Review

Three dimensional printed nanostructure biomaterials for bone tissue engineering

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A B S T R A C T

The suffering from organ dysfunction due to damaged or diseased tissue/bone has been globally on the rise. Current treatment strategies for non-union bone defects include: the use of autografts, allografts, synthetic grafts and free vascularized fibular grafts. Bone tissue engineering has emerged as an alternative for fracture repair to satisfy the current unmet need of bone grafts and to alleviate the problems associated with autografts and allografts. The technology offers the possibility to induce new functional bone regeneration using synergistic combination of functional biomaterials (scaffolds), cells, and growth factors. Bone scaffolds are typically made of porous biodegradable materials that provide the mechanical support during repair and regeneration of damaged or diseased bone. Significant progress has been made towards scaffold materials for structural support, desired osteogenesis and angiogenesis abilities. Thanks for innovative scaffolds fabrication technologies, biodegradable scaffolds with controlled porosity and tailored properties are possible today. Despite the presence of different bone scaffold fabrication methods, pore size, shape and interconnectivity have not yet been fully controlled in most of the methods. Moreover, scaffolds with tailored porosity for specific defects are still difficult to manufacture. Nevertheless, such scaffolds can be designed and fabricated using three dimensional (3D) printing approaches. 3D printing technology, as an advanced tissue scaffold fabrication method, offers the opportunity to produce complex geometries with distinct advantages. The technology has been used for the production of various types of bodily constructs such as blood vessels, vascular networks, bones, cartilages, exoskeletons, eyeglasses, cell cultures, tissues, organs and novel drug delivery devices. This review focuses on 3D printed scaffolds and their application in bone repair and regeneration. In addition, different classes of biomaterials commonly employed for the fabrication of 3D nano scaffolds for bone tissue engineering application so far are briefly discussed.

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1. Introduction

1.1. Bone tissue engineering

Each year, the number of people suffering from organ dysfunction or organ failure due to damaged or diseased tissue is increasing, because of the aging global population, traumas and illnesses such as; heart attacks, strokes and joint degeneration. The existing pharmacological treatment options are also incapable to adequately repair tissue damage and may also drastically reduce the quality of life of the victims. This lack of therapeutic intervention is principally due to the fact that current treatments focuses mostly on prevention or reducing further tissue damage rather than contributing to the repair or regeneration of the damaged tissue [1]. Associated with increasing global population mobility, aging, accidents, war and displacement, trauma and organ/tissue damages such as, organ degeneration, bone fracture and maxillofacial complications have been still rampant [2]. In addition to the physical trauma and functional impairment, such damages do have critical psychosocial and emotional implications [3,4]. Transplantation using donated organs has been practiced in different settings in addition to available pharmacological and supportive treatment modalities. However, inadequate access to donated organs and immunological complications has been major challenges in the field [5–7].

Encompassing multidisciplinary experts including molecular biology, chemistry, cell biology, biomaterial science, immunology, engineering and medicine, tissue engineering has played indispensable roles in the design, fabrication and transplantation of 3D tissues constructs by combining scaffolds, cells and other bioactive materials [8]. Actually, bone is well known for its self-healing abilities; however, in case of large-scale bone defects, self-healing may be extensively delayed or may not be possible at all and external intervention is needed to restore normal tasks [9–11]. Non-union fractures are commonly characterized by a substantial gap between the fractured bone ends. To bridge this distance, a platform is necessary and also serve as a temporary support at the defect zone [12]. Current treatment strategies for non-union bone defects include: the use of autografts, allografts, synthetic grafts, free vascularized fibular grafts, osteoconductive scaffolds, osteoprogenitor cells and growth factor [13]. Autografting remains to be the gold standard strategy, because of its osteoconductive and osteoinductive environment and non-immunogenicity [14,15]. However, limited quantities for harvest and donor morbidity has been the limiting factors for the use of autografting driving the search for alternative solutions [16]. While allografts and synthetic grafts could avoid the mentioned limitations compared to autografts, they do not provide the necessary osteoinductive signals and vascularity required for bone healing [17]. In addition, possibility of graft rejection by the host immune system and disease transmission from donor to host are problems associated with allografts [14]. Synthetic grafts on the other hand are subjected to fatigue and wear upon aging [18]. Most grafts used in clinical settings have also limitations because of the lack of integration with bone substitution at the ends of grafts leading to non-unions, and late graft fracture occurring in as high as 60% of the cases at 10 years [19,20].

Bone tissue engineering has emerged as an alternative for fracture repair to satisfy the current unmet need of bone grafts and to alleviate the problems associated with autografts and allografts [2,18]. It focuses on methods to synthesize and/or regenerate bone to restore, maintain or improve its functions in vivo [21]. The technology offers to possibility to induce new functional bone regeneration using synergistic combination of functional biomaterials, cells, and growth factors [22]. As a result of its desirable attributes, bone tissue engineering is gaining momentum in orthopedic and regenerative medical practices. According to a review by Amini and Laurencin [2012], over half a million people in the US receive bone defect repairs yearly with an estimated cost of 2.5 billion, a figure predicted to be doubled by 2020 [23]. In the US, bone is the second most transplanted tissue next to blood, and the over increasing demand for bone grafts and substitutes was estimated to be $3.3 billion of revenues by 2013, with a compound annual growth rate of 13.8% between 2006 and 2013 [2].

