels. Gelatin can be functionalized with unsaturated methacrylamide to create covalently bound hydrogels for encapsulating stem cells. Of note, gelatin presents a better performance in biomechanical and biochemical properties when compared with other frequently-used hydrogels like alginate and agarose.

Matrigel is a soluble basement membrane extract from Engelbreth-Holm-Swarm (EHS) murine sarcoma. It is composed of laminin, type IV collagen, nestin, heparin sulfate glycoprotein, as well as growth factors and matrix metalloproteinases. At room temperature, matrigel polymerizes to form a biologically active 3D matrix, which mimics the structure, composition, physical properties and functions of the cell basement membrane in vivo, benefitting the culture and differentiation of cells in vitro.

Another classic tissue-derived biomaterial scaffold is made of fibrin, which presents superior properties for providing a microenvironment for stem cells. For instance, nerve growth factor β-NGF was covalently incorporated with fibrin scaffold to produce neurons and oligodendrocytes. However, plasmin inhibitor had to be co-operated to avoid unexpected degradation of the 3D scaffold caused by the ESCs.

3.2. Synthetic biomaterials

Although natural biomaterials have favored biocompatibility and self-existing biosignals, the frail mechanical strength and difficulty in modification limit their broader applications. To overcome these obstacles, synthetic scaffolds have become a solution. As a designed
provide sufficient stiffness for the bone recovery device, hydroxyapatite-based calcium phosphate and bioactive glasses are usually added to form a composite for optimized osteo-fate directing [52,53].

The addition of bioceramics can be also used to enhance the porosity of a polymer implant for improved nutrition and waste transport throughout the scaffolds by introducing micro-/nanoscale voids characteristic of the ceramics [54]. However, the use of a ceramic scaffold can be sensitive. For example, while calcium phosphate cements possessed a unique injectable property and brilliant biocompatibility in vivo, the release of phosphate from an unmodified scaffold lowered medium pH and hampered cell proliferation in vitro [55]. By sintering the scaffolds at a very high temperature, Link et al. [55] altered the calcium phosphate content to a more stable form and the resulting scaffolds improved the specification of osteoblast-like cells in vitro. These results indicated that the physiochemical state of bioceramics played a crucial part in cytotoxicity and both in vitro and in vivo performance needed to be taken into account to comprehensively understand the function of bioceramics for clinical application.

### 3.2.3. Synthetic metals

Titanium, titanium alloys, stainless steels and cobalt alloys are the commonly used metals that contribute to bone regeneration, especially for orthopedic and dental treatments. Stainless steel implants serve as the most accepted materials because of easy procurement; however, it is hard to control the cell-metal interactions on the surface of the materials due to non-specific protein adsorption and cell adhesion, which leads to sub-optimal integration with the host tissue. A recent study aimed to overcome these deficiencies by covalently tethering adhesive peptides to functionalized stainless steel to control stem cell attachment [56]. Another study for improving the biocompatibility of stainless steels by utilizing ZrO2 and SiO2/ZrO2 coating showed that the proliferation of stem cells was dependent on stainless steel scaffold surface properties [57]. Titanium and tantalum demonstrated satisfactory biocompatibility, anticorrosion and excellent mechanical properties which could improve MSC multilineage differentiation in vitro by providing adequate plots for cell adhesion in 3D porous scaffolds [58]. In addition to increasing cell viability, titanium and tantalum could also conserve the immunophenotypic features of MSCs [59]. Moreover, a tantalum film may be deposited onto titanium alloy (Ti6AlV) by filtered cathodic vacuum arc deposition (FCVAD) to enhance the mechanical and anticorrosion properties as well as cytocompatibility for mammalian bone MSCs [60]. Another innovative study exploited a two-layer coating comprised of a tantalum layer and a polymer-titanium hybrid layer on medical devices to enhance the biocompatibility for MSCs [61]. These results support that the titanium, tantalum, and their alloys are suitable for the biomaterial scaffolds.

While metallic biomaterials stand out in compressive strength and fatigue resistance, potential risks exist for current applications include toxic ion release and non-degradable fixtures. Promisingly, magnesium (Mg) and its alloys could serve as a novel substrate to settle these issues. The elastic modulus and compressive yield strength of Mg is similar to that of natural bone, so it can provide an imitative physical microenvironment for cell culture. Furthermore, the magnesium cation is one of the natural trace elements in human body and thus it is relatively harmless upon degradation [62,63]. However, several parameters for 3D scaffold design in MSC differentiation need to be investigated in detail, and more efforts are required to unveil the complex mechanism of the Mg-based scaffolds.

### Table 2

| Types                | Examples                  | Properties                        | Application                      |
|----------------------|---------------------------|-----------------------------------|----------------------------------|
| Natural biomaterial  | Collagen, hyaluronic acid, gelatin, laminin, fibrin | Good biocompatibility, Less immune responsive, Self-existing biosignal, Short degradation period, Poor mechanical strength, Easy modification, Properties can be designed, Good mechanical strength, Poor degradability, Poor tensile property | Cartilage/bone repair, Osteochondral repair, Cornea repair, Nerve regeneration, Coating matrix, All kinds of stem cell culture and tissue repair, Additives in bone tissue engineering |
| Synthetic biomaterial | Polymer: PLA, PLGA, PCL, PEG, PVA, PHEMA, PMMA, Ceramic: HA, TCP, bioactive glass, Metal: Titanium, titanium alloy, stainless steel, cobalt alloy | Good compressive strength, Good fatigue resistance, Non-degradable, Non-bioadhesive | Orthopedic and dental treatment |

Component, the structure and relative mass of a synthetic biomaterial can be controlled at will. Nevertheless, synthetic biomaterials are not consummate for this application since they lack cell adhesion properties and biological signals and thus cannot direct cell fate on their own. Notably, biocompatibility and biodegradability of the synthetic composite frequently acts as the most essential hurdle in stem cell culture, and many studies are being conducted to solve these issues.
3.2.4. Synthetic graphene

Recently, one of the 2D complanate structure biomaterials, graphene, has come into sight. Graphene has plenty of marvelous characteristics such as adequate reaction area, active surface chemistry and functional electrical/thermal conductivity [65,66]. Graphene and its derivatives could serve as a biocompatible and biodegradable pre-concentration platform for MSC growth and osteogenesis, through firm non-covalent binding. The differentiation could be modulated by interactions of growth factors and π–π stacking as well as electrostatic and hydrogen bonding [67]. Interestingly, graphene and its derivatives could also act as ultrasensitive bio-detection materials for immunosensors of the Nanog protein to quantify the pluripotency of stem cells [68]. There have been studies demonstrating the merits of graphene in stem cell culture; nevertheless, we have to admit that the mechanism of stem cell-graphene interactions are barely understood and there is still a demand for further studies to shift the graphene-based stem cell niche from 2D planes to 3D structures.

4. Biomaterials & stem cell applications

Biomaterials play an essential role in the field of tissue engineering and regenerative medicine. The diversity and versatility of these biomaterials have promoted a bloom of development in the field. With the increasing demand for tissue repair, a plethora of biomaterials are modified and exploited in various applications with regard to specific tissue type, as illustrated in Fig. 2 and listed in Table 3. In this section, detailed applications of novel biomaterials to direct stem cells into different lineages for tissue repair will be discussed accordingly.

4.1. Nerve tissue

Neurological diseases are a severe threat to human health: about 46.8 million people suffer from dementia and Alzheimer’s Disease (AD) and over 10 million patients were diagnosed with Parkinson’s Disease (PD) [95,96]. However, lacking the ability of spontaneous regeneration, the repair/reconstruction of nervous system needs to be associated with therapeutic intervention, in which nerve tissue engineering scaffolds are designed to direct the formation of new neurons from stem cells in vitro, for the replacement of damaged or dysfunctional nerve tissue.

Natural biomaterials, such as collagen, alginate, gelatin, chitin, elastin and hyaluronic acid, are widely used scaffold matrices for nerve tissue repair due to their excellent biocompatibility and cell-adhesive properties. For example, aligned and oriented 3D collagen hydrogels, prepared by mechanical strain stimulation after gelation, were proven to enhance neuronal growth compared to 2D culture and proven to direct an elongated morphology of both single-cell neuron and nerve tissue explant [97]. Scaffolds compatible with stem cells are favored as they hold the potential of directing differentiation of stem cells into any neural and glial cell type that are required in the treatment. Winter et al. [69] devised transplantable tubular hydrogel-collagen micro-columns, mimicking glial tubes, to guide nerve progenitor cells for central nervous system regeneration. Moreover, collagen-based scaffolds have been shown to preserve in a hypothermic condition for up to 4 days without affecting cell viability and metabolic activity, indicating a promising feasibility of clinical and commercial translation of this system [98]. Nonetheless, natural scaffolds suffer from relatively-weak mechanical properties, and thus synthetic biomaterials were introduced into natural scaffold. For example, poly-(L-lactic acid)-co-poly-(3-caprolactone) was mixed with collagen to form a nerve tissue-targeted nanofibrous scaffold that enhanced the differentiation of MSCs into neurons [70]. The incorporation of hyaluronic acid doped-poly-(3,4-ethylenedioxythiophene) (PEDOT-HA) nanoparticles into chitosan/
Table 3
Biomaterials used in different tissue types for specific functions.