Globally, approximately 15 million fracture incidences are reported (of which up to 10% are complicated by non-unions [18,24]. USA contributes 48.6% of the global market revenue to tissue engineering solutions and is the leading country by investing about 60% of the global tissue engineering expenditure to research and development (R&D) [2]. The process of tissue engineering often begins with the development of a three-dimensional (3D) scaffold to be supported in the medium and essential for the appropriate proliferation and differentiation of cells embedded in or infiltrating within it [1].

1.2. Bone scaffolds

Tissue engineering is a multidisciplinary science employing the principles of engineering and biological sciences for the fabrication of functional constructs used to restore, maintain or improve tissue functions [25,26]. The technology requires three basic elements: cells, factors for tissue induction, and a matrix for seeding a cell. This combination is designed to grow tissue in vitro, prior to implantation in to the subject [25]. The design of the scaffold prior to exposure to cells is of vital importance. The scaffold must present a surface that promotes cell attachment, growth and differentiation, while providing a porous network for tissue growth [27].

When designing a scaffold, the material of choice is of a great concern. For optimum tissue regeneration, biocompatible scaffolds with comparable degradation rate to tissue regeneration are best recommended [28]. After implanted, the scaffold must have the mechanical properties required to temporarily offer structural support until the formation of a new tissue. In addition to being biocompatible and biodegradable, the scaffold must possess key morphological characteristics; it must be highly porous and offer a suitable path for nutrient transmission and tissue ingrowths. To achieve these requirements, tissue engineering scaffolds are often designed to mimic the structure of the naturally occurring extracellular matrix (ECM) [29,30].

Bone scaffolds are typically made of porous biodegradable materials that provide the mechanical support during repair and...
regeneration of damaged or diseased bone [31]. Researches on bone tissue engineering over the past decades have encouraged innovation in new materials, new processing techniques, and applications. Significant progress has been made toward scaffold materials for structural support, desired osteogenesis and angiogenesis abilities [32,33]. Nowadays, biodegradable scaffolds with controlled porosity and tailored properties are possible as a result of innovation in scaffold fabrication using advanced technologies. The exceptional mechanical properties of natural bone are derived from an architectural design that spans nanoscale to macroscopic dimensions, with precisely and carefully engineered interfaces [34]. Different research groups have tried to manipulate the mechanical properties (stiffness, strength, and toughness) of scaffolds through the design of nanostructures (the inclusion of nanoparticles or nano fiber reinforcements in polymer matrices) to mimic the natural nanocomposite architecture of bone [34].

There has been an increasing interest in scaffold-based strategies for bone tissue engineering as represented by the exponential rise in the number of scientific articles over the past decade (Fig. 2A). Various scaffolds used in conjunction with stem cells and gene therapy strategies have demonstrated promising results of new bone formation and repair of segmental defects in both small and large animal studies [3]. Justifying that scaffolds are an integral part of bone tissue engineering. Scaffolds are 3D biocompatible structures which can mimic the ECM properties (such as mechanical support, cellular activity and protein production through biochemical and mechanical interactions), and provide a template for cell attachment and stimulate bone tissue formation in vivo [21,35]. In addition to chemistry, other critical parameters which define the performance of a scaffold's are: pore size, pore volume and mechanical strength. At an early phase, bone ingrowth takes place at the periphery of scaffolds with a negative gradient in mineralization toward the inner parts. For continuous ingrowth of bone tissue, interconnected porosity is important as it can allow nutrients and molecules to transport to inner parts of a scaffold to facilitate cell ingrowth, vascularization, as well as waste material removal [36,37].

As surface area per unit volume can be increased with higher porosity, the biodegradation kinetics of scaffolds can be influenced by varying pore parameters. Biodegradation through a cell mediated process or chemical dissolution are both important to ascertain stabilized repair and scaffold replacement with new bone without any remnant [37]. A minimum pore size between 100 and 150 μm is needed for bone formation [38,39]. However, enhanced bone formation and vascularization are reported for scaffolds with pore size larger than 300 μm [40,41]. The permeability of nutrients through the scaffold can also be controlled by pore volume and the mechanical properties. Besides the biological performance, the initial mechanical properties and strength, degradation rate should match that of the host tissue for optimum bone healing [42]. The pore size, geometry, and strut orientation with respect to the loading direction can highly affect the degradation kinetics of porous scaffolds [43]. Finally, surface properties such as chemistry, surface charge and topography also influence hydrophility and in turn cell—material interactions for bone tissue ingrowths [44].

There are different methods that can be applied for bone scaffold fabrication. Chemical/gas foaming [45], solvent casting, particle/salt leaching [46], freeze drying [47], thermally induced phase separation [48], foam-gel technology [49] and electrospinning [50] are some of the extensively used techniques in the field. But, it is not possible to fully control, pore size, shape, and its interconnectedness in these approaches. Moreover, scaffolds with tailored porosity for specific defects are difficult to manufacture with most of these approaches [51]. Such scaffolds can be designed and fabricated using additive manufacturing (AM) approaches. A variety of AM methods, including: 3D printing (3DP), solid freeform fabrication (SFF), rapid prototyping (RP), are approaches that allow complex shapes for scaffolds' fabrication directly from a computer-aided design (CAD) file [9]. This review focuses on 3D printed scaffolds and their application in bone repair and regeneration.