| Tissue type            | Biomaterials                  | Stem cell types | Applications                                       | Ref. |
|------------------------|-------------------------------|-----------------|----------------------------------------------------|------|
| Nerve tissue           | Collagen                      | Nerve progenitor cell | Central nerve regeneration                          | [69] |
|                        | Poly (l-lactic)-co-poly-(3 caprolactone) | MSC              | To enhance differentiation into neurons             | [70] |
|                        | PLGA                          | KT 98           | To increase proliferation rate                      | [71] |
|                        | PEDOT-PA                      | PC12            | To enhance synapse growth                           | [72] |
|                        | Collagen-fibrin               | Schwann cell     | Peripheral nerve regeneration                       | [73] |
| Cardiovascular tissue  | Alginate                      | MSC             | To encapsulate cells                                | [74] |
|                        | RGD peptide                   | MSC             | To promote cell attachment                          | [74] |
|                        | GFOGER hydrogel               | Cardiac progenitor cell | To induce cardiomyocyte differentiation            | [75] |
| Pancreatic islet        | Alginate, agarose, PEG, PLGA, PLLA | Any iPSC, ESC    | To encapsulate cells                                | [76-79] |
| Hematopoietic system    | Tropoelastin                  | HSC             | To enhance proliferation                            | [80] |
|                        | Collagen                      | HSC, MSC         | To modulate matrix elasticity                       | [81] |
|                        | Collagen, PEG, PCL            | HSC, MSC         | To support co-culture of HSC and MSC                | [82-84] |

gelatin (CS/Gel) scaffold led to better mechanical properties and improved substrate conductivity, enhancing the direction of synapse growth [72].

The expanding demand for nerve injury treatment poses an increase in the study of synthetic scaffolds, as they can overcome the problems encountered by traditional allografts and autografts, including limited availability and immune rejection. Synthetic scaffolds may be rationally designed and engineered with supported biodegradability, biocompatibility, porosity, hydrophilicity and mechanical strength for the injured site. The availability of different synthetic biomaterials with different functions [99], including poly l-lactic acid (PLLA), PLA, PCL, poly-glycerol sebacate, PLGA, poly-3-hydroxybutyrate (PHB), polyamide, polydioxanone, poly-e-caprolactone-co-ethyl ethylene phosphate (PCLEEP), poly-D, l-lactide-co-caprolactone (PDLLCL), PVA, poly acrylonitrile-co-methylacrylate (PAN-MA) and a copolymer of methyl methacrylate and acrylic acid (PMMAAA), has enabled the incorporation of various neurotropic factors for neuronal repair. Of note, a conduit-based technology, a specific scaffold that could provide an appropriate microenvironment for peripheral nerve regeneration over long nerve gaps or in the case of large-sized nerves, has been an active research area in both fundamental studies and clinical trials. In this nerve conduit, the proximal and distal nerve stumps are plugged into the two ends of the tube respectively, and axons are formed from the proximal nerve and eventually germinated maturely at the end of the distal nerve. Due to the restricted growth by encapsulated conduit, the probability of immune rejection and neumoramic formation will decrease. For example, a gelatin cryogel 3D conduit was shown to recover the transected peripheral nerve and reduce the risk of scarring other tissues [100]. A hybrid structure guidance conduit by wrapping PLGA micro-fiber bundles in a flat micro/nanostructured PLGA membrane was also shown to increase cell proliferation and promote neurite outgrowth [71]. In another in vivo test [73], collagen-fibrin conduit rods containing Schwann cells increased axonal differentiation in the midsection and distal part of the injured nerve after 4 weeks, providing strong proof of peripheral nerve regeneration using conduit biomaterials.

4.2. Cardiovascular tissue

Cardiovascular disease is a leading cause of death in the world. Patients suffering from ischemic heart disease including myocardial infarction (MI), cardiac arrests and stable/unstable angina, increase significantly along with the progress of tobacco, unhealthy living habits, overdrinking and other detrimental factors. Tissue engineering serves as a reliable option to cure these disorders by exploiting bone marrow-derived stem cells, ESCs, parthenogenetic stem cells and iPSCs. While direct injection of cardiac stem cells have achieved some progress in remuscularization of the heart, difficulties in increasing cell survival and efficient tissue integration [101] call for functional biomaterials to facilitate heart regeneration. The ideal scaffolds for cardiac tissue should follow the nature of ECM and porous architecture for musculuarization and vascularization. The porous 3D architecture provides attachment sites for stem cells, promoting the interactions between implanted cells and the host tissue for vascularization [102].

The earliest attempt to retrieve normal heart tissue in vitro was attempted in 1994 by culturing chicken embryonic heart cells in collagen I [103]. Aftewords, multiple natural (matrigel, collagen, fibrin, alginate, hyaluronan, chitosan etc.) and synthetic (peptide-ampiphile nanofibers, PEG hydrogels etc.) biomaterials were developed for treating injured heart [104]. Even without cells, alginate [105], fibrin [106], methycellulose [107] and hyaluronic acid [108] themselves were reported to restore heart function. Incorporation of follistatin-like 1 into a collagen scaffold stimulated de novo cardiomyogenesis after MI in vivo [109], further indicating that these acellular scaffolds could direct cardiac fate and become a potential therapy for cardiac repair.

Considering the complexity of heart tissue, stem cell-based scaffold is more efficient and attractive, and the field has become flooded by the combination of hPSC-derived cardiomyocytes (CMs) and biomaterials. Hydrogels were often applied for cardiac repair. Particularly, alginate hydrogels were used to encapsulate stem cells for CM differentiation with enhanced cell attachment and survival. Yu et al. [74] produced Arg-Gly-Asp (RGD, major recognition receptor from adhesive ECM, blood and cell surface proteins) modified alginate microspheres to encapsulate MSCs for cardiac regeneration. While both the encapsulated MSCs and CMs alone could induce similar angiogenesis, the former showed long-term survival in the infarcted area. A study of ECM-derived GFOGER hydrogels mimicking collagen adhesive site revealed that induced cardiomyogenesis of cardiac progenitor cells was accompanied by the reduction of reparative growth factor release in vitro [75]. However, these hydrogels did not rescue cardiac function or reduce reparative growth factor level in rats undergoing ischemia-reperfusion, suggesting different mechanisms between in vivo and in vitro tissue regeneration. Interestingly, 3D PEG hydrogels containing thiosulfate cyanide sulfur transferase (TST) were fabricated to catalyze H2S production for a minimal ischemic effect and reperfusion damage in the implant site while stimulating angiogenesis [110]. The increased proliferation of human cardiac progenitor cells suggested the possibility of endogenous gasotransmitter on cardiac regeneration. Likewise, a plenty of studies aim to identify reliable stem cell-materials grafts for cardiac repair, but a lack of mechanistic understanding of cardiac regeneration slows down the pace in clinic study.

4.3. Pancreatic islet

Diabetes mellitus is one of the major non-communicable diseases characterized by hyperglycemia. It is a serious and chronic disease that could further lead to the dysfunction and damage of many other organs, including eyes, ears, blood vessels, kidneys and nerves. There are three forms of diabetes - Type 1, Type 2 and Gestational diabetes. Type 1 diabetes is mainly caused by the failure of pancreas to produce insulin due to the loss of β cells from an autoimmunity, and therefore, the
regeneration of insulin-producing $\beta$ cells holds the key in Type 1 diabetes treatment. Conventional methods including allo- and xeno-pancreatic islet transplantation often encounter with insufficient donors or severe immune rejection. Alternatively, stem cell-derived pancreatic $\beta$ cells are considered as an unlimited source, and autologous patient-specific iPSC-derived $\beta$ cells are supposed to generate minor immunoreaction. Furthermore, semi-permeable porous scaffolds provide better seeding and a protective microenvironment for pancreatic cells/tissues. In conclusion, incorporation of pancreatic progenitor cells into biomaterials for grafting without immunosuppression becomes an ideal method in future diabetes treatment.

Encapsulation of islet cells in biomaterials, in which the encapsulated cells were isolated from the immune system, was widely investigated to avoid auto-immune attack. Both Bratlie et al. [103] and Scharp et al. [102] reviewed structural approaches for islet encapsulation, and summarized the prevalent use of alginate, agarose, tissue-engineered chondrocytes, polyacrylates and PEG to produce macro-/micro-/nano-devices, conformal coating and layer-by-layer coating for encapsulation. Upon linked with directing factors, these scaffolds could specify pancreatic fate. A recent study invented an activin A-grafted gelatin-PLGA nanoparticle (PLGA NP) scaffold to induce endoderm formation from iPSCs in the presence of 2-(4-morpholinyl)-8-phenyl-4H-1-benzopyran-4-one (LY294002) prior to pancreatic differentiation [78]. This pioneering work proposed a guidance for the production of pancreatic cells from iPSCs in polymer scaffolds. For example, stem cell-derived $\beta$ cells were encapsulated in TMTD (triazole-thiomorpholine dioxide) modified alginites for in vivo glycemic control for 174 days without exploiting immunosuppression [79].

Numerous pre-clinical trials using encapsulated stem cell-derived human $\beta$ cells were performed in non-immunosuppressed diabetic animal models, yet few of them were successfully confirmed in clinical human patients for the past three decades [111]. Scharp and Marchetti [76] owed the difficulties in encapsulated islet clinical therapy to the following cues: a) poor translation of treatment results obtained from rodents to human, b) acute loss of encapsulated cells due to hypoxia, c) transplantation antigen sensitization to the implant recipient, d) not enough cell quantities implanted. While the first reason could only be solved upon plenty of clinical human trials, the other three could be overcome by proper design of the scaffold/cell composite. Most of the unsuccessful islet implantations were caused by the low survival rate of the cells due to poor vascularization around the implanted site [112]. Proper vascularization between encapsulated cells and host tissue can be achieved by the directing function of engineered porous tunnels, or by co-encapsulation of vascular cells [113,114]. Kaufman-Francis et al. [115] co-cultured pancreatic islets, human umbilical vein endothelial cells (HUVECs) and human fibroblasts on highly porous PLLA/PLGA scaffolds, leading to an increased insulin secretion by 50%, an enhanced islet survival and the formation of functional tube-like endothelial vessels.