1.3. Three-dimensional (3D) printing

Over the last three decades, significant advances have been made on 3D printing technology and it has been used in various industries including; electronics, robotics and healthcare [52]. 3D printing follows an additive principle where by solid objects can be created by printing successive layers using a computer aided modeling. Three-dimensional (3D) printing is becoming a research and development focus in many field including both traditional industries advanced biomedicine as it can quickly and accurately fabricate any desired 3D model only if its size is appropriate. It is conceptually defined as a method for direct digital manufacturing that provides capabilities for creating a wide range of object geometries (including internal channels) using a broad variety of materials such as ceramic, metal, metal-ceramic composite, and polymeric materials [2].

The conventional techniques used for the fabrication of tissue scaffold such as solvent casting, gas foaming, phase separation, particulate leaching, and freeze drying lack the unique features of native tissue responsible for coordination of specialized cell and tissue functions. 3D printing technology, as an advanced tissue scaffold fabrication method offers the opportunity to produce complex geometries with distinct advantages such as fitting into irregular defect sites and mimicking tissue complexity through precise positioning [53,54].

1.3.1. What is 3D printing?

Three-dimensional printing (3DP) is a manufacturing process in which objects are made by fusing or depositing materials such as: plastic, metal, ceramics, powders, liquids, or even living cells in layers to produce a 3D object [55]. This process is also referred to as additive manufacturing (AM), rapid prototyping (RP), or solid freeform technology (SFF). 3D printers function in a similar fashion to traditional inkjet printers; however, three-dimensional objects are built in 3D unlike printing layers of ink on papers [56]. 3D printing is expected to revolutionize the healthcare technology through the provision advanced diagnostic, imaging and therapeutic options [55,57]. Despite the presence of dozens of 3D printing process with varying technological platforms, resolution capacities, production efficiencies and input material requirement, all can build a 3D object in almost any shape imaginable as defined in a computer-aided design (CAD) file [58,59]. 3D printing process is founded on the development of virtual blueprints of objects using CAD which can later be scanned by the printing machine, built on from a series of layers and finally fused to generate the desired shape. The instructions from the CAD system guides the movement of the 3D print head along the x-y-z plane to build the object vertically layer by layer. The technology offers the opportunity to convert two-dimensional (2D) radiographic images such as x-rays, magnetic resonance imaging (MRI), or computerized tomography (CT) scans to digital 3D print files for creating complex, customized anatomical and medical structures [60,61].

1.3.2. History of 3D bioprinting

The conception of 3D printing, also referred to as additive manufacturing (AM), rapid prototyping (RP), or solid-freeform technology (SFF), was first developed by Charles Hull in 1980s and he patented the first 3D printing technology, stereolithography in 1986 by the US Patent Office [62,63]. His basic training in physics and his work...
on hotopolymers for the production of plastic objects at the Ultra Violet Products company helped him develop the technology [64].

Under the stereolithography technology, an STL file format is used to interpret the date from CAD and that can be electronically communicated to the printers for the manufacturing of 3D objects with the desired color, texture, and thickness [64,59].

The initial technology had several limitations including lengthy fabrication process and design imperfections. Subsequently, Hull and other investigators have made significant developments like the STL file format, the CAD software and data transmission systems. Hull also developed the first commercial 3D printer, commonly referred as “Stereolithography Apparatus”. 3D printing technology was further revolutionized following the development of Fused Deposition Modeling (FDM) by Scott Crump at Stratasys in 1990 [65]. Michael Cima and Emanuel Sachs from MIT also patented the first apparatus termed “3D printer” in 1993 to print plastic, metal, and ceramic parts [66]. Companies such as Helisys Organovo have developed technologies to print objects from living human tissue [67].

3D bioprinting is based on the same principles of earlier 3D printing technologies, but has been customized to manufacture permanent implants, biomimetic scaffolds and drug delivery platforms using cells, growth factors and biomaterials as input materials. The technology can produce objects with controlled morphology and internal structure having highly similar structure to the human body [68,69]. Currently, the technology is used in various aspects of tissue engineering and regenerative medicines applications including hard and soft tissue printing, cartilage printing, skin printing and tumorous tissue model printing [69]. The commonly used printing technologies for 3D bioprinting include laser printing, inkjet printing and extrusion printing with unique features as presented under Table 1. The control and optimization of key factors such as input materials properties, scaffold structure, printing precision and environmental control are essential elements for successful 3D bioprinting [70–72].

According to the comprehensive review by Ventola [59], the major steps in 3D bioprinting include: (i) creating a blueprint of the desired organ with its vascular architecture; (ii) generating a bioprinting process plan; (iii) isolating stem cells and differentiating them into organ-specific cells; (iv) preparing bioink reservoirs with organ-specific cells, blood vessel cells, and support medium to be loaded into the printer; (v) bioprinting the required product; and (vi) placing the bioprinted organ in a bioreactor prior to transplantation (Fig. 1).

### 2. Medical application of 3D printing

The medical applications of 3D printing date back to the early 2000s following the production and use of dental implants and prosthetics [75]. Since then, the medical applications for 3D printing have evolved considerably. Medical uses for 3D printing, both actual and potential, can be organized into several broad categories, including: tissue and organ fabrication; creation of customized prosthetics, implants, and anatomical models; and pharmaceutical research regarding drug dosage forms, delivery, and discovery [76]. Some reviews describe the use of 3D printing to produce bones, ears, exoskeletons, windpipes, a jaw bone, eyeglasses, cell cultures, stem cells, blood vessels, vascular networks, tissues, and organs, as well as novel dosage forms and drug delivery devices [77]. Its application in medicine can provide many benefits, including: the customization and personalization of medical products, drugs, and equipment; cost-effectiveness; increased productivity; the democratization of design and manufacturing; and enhanced collaboration [78].