4.4. Hematopoietic system

The hematopoietic system, comprised of bone marrow, spleen, tonsils and lymph nodes, serves as a blood producer. Hematopoietic stem cells (HSCs), residing in the bone marrow (main source) and cord-blood, hold the proliferation and immune potential to generate all cellular components to maintain the blood circulatory function. This renders HSCs as an essential core for the transplantation treatment of hematopoietic disorders and cancers [116]. However, the fate of a long-term HSC includes four stages: self-renewal, differentiation, emigration from bone marrow to blood and apoptosis [116]. Therefore the organization of hematopoiesis signaling pathways is rather complicated. Cell to cell, cell to cytokine and cell to ECM interactions, as well as the microenvironment parameters, such as $O_2$ concentration, permeability, transcription factors, cytokines, stability, etc., all devoted to the ex vivo expansion of HSCs [117], shaping great challenges for the design of biomaterials for hematopoietic system therapy.

HSCs were considered strongly related to bone component and Calvi et al. [118] confirmed that osteoblastic cells were the regulatory component in the HSC niche in vivo. In this sense, the ECM properties of HSC were quite similar to that of osteoblasts. Nonetheless, while stiffness was widely studied for bone scaffolds, recent research revealed an emphasized impact of elasticity for the HSC expansion in biomaterials. Hoist et al. [80] found that elimination of elasticity sensing ability of HSCs could neutralize the proliferation enhancement on elastic biomaterials. More recently, Choi et al. [81] fabricated collagen-coated polycrylamide scaffolds and discovered the spreading and morphology of hematopoietic stem and progenitor cell (HSPC) were responsive to the substrate elasticity, validating the important role of matrix biophysical properties in HSC survival. In addition, a 3D structure was proved to be more beneficial for HSC expansion as compared to 2D culture, since higher ratio of surface area to volume provided more efficient cellular interactions [119].

Apart from substrate mechanical properties, mesenchymal stromal cells also played a crucial role in HSC function [120], and therefore attempts to co-culture HSC with stromal cells were broadly studied. Collagen [82], PEG [83] and PCL [84] scaffolds were fabricated for the co-culturing of HSC with stromal cells, regardless of their source, all of which led to an improved proliferation of HSCs. Stromal cells were often conceived as support biomaterial, mimicking the biological component of HSC niche in bone marrow. Biological cues were further incorporated in the scaffolds to direct HSC activity. Mahdik et al. [121] used a methacrylamide-functionalized gelatin hydrogel to covalently immobilize stem cell factors (SCF), which could selectively maintain HSC multipotency. The incorporation of SDF-1α into PLGA scaffolds was shown to improve HSC recruitment and reduce subsequent inflammatory responses [122]. In conclusion, specific biomaterials can be tuned mechanically, chemically or biologically to meet the criteria for the hematopoietic therapy.

4.5. Bone/cartilage tissue

Nowadays, aging and bone injuries have led to a prominent problem of bone tissue loss. To address the clinical demand, a plethora of efforts have employed tissue engineering approaches, comprised of biomaterials and stem cells, to promote bone and cartilage regeneration [123-125].