Following the development of fast and precise prototyping 3D printing machines, companies have started commercialization of various medical technology products. Availability of open access software technology sources helped rapid proliferation and commercialization of the platform for medical application [64]. The growing application of 3D printing in medicine is primarily associated with the efficient manufacturing process, and the flexibility to manufacture medical products with customized size and shape. The technology has been used for the production of various types bodily constructs such as blood vessels, vascular networks, bones, cartilages, exoskeletons, eyeglasses, cell cultures, tissues, organs and novel drug delivery devices [55,79]. Generally, the medical applications of 3D printing technology can be broadly categorized as: (i) tissue and organ fabrication; (ii) manufacturing of prosthetics, implants and anatomical models; and (iii) development of novel drug delivery platforms and advanced dosage forms [59]. Because of the increasing aging population, natural and manmade crisis, accidents, birth defects and related medical problems, the demand for replacement and transplant products is significantly growing. In the light of increasing demand and dire shortage of donors, 3D bioprinting will continue to play major roles in the field [80,81].

As the latest advancement in tissue engineering, 3D bioprinting has been predicted to revolutionize the production of tissue constructed using cell-based material inputs and well-established bioink technologies. Because of their desired features such as; greater differentiation capacity and self-renewal, stem cells are

### Table 1

Unique features of the major 3D bioprinting technologies [73].

| Print methods        | Bioinks                        | Resolution | Cell viability | Cell density | Print speed | Target tissue   |
|----------------------|--------------------------------|------------|----------------|--------------|-------------|-----------------|
| Laser-assisted Printing | Fibrinogen, collagen, GelMA | 1–50 μm   | 97%            | 10⁶ cells/ml | 100–1600 mm/s | Skin, vesse    |
| Inkjet Printing     | Collagen, poly(ethylene glycol) dimethacrylate (PEGDMA), fibrinogen, alginate , GelMA | 50–500 μm  | 85–98%         | <5 × 10⁶ cells/ml | 1000–5000 droplets/s | Skin, cartilage, bone, tumor, liver |
| Extrusion Printing  | Gelatin, poly-caprolactone (PCL), polyethylene glycol (PEG), alginate, hyaluronic acid (HA), polyamide(PA), polydimethyl-siloxane (PDMS) dECM, nanocellulose | >50 μm     | 80–96%         | Cell spheroid | 5–20 mm/s       | Skin, cartilage, vessel, bone, muscle, tumor, heart |
gaining momentum for 3D biofabrication of precise tissue scaffolds and replacements organ constructs [82,83].

2.1. 3D printed bone nanoscaffolds

As a scaffolding tissue, bone is responsible for support, protection, load bearing and hematopoietic functions. The human bone is well known for its ability to continuously remodel and rebuild itself [84]. However, large scale defects, degenerations or inflammations caused by accidents, physical traumas, musculoskeletal maladies, infections or tumors may often be difficult to be healed by the natural process demanding for external interventions [85,86]. On the other hand, growing shortage of donors, transplant rejection, donor site morbidity, failure upon aging and mechanical wearing associated with the conventional tissue transplantation and use prosthetic supportive implants calls for cytocompatible and lasting solutions [87,88]. Advances in biomaterial sciences and nanotechnology enabled the application of 3D printing in tissue engineering and regenerative medical practices with better flexibility and clinical outcomes [2,89].

3D bioprinting method was used to fabricate more ideal structural scaffolds with better control of pore morphology, pore size, and porosity. The technology is being used for fabrication of versatile solid free-form structures that can offer an unprecedented flexibility in both material selection and geometry to produce customized scaffolds for growing irregular tissues [90]. While engineering hard tissues like bone, a high degree of porosity together with high mechanical strength is critical which can be difficult to attain using traditional techniques [91]. An ideal 3D scaffold is expected to have important features such as: high porosity, well-interconnected pore networks, and consistent and adequate pore size which can augment cell migration and infiltration [92]. These parameters are essential elements of scaffold geometry which determine the level of access to cell recruitment, vascularization and nutrients which intern influence cell adhesion, proliferation and distribution. There are different views by researchers on the size range of these parameters of which some do have contradicting views where pore sizes ranging from 20 μm to 1500 μm have been reported [39,40,93]. A pore size range of 100–135 μm has been recommended as optimum for effective bone growth [94,95]. Degree of porosity in excess of 90% was recommended for adequate nutrient diffusion and cell–biomaterial interactions [96]; however, it should not be extremely high as it may affect the desired mechanical properties of the scaffold [95]. The selection of input materials is hence a critical component in 3D bio-printing. The different class of biomaterials commonly employed for the fabrication of 3D nanoscaffolds for bone tissue engineering application are presented in Table 2.

Because of the relative similarity with body proteins and receptors, the fabrication such biomaterials as nanoscaffolds could enable them to freely interact with receptors and easily integrate with the membrane matrix structures. Investigations have shown that nanoscaffolds demonstrated better cell functionality than micro and macro level constructs [114]. This is associated with the fact that bone is composed of different proteins like collagen, fibronectin, laminin, and vitronectin which form soft hydrogel nanocomposites in the presence of water; and the bioactive nanoscaffolds can easily mimic the natural environment and augment osteocyte differentiation and mobility pathways [115]. Silk fibroin-hydroxybutyl chitosan blended nanofibers, apatite-collagen-polycaprolactone nanocomposites, Bone Morphogenetic Protein 2 (BMP-2) based gold nanoarrays, BMP-2 based silk fibroin/chitosan/Nanohydroxyapatite nanocomposites have demonstrated promising results in animal models [116,117].