Different cell types have been investigated for bone repair. Straightforwardly, ESCs were cultivated on 3D osteoconductive scaffolds to form large and compact bone constructs primed for transplantation [126]. Osteogenic potential of bone marrow-derived mesenchymal stem cell MSC was also validated to become any bone/cartilage component, including osteoblasts, chondrocytes, myocytes and adipocytes [128]. While whether or not the age of the MSC donor affects the cell viability and functionality still under debate [129,130], the ‘aging’ of MSC itself in terms of reduced proliferation and differentiation capacity [129], and its limited quantity in a body restricts its large-scale applications. On this account, iPSCs [131] or adipose-derived stem cell (ADSC) [132] were identified as a more reliable cell source for bone regeneration. iPSCs surpasses the limitation of cell quantities and age and may be applied to three strategies for bone regeneration [131]: a) generating MSCs through an embryoid body (EB); b) generating MSCs without an EB; c) generating bone tissue bypassing MSC or EB. iPSCs were demonstrated to have similar or even superior osteogenic capacity to MSC itself in terms of reduced proliferation and differentiation capacity [129], and its limited quantity in a body restricts its large-scale applications. On this account, iPSCs [131] or adipose-derived stem cell (ADSC) [132] were identified as a more reliable cell source for bone regeneration. iPSCs surpasses the limitation of cell quantities and age and may be applied to three strategies for bone regeneration [131]: a) generating MSCs through an embryoid body (EB); b) generating MSCs without an EB; c) generating bone tissue bypassing MSC or EB. iPSCs were demonstrated to have similar or even superior osteogenic capacity to MSC. On the other hand, ADSC is a specific type of MSC but with distinct advantages over those derived from bone marrow: better angiogenic and osteogenic properties, higher proliferation and differentiation capacity, larger percentage of stem cell progenitors in tissue [133-135] and most importantly, less affected by the donor age [132,136]. The elevated angiogenesis and osteogenesis of ADSCs rendered better performance in a pre-clinical bone allografting trial [137].
The basic design of biomaterials for bone/cartilage tissue engineering emphasizes pore morphology and structure to direct osteo-differentiation, which happen to be the most crucial characteristics of a scaffold. Cells tend to attach, proliferate and migrate on pores larger than 100 μm, and pores smaller than 50 μm are beneficial for nutrient and waste transport [138,139]. Salerno et al. [140] found macro-porosity was favored by colonization of osteoblast MG63 in bi-modal porous PCL scaffolds. Particularly, diameters in the range of 380–450 μm could promote chondrocyte and osteoblast growth, while sizes ranging from 290 to 310 μm could accelerate bone differentiation [141]. The introduction of osteo-inductive and osteo-conductive structures into biomaterials has recently become a promising method to produce heterogeneity in the structures, mimicking that of natural bones [142]. For example, Rodrigues et al. [143] constructed a starch/PCL scaffold containing an osteogenic layer and a chondrogenic layer for improved osteoarthritic treatment.

Likewise, natural biomaterials, including collagen, gelatin [85], hyaluronic acid [87], fibrin [88], alginate [144] et al., are widely reported for bone tissue engineering. However, natural biomaterials are often taken as a composite component merely to enhance biocompatibility due to a strict demand of mechanical properties for bone tissue engineering. Bioceramics such as HA, tricalcium phosphate and bioactive glasses [89] are often used as additives to enhance their mechanical properties, osteo-inductivity and osteo-conductivity. In addition, GAG [90] was introduced to promote scaffold, cellular and biological cue interactions as it has strong affinity to growth factors for directing cell fate. PLA [91], PLGA [92], PCL [93] and polyurethane (PU) [94] are popular synthetic biomaterials for bone/cartilage tissue engineering. Among them, PCL stood out for its superior short-term and long-term biocompatibility due to non-inflammatory and nontoxic degradation products [145–147], unlike acid-releasing PLA and PLGA. Moreover, the degradation of PCL was rather slow (> 1 year), matching the regeneration of natural bones and its medical application was approved by FDA [141].

5. Stem cell biomanufacturing

Biomanufacturing uses biological systems including enzymes, microorganisms, cells, tissues, plants and animals to manufacture commercial products for agricultural, food, energy, material and pharmaceutical industries [148]. Biomanufacturing can be traced back to 1860s, when fermentation products appeared on the market [148]. Today, biomanufacturing can cover a wide range of chemical products [149], regenerative medicine [150] and even metal biofimery [151]. However, most of stem cell-associated biomanufacturing is limited to the manufacturing or fabrication of tissue engineering devices. An exception was conducted by Google who invested $250,000 for a program to make hamburgers from cow stem-cell-derived muscles [152]. While the majority of literature relevant to biomanufacturing and stem cells concentrates on the design of bioreactors [153–155] or product quality control [156], those focusing on the large-scale production of stem-cell-associated therapeutic products brought in the term biofabrication.

Biofabrication is a vigorous and young field that is capturing attention in tissue engineering and regenerative medicine. Biofabrication utilizes automated processes such as additive manufacturing technologies to produce 3D biomaterial-based cell culture systems [157]. Biofabrication is characterized by cell-based building blocks, bio-inspired fabrication methods and biological products [158]. Accordingly, the bio-products of biofabrication are mostly hybrids of stem cells and biomaterials, including scaffolds, microcarriers, microgels, etc. Because the ultimate goal of biofabrication in tissue engineering and regenerative medicine is to achieve a favored spatial organization of constructs for stem cells to differentiate and mature on after implantation, all of the cell types and biomaterials mentioned above can be employed, and the focus of biofabrication becomes more of the methodologies of ‘fabrication’ than on the selection of building blocks. Bajaj et al. [159] reviewed common biofabrication techniques, including solvent casting, particle leaching, freeze drying, gas foaming, bioprinting and photolithography. Among these, novel techniques like bioprinting and photolithography are legitimate approaches for biofabrication as they allowed for the involvement of live cells. Likewise, Groll et al. [157] categorized the biomanufacturing (additive manufacturing methodologies) into bioprinting and bioassembly based on the organization mechanism of building blocks.