2.1.1. Calcium phosphate (CaPs) based bioactive ceramic scaffolds

Calcium phosphate (CaP) ceramics are commonly used in bone tissue engineering because of their biocompatibility, excellent bioactivity, osteoconductivity, availability and cost-effectiveness [9,114]. The incorporation of biomimetic CaP nanomaterials such as nano-hydroxyapatite (nHA), TCP and CaP are at the forefront of 3D printing research. As a result of its excellent cytocompatibility, osteoconductive and bioactive nature, nHA has been targeted as the future bone nanomaterial to be considered in 3D printing systems, and even used as the main constituent. In addition to nHA, TCP is also utilized in 3D printing. The fine powder form of nHA-TCP were used for the fabrication of nanoscaffolds by a novel 3D sintering method.
Sintered TCP scaffolds are characterized by higher compressive strength and more optimal microporosity to macroporosity ratio which in turn can facilitate the formation of new bone in vivo [51]. Controlled biodegradability is an essential feature of bone scaffolds as it can progressively create space for new tissue growth during regeneration. In this regard, CP scaffolds, particularly TCPs care capable of tunable bioresorption [95]. The degradation products of CP scaffolds also participate in bio-mineralization and can facilitate bone formation and bioactivity [119,120]. The limitation of CP scaffolds; however, is weak and brittle property associated with the porosity limiting their use only in none load-bearing bone repairs. In addition, CP scaffolds lack osteoinductive activity which is important in bone healing process [121]. Combination of CP ceramics with biopolymers has demonstrated improved mechanical and biochemical performances [122–124].

During the process of bone and tissue repair, capillaries and vessel formation, and homogeneous osteoconduction can be enhanced from central channels [9,97]. The effect of pore size on human fetal osteoblasts (hFOB) was studied with 3D-printed TCP scaffolds [99]. The decrease in designed pore size from 1000 to 750 and 500 μm resulted in an increase in proliferated cell density. As can be depicted in Fig. 2, the 3D printed and microwave sintered β-TCP scaffolds show interconnected macro porosity across the sample. The study done by Bose et al. (2013) on the morphologies of hFOB cells on scaffold surfaces and pore walls showed good cell

| Types of Materials | References |
|--------------------|------------|
| 1. Calcium phosphate (CaPs) based bioactive ceramic scaffolds |  |
| - Hydroxyapatite (HA) | [97] |
| - Hydroxyapatite (HA) | [98] |
| - Biphasic calcium phosphate (BCP) | [98] |
| - Tri calcium phosphate (TCP) | [98] |
| - Tri calcium phosphate (TCP) | [99] |
| - CaP mixture with Ca/P ratio of 1.7 | [100] |
| - TTCP/b-TCP | [101] |
| - Tetracalcium phosphate (TTCP), dicalcium phosphate and TCP | [100] |
| - TTCP/calcium sulfate dehydrate | [101] |
| - TTCP/calcium sulfate dehydrate | [101] |
| - α/β-TCP (final product: dicalcium phosphate dihydrate (DCPD)) | [102] |
| - Mesoporous bioactive glasses (MBG) modified β-tricalcium phosphate (MBG-β-TCP) | [91] |
| 2. Polymer based Scaffolds |  |
| - High density PE (HDPE) | [103] |
| - Poly lactic acid (PLA) | [104] |
| - Poly lactic acid (PLA) | [105] |
| - poly (propylene fumarate) | [106] |
| - Starch/PLLA + PCL | [107] |
| - Poly ethylene (PE) or HDPE | [108] |
| - poly (ether- ketone-ketone) | [109] |
| 3. Composite scaffolds |  |
| - α/β-TCP modified with 5 wt% hydroxyl propyl methyl cellulose | [100] |
| - HA/ε-polycaprolactone (PCL) | [110] |
| - HA/maltodextrin | [100] |
| - Polycaprolactone-hydroxyapatite | [111] |
| - PCL/PLGA/β-TCP | [112] |
| - Biodentine/polycaprolactone | [113] |

**Fig. 2.** Photograph of the sintered 3D printed β-TCP scaffolds for mechanical strength and in vivo testing (small samples) [99].
adherence and cell ingrowth into the pores, suggesting that the scaffolds were non-toxic [9].

New bone formation was observed at the implant/host bone interface and inside the interconnected macro and intrinsic micro pores after 4 and 8 weeks in both pure and doped TCP. Tartrate resistant acid phosphatase (TRAP) staining, lacunae formation and microscopic images conducted using osteoclasts also confirmed monocytes differentiation to multinuclear osteoclast-like cells on a wide range of compositions ensuring the biocompatibility of the scaffolds [98]. HA scaffolds with high surface areas demonstrated cytocompatibility and adequate cell adhesion with MC3T3-E1 fibroblast cells in vitro [125]. In vivo biocompatibility and osteoconductivity of 3D-printed scaffolds showed that, the 3Dprinted brushite and monetite cements with controlled open porosity increased osteoconduction in vivo in a goat model [37]. It has been shown that the use of phosphoric acid instead of polymeric binders can improve both resolution and compressive strength [100].