5.1. Bioprinting

The evolution of biofabrication was parallel with the development of bioprinting [157]. The intimate relationship between the two dimmed the inconsistent concepts and often misguided the public to consider them as interchangeable. Actually, bioprinting is a frequently-used strategy of biofabrication, defined as the process for patterning and assembling cells, tissues, molecular cues and biomaterials [160]. The minimum fabrication unit of bioprinting is down to molecular level and when cells, cell aggregates, biomaterials or bioactive molecules are small enough, they are considered ‘bioink’ [161,162] and can be bioprinted. There are two types of bioprinting featured by the printing equipment: drop-based and extrusion. Drop-based bioprinting uses an inkjet and deposits bioink dropwise to form a gel or solid scaffold [161]. Extrusion bioprinting employs a mechanical extruder to continuously deposit bioink [163].

Bauwens et al. [164] bioprinted different sizes of hESC/poly (d-methylsiloxane) (PDMS) composite and found that colonies containing different germ-layer-based hESCs had distinct differentiation capacity, indicating a size-motivated effect of the bioprinting product. The size control idea was further progressed by Dias et al. [165] using laser direct-writing to print ESCs onto gelatin. Other than ESCs, iPSCs could also be bioprinted into gelatin methacryl (GelMA) microfibrous scaffolds for formation of an endothelial-integrated myocardium by an extruder [166] or into alginate hydrogels for hepatocyte differentiation by a drop-based printing platform [167]. For the broad application in a variety of tissue repairs, MSCs and tissue-specific progenitor cells were bioprinted with regard to the target tissue. Gaeble et al. [168] co-seeded human umbilical vein endothelial cells (HUVEC) with hMSCs onto a urethane urea (PEUU) cardiac patch in a programmed pattern by a laser-induced-forward-transfer (LIFT) cell printing technique and found restored vessel function to the infarct area in mice. Koch et al. [169] used the same technique to deliver fibroblasts with hMSCs onto a gold sheet for skin formation. Gao et al. [170] bioprinted hMSCs together with bioceramics to form poly (ethylene glycol) dimethylacrylate (PEGDMA) scaffolds for bone/cartilage tissue repair. Ma et al. [171] bioprinted hepatic progenitors, HUVECs and hMSCs onto GelMA and glycidol methacrylate-hyaluronic acid (GMHA) chips to generate a liver model for drug screening and pathophysiology studies. Huang et al. [172] bioprinted epithelial progenitors with EGF onto gelatin alginate substrate for sweat gland regeneration.

Bioprinting techniques are versatile for pluripotent, multipotent and unipotent stem cell, and have a broad application. Gel-forming biomaterials (alginate, PEG), together with solid-forming materials (PLGA, PCL), bioactive ceramics and glasses, as well as growth factors can all be used as bioink for a programmed printing process.

5.2. Bioassembly

Another biofabrication strategy is bioassembly. Unlike bioprinting, bioassembly starts with large pre-formed cellular constructs such as large cell aggregates, cell fibers, cell sheets and even microtissues. Bioassembly refers to the automated assembly of pre-generated cellular devices produced by cell driven self-organization or cell-material hybridization technologies [157]. The minimum building block unit of bioassembly is a pre-formed cellular device large enough for automated assembly.
The assembling unit can come in the form of microcarriers or microgels. Many researchers [173–175] demonstrated the long-term maintenance, self-renewal capacity and pluripotency of hESCs cultured on microcarriers. These microcarriers may be further assembled by different methods for tissue engineering applications. For instance, Xie et al. [176] employed a pressurized-CO₂, assisted technique to assemble PLGA microscaffolds containing ESCs layer by layer for tissue engineering. The cells in microcarriers were not restricted to ESCs. The same group also bioassembled hMSC microscaffolds and demonstrated preserved protein and DNA activity [177]. Furthermore, Guduric et al. used a PLA membrane with human bone marrow stromal cells and endothelial progenitor cells for a layer-by-layer bioassembly [178]. After 7 days of culture, cell migration and osteoblastic differentiation were observed, suggesting a potential application in bone tissue engineering. In addition to cells, microtissues were also applied as building blocks for bioassembly. Mekhileri et al. [179] pre-differentiated chondrogenic microtissues, and sequentially plotted the microtissues and PEGT/PBT polymer scaffold layer-by-layer using a fluidic-based bioassembly system. Comparable cell viability was obtained between the bioassembled cell-biomaterial hybrids and microtissues.

In summary, the evolution of bioassembly extends the working unit of bioprinting to larger blocks, broadens the methodologies to organize complex structures with a scaffold, and paves the way for scale-up in biofabrication, thus making up to the stem-cell-associated biomanufacturing.

5.3. Scale-up production of stem cells

Stem cells outstand in future therapeutic applications. Nonetheless, the realization of their full potential requires sufficient amount of stem cells, and thus their large-scale production becomes an important issue in stem cell engineering. As mentioned above, the manufacturing of stem cell products often associates with bioreactors, while the complex nature of stem cell culture has limited the direct transfer of 2D culture conditions to 3D bioreactor production. Stem cell fate to either maintain pluripotency or differentiate is specified by the combinatorial influence of biochemical factors, cell-cell interactions, and cell-matrix interactions in the culture environment. Stem cell fate can also be impacted by the mechanical stimuli from its surrounding fluid when the cells are introduced to bioreactors for scaling up. Kinney et al. [180] reviewed and summarized the effect of hydrodynamic environments on stem cell aggregation, metabolism and phenotype in suspension culture. Importantly, they envisioned cell aggregates as a promising method for large-scale cell production [181], while it remains challenging to maintain the differentiation homogeneity of the stem cell aggregates due to the altered cue profile caused by size effect. Biomaterials are then introduced for more homogenous cell assembly [182] as they could regulate stem cell expansion and differentiation by mechanically interacting with cytoskeleton [183], or by harnessing the binding and release of both exogenous and endogenous growth factors [184]. Of note, the controlled spatial delivery of biochemical cues through biomaterials could overcome the diffusion limitation and eliminate heterogeneity of stem cell aggregates [185]. In particular, natural or synthetic microcarriers, with or without functional modification via bioactive groups (DEAE, collagen) for better cell attachment [186], were often applied in bioreactors to engineer stem cell proliferation and differentiation [187,188]. Microencapsulation of stem cells using alginate, agarose, or hyaluronic acid hydrogel was another biomaterial application to avoid agglomeration while preserving cell-cell and cell-matrix contact [187]. For instance, a series of scalable thermoreversible PEG-based or hyaluronic acid-based hydrogel systems [189–191] have been developed for cell spheroid encapsulation to protect cells from hydrodynamic stresses and prevent excessive agglomeration, leading to a 20-fold higher cell yield as compared to traditional suspension culture. While the introduction of microcarriers or microencapsulation could solve one or several problems, they were not perfect solutions as the biomaterials used may exert new problems, such as microcarrier clumping, difficult monitoring and observing of culture condition, and additional separation step of materials and cells [192].

Apart from cell expansion, large-scale generation and manufacturing of iPSCs from patient's somatic cells is another challenge for personalized medicine [193,194]. The current manipulation process to produce clinical-grade iPSCs is time-consuming, labor-intensive, and expensive [195]. For example, the total cost reached up to $50,000 to produce iPSCs for one patient to meet the clinical needs [196]. On this account, Lin and his colleagues developed an alginate hydrogel tube system (AlgTubes) to efficiently reprogram human fibroblasts into iPSCs [195], which resulted in high purity of iPSCs that express pluripotent markers OCT3/4, NANOG, ALP and SSEA4 on day 30 and maintained a long-term (20 passages) pluripotency. This AlgTube system was also applicable to other bio-manufacturing processes, such as biofabrication [197]. The simplicity, good reproducibility, cost-effective and feasible integration of induction, expansion and differentiation of AlgTubes enabled it a promising tool for future manufacturing of iPSCs to meet the various demand for personalized medicine at larger-scale.

The preservation and transportation of large quantities of stem cells represents another challenge in stem cell manufacturing. The aforementioned AlgTubes proposed a simple transportation pattern in hydrogels but its preservation and storage capacity was not evaluated. Li and co-workers discussed various cryopreservation bioprocesses [198] and formulations [199] for large-scale banking of iPSCs, but similar work was not conducted in other studies. Additionally, the online monitoring of cell aggregates in bioreactors and the fact that different cells require distinct culture patterns and quality control measurements [180] make the design of a universal large-scale bioreactors an even more challenging task, which requires a comprehensive master of the stem cell production process, the follow-up transportation and preservation process, and the rational bioreactor design. Even with these challenges, the promise biomaterials hold for the stem cell manufacturing offers tremendous potential for expanding the scope of stem cell-based therapies.

6. Conclusion

Stem cells are gaining attention due to their self-renewal ability and capacity of differentiating into any specific cell type. Pluripotent, multipotent and unipotent stem cells draw different interests in a variety of applications, including drug screening, disease modeling and regenerative medicine. However, immune sensitivity limits the application of stem cells in clinical trials. With the aid of biomaterials, these barriers can be overcome. Natural biomaterials are highly biocompatible, while synthetic biomaterials can be rationally designed for a particular purpose. The incorporation of stem cells into structured and modified biomaterials increases the competence of restoring and repairing dysfunction tissues. The well-organized spatial properties of a biomaterial or scaffold in turn can provide a protective and sometimes inducible microenvironment for the stem cells, mimicking the natural ECM. The stem cell-biomaterial system has been applied to various tissue treatments, including heart, nerve, pancreatic islet, hematopoietic system and bone. The appearance of biofabrication technologies enables the precise design and scalable biomanufacturing of cell-based scaffolds using program-controlled bioprinting or bioassembly.

Declaration of competing interest

None.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.biomac.2019.11.002.

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