Biphasic calcium phosphate (BCP) and β-TCP hydroxyapatite scaffolds at different concentration of stable phases have been better recommended because of their controlled bioactivity and better balancing between resorption and solubilization which can maintain biomaterial stability while enhancing bone ingrowth [118]. 3D-printed TCP samples with micro and macro-porosity were also facilitated osteogenesis in a rat femur model [126]. To further improve the mechanical properties of the scaffolds, Khalyfa et al. (2007) have utilized tow post-fabrication procedures: sintering and a polymer infiltration on TTCP/β-TCP and TTCP/sodium carbonate dehydrate bone cements, and demonstrated cytocompatibility on MC3T3-E1 cells model. According to their report, the shortest hardening time obtained was between 20 and 40% for citric acid, and 30–40% for lactic acid while used as binders. It was also reported that lower binder concentration in the presence of sodium hydrogen phosphate and phosphoric acid can prolong the hardening time for the cements [101]. 3D printed mesoporous bioactive glasses (MBG) modified β-tricalcium phosphate (MBG-β-TCP) scaffolds with hierarchical pore structure and functional strut surfaces also demonstrated better compressive strength and apatite-mineralization ability with enhanced new bone formation in vivo as compared to BG-β-TCP and β-TCP scaffolds [117].

2.1.2. Polymer based 3D printed bone scaffolds

Bone scaffolds are designed to offer a number of desired functions including: (i) promoting cell–scaffold interactions, cell adhesion and ECM deposition, (ii) facilitating transport of gases, nutrients and regulatory factors that are essential for cell survival, proliferation and differentiation; and (iii) provoking minimal degradation to provide the necessary support for cell attachment and proliferation. Ideally, composite scaffolds are required to have an approximate compressive strength of 100–230 MPa; elastic modulus closer to 7–30 GPa; tensile strength of 50–151 MPa; porosity between 60% and 90%; and an average pore size of >150 μm [133,134]. Besides, smart combination of biodegradable polymers and bioactive ceramics have the ability to control fast autocalytic degradation effect of acidic end groups resulting from hydrolysis of some polymer chains through buffering the pH of the surrounding solution [135].

Combination of polymers with bioceramics or bio glasses have been widely used as composite scaffolds in bone tissue engineering [21,31,136]. Schantz et al. [137] fabricated a biodegradable polymer-ceramic scaffold via 3D printing and testing in vitro using human MSCs within fibrin glue. The results revealed that the cells were able to attach, migrate and osteogenically differentiated within the biomimetic bone scaffold. 3D printed mineral trioxide aggregate/ polycaprolactone (MTA/PCL) hybrid scaffolds with controlled size, high-porosity (70%), and a compressive strength of 4.5 MPa showed effective adhesion, proliferation, and differentiation on human dental pulp cells with excellent physical and chemical properties suitable for bone tissue engineering (hDPSCs). These scaffolds have not only excellent physical and chemical properties but also enhanced osteogenic differentiation, making them useful for bone tissue engineering [90]. Similarly, Yao et al. [111] demonstrated improved cell adhesion, proliferation and chondrogenic differentiation by 3D polycaprolactone-hydroxyapatite scaffolds upon in vitro and in vivo testing. 3Dprinted PCL/PLGA/β-TCP scaffolds also showed enhanced osteogenic potential when tested on human nasal inferior turbinate tissue-derived mesenchymal stromal cells [112]. 3D printed Biodentine/polycaprolactone composite scaffolds with controlled macropore sizes and structures fabricated using extrusion technology for orthopedic and dental applications were proved to have good apatite-forming ability, and enhancing cell proliferation and differentiation when tested on human dental pulp cells [113].

2.1.3. Composite scaffolds

Even though a variety of materials including ceramics, polymers and hydrogels have shown promising results for the fabrication of bone and tissues scaffolds, each of them has limitations in giving all required material attributes, and also in mimicking the natural processes of bone growth when used individually. Combination of materials in the form of 3D composite scaffolds has been widely used with to address the stated limitations [131]. Composites are materials made from two or more constituents with significantly different physical or chemical properties, and produce superior characteristics when combined than the individual constituents [31,132]. Biodegradable composites should have mechanical competence characterized by suitable fracture strength and elastic modulus values, as well as controlled strength and modulus degradation to provide the necessary support for cell attachment and proliferation. Ideally, composite scaffolds are required to have an approximate compressive strength of 100–230 MPa; elastic modulus closer to 7–30 GPa; tensile strength of 50–151 MPa; porosity between 60% and 90%; and an average pore size of >150 μm [133,134]. Besides, smart combination of biodegradable polymers and bioactive ceramics have the ability to control fast autocalytic degradation effect of acidic end groups resulting from hydrolysis of some polymer chains through buffering the pH of the surrounding solution [135].
(HAp) by treatment with a hydrothermal reaction in an NiH4H2PO4 solution and coated with a ε-poly-L-lysine (PLL) polymer solution (5 and 10 w/v) showed significantly increased compressive strength by about 2-fold and 4-fold, respectively, compared with uncoated scaffolds. In another study, 3D scaffolds coated with PCL improved MG-63 cells adhesion and proliferated with improved osteoblast differentiation [110]. Similarly, β-tricalcium phosphate (β-TCP) and polycaprolactone (PCL) composites scaffolds, at 50:50 and 70:30 composition showed an improved cell adhesion and proliferation, and higher alkaline phosphates activity making them ideal for dental applications or regeneration therapies [138].

3. Conclusion

Advances in biomaterial sciences and nanotechnology enabled the application of 3D printing in tissue engineering and regenerative medical practices with better flexibility and clinical outcomes. Over the last decades, 3D bioprinting method has been used to fabricate more ideal structural scaffolds with better control of pore morphology, pore size, and porosity. The selection of input materials is a critical component in 3D bioprinting. Different classes of biomaterials were employed for the fabrication of 3D nanoscaffolds for bone tissue engineering application. Calcium phosphate based bioactive ceramic scaffolds, polymer based scaffolds, and composite of Calcium phosphate based bioactive ceramic and polymer scaffolds have not only excellent physical and chemical properties but also enhanced osteogenic differentiation, making them ideal for dental applications or regeneration therapies [138].

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References

[1] Do AV, Khorsand B, Geary SM, Salem AK. 3D printing of scaffolds for tissue regeneration applications. Adv. Healthc. Mater. 2015;4:1742–62.

[2] Liu W, Li Y, Liu J, Niu X, Wang Y, Li D. Application and performance of 3D printing in nanobiomaterials. J Nanomater 2013;13. https://doi.org/10.1155/2013/681050; ID 681050.

[3] Patrick CW. Tissue engineering strategies for adipose tissue repair. Anat Rec 2001;263:361–6.

[4] Lahiry MR, AS, Skalska AR, Thomas SS, Nejad AG, Cha YS, Park CH, et al. Skeletal design and fabrication of biomimetic 3d scaffolds: unique architectures of extracellular matrices for enhanced adipoogenesis and soft tissue reconstruction. Nature Sci. Rep. 2018;8:5969. https://doi.org/10.1038/s41598-018-23996-w.

[5] Evans RW, Manninen DL, Garrison LJP, Maer AM. Donor availability as the primary determinant of the future of heart transplantation. J Am Med Assoc 1985;255:1892–8.

[6] Wood R, Goto R. Mechanisms of rejection: current perspectives. Transplantation 2012;93:1–10.

[7] Maggiore U, Oberbauer R, Pascual J, Viklicky O, Dudley C, Budde K, et al. Strategies to increase the donor pool and access to kidney transplantation: an international perspective. Nephrol. Dial. Transpl. 2015;30:217–22.

[8] Castells-Sala C, Alemayehu-Rikes M, Fernandez-Muinos T, Rech-Sancho L, Lopez-Chicon P, Aloy- Reverte C, et al. Current applications of tissue engineering in biomedicine. J. Biochir. Tissue. Chip 2013;52.004. https://doi.org/10.4172/2153-0775.p-004.

[9] Bose S, Hahabzadeh S, Bandyopadhaya A. Bone tissue engineering using 3D printing. Mater Today 2013;16:496–504.

[10] Gao C, Schilling AF, Yonezawa T, Wang J, Dai C, Cui X. Bioactive nanoparticles stimulate bone tissue formation in bioprinted three-dimensional scaffolds and human mesenchymal stem cells. Biotechnol J 2014;9:1304–11.

[11] Garcia P, Histing T, Holstein JH, Klein M, Laschke MW, Matthys R, et al. Rodent animal models of delayed bone healing and non-union formation: a comprehensive review. Eur Cell Mater 2013;26:1–14.

[12] Langer R, Vacanti JP. Tissue engineering. Science 1993;260:920–6.

[13] Emara KM, Diab RA, Amara AK. Recent biological trends in management of fracture non-union. World J. Orthop. 2015;6:623–8.

[14] Bose FR, Orofo RD. Bone tissue engineering: hope vs hype. Biochem. Bioph. Res. Co. 2002;92:1–7.

[15] Schroeder JE, Moshref R. Tissue engineering approaches for bone repair: concepts and evidence. Injury 2011;42:609–13.

[16] Sreebna M, Sattah K, Moorhi A, Srinivasan N, Ramsamy K, Selvanayagam N. Biocomposites containing natural polymers and hydroxyapatite for bone tissue engineering. Int J Biol Macromol 2010;47:1–4.

[17] Lane JM, Tomin E, Bostrom MP. Biosynthetic bone grafting. Clin Orthop Relat Res 1995;367:5107–17.

[18] Salgado AJ, Coutsinho OP, Reis RL. Bone tissue engineering: state of the art and future trends. Macromol Biosci 2004;4:743–65.

[19] Wheeler DL, Enneking WF. Allograft bone decreases in strength in vivo over time. Clin. Orthop. Relat. R. 2005;435:36–42.

[20] Soucacos PN, Daliaina Z, Ceris AE, Johnson EO. Vascularised bone grafts for the management of non-union. Injury 2006;37:541–50.

[21] Rezvan K, Chen QF, Blaker JF, Bocaccini AR. Biodegradable and bioactive porous polymer/inorganic composite scaffolds for bone tissue engineering. Biomaterials 2006;27:3431–32.

[22] Akter F, Ibanez J. Bone and cartilage tissue engineering. In: Akter J, editor. Tissue engineering made easy. Cambridge: Elsevier Inc.; 2016. p. 77–99.

[23] Amini AR, Laurencin CT. Nukavarapu SP. Bone tissue engineering: recent advances and challenges. Crit Rev Biomed Eng 2012;40:363–408.

[24] O’Keefe RJ, Mao J. Bone tissue engineering and regeneration: from discovery to the clinic-an overview. Tissue Eng Part B. Rev. 2011;1:77–89.

[25] Vacanti JP, Langer R. Tissue engineering: the design and fabrication of living replacement devices for surgical reconstruction and transplantation. Lancet 1999;354:324–4. https://doi.org/10.1016/s0140-6736(99)0247-7.

[26] Langer R. Tissue engineering: perspectives, challenges, and future directions. Tissue Eng 2007;13:1–2.

[27] Liu X, Ma PX. Polymeric scaffolds for bone tissue engineering. Ann Biomed Eng 2004;32:477–86.

[28] Lee KY, Mooney DJ. Hydrogels for tissue engineering. Chem Rev 2001;101:1869–79.

[29] Smith IO, Liu XH, Smith LA, Ma PX. Nanostructured polymer scaffolds for tissue engineering and regenerative medicine. Wiley Interdiscip. Rev. Nanomed. Nanobiotechnol. 2009;1:226–36.

[30] Dhadayuthapani I, Yoshida Y, Mekawaka T, Kumar DS. Polymeric scaffolds in tissue engineering application: a review. Int. J. Polym. Sci. 2011. https://doi.org/10.1155/2011/20602.

[31] Bose S, Roy M, Bandyopadhaya A. Recent advances in bone tissue engineering scaffolds. Trends Biotechnol 2012;30:546–54.

[32] Kanzelner JM, Orofo ROC. Osteoinduction and osteogenesis: the potential for engineering bone. Eur Cell Mater 2008;15:100–14.

[33] Chen Y, Chen C, Kawaeze N, Chen G. Promoted angiogenesis and osteogenesis by dexamethasone-loaded calcium phosphate nanoparticles/ collagen composite scaffolds with microgroove networks. Nature Sci. Rep. 2018;8:14143. https://doi.org/10.1038/s41598-018-23995-y.

[34] Gong T, Xie J, Liao J, Zhang T, Lin S, Lin Y. Nanobiomaterials and bone regeneration. Bio Res 2015;3:15029. https://doi.org/10.1038/bomeres.2015.29.
[35] Müller B, Deyhle H, Fierz FC, Irseck SH, Yoona JY, Mushkolaj S, et al. Bio-
imetic hollow scaffolds for long bone replacement. Proc SPIE Int Soc Opt Eng 2009;7401:1.

[36] Jones AC, Anis CH, Sheppard AP, Hutmacher DW, Milsorpe BK. Knackstedt MA. Assessment of bone ingrowth into porous biomaterials using MICRO-CT. Biomaterials 2007;28:2491–504.

[37] Hadzovic P, Chudoba D, Olsman CJ, Assent DC, van Blitterswijk CA, Barrallet J. Osteoconduction and osteoinduction of low-temperature 3D printed bio-
ceramic implants. Biomaterials 2008;29:944–53.

[38] Mour M, Das D, Winkler T, Hoeng E, Mielge K, Morlock MM, et al. Advances in porous bio-
materials for dental and orthopaedic applications. Materials 2010;3:2947–74.

[39] Murphy CM, O’Brien FJ. Understanding the effect of mean pore size on cell ac-
tivity in collagen-glycosaminoglycan scaffolds. Cell Adv. Mghr. 2010;4:377–81.

[40] Pascualgourov V, Bandyopadhyay A. Porosity of 3D bioprinted biomaterials and osteo-
genesis. Biomaterials 2005;26:5474–91.

[41] Otsuki B, Takemoto M, Fujibayashi S, Neo M, Kukubo T, Nakamura T. Pore throat size and connectivity determine bone and tissue ingrowth into porous implants: three-dimensional micro-CT based structural analyses of porous bioactive titanium implants. Biomaterials 2006;27:5892–900.

[42] Bandypadhyay A, Bernard S, Xue W, Bose S. Calcium phosphate-based resinable ceramics: influence of MgO, ZrO, and SiO2 dopants. J Am Ceram Soc 2006;89:2673–88.

[43] Base S, Tarafder S, Banerjee SS, Davies NM, Bandypadhyay A. Understanding in vivo and regional property and mechanical variation in MgO, SiO2 and SiO2 doping on CaP-3D. J ICM. 2009;24:1282–90.

[44] Tarafder S, Banerjee S, Bandypadhyay A, Bose S. Electrically polarized bipolar calcium phosphates: adsorption and release of bovine serum al-
umin. Langmuir 2010;26:16625–9.

[45] Knoepfler P, Barash J, Walenski P, Kynk T, Ciach T. Fabrication of in-situ foamed chitosan/lactose-TPC scaffolds for bone tissue engineering application. Mater Lett 2012;85:124–7.

[46] Stoppato M, Carletti E, Sidarovich V, Quattrone A, Unger RE, Kirkpatrick CJ. Cui X, Boland T, D’Lima DD, Lotz KM. Thermal inkjet printing in tissue en-
thinking.” Q IEEE Pulse 2013;4:12

[47] Murphy CM, Holmes B, Faucett S, Zhang LG. Three-dimensional printing of biological materials. United States: USPaT Of 2000;21:2529

[48] Nandakumar S, Bandyopadhyay A, Bose S. Calcium phosphate/chitosan nanocomposites: a novel approach for bone regeneration. J. Bioact. Compat. Pol. 2013;28:16

[49] Nandakumar S, Bandyopadhyay A, Bose S. Electrically polarized bipolar calcium phosphates: adsorption and release of bovine serum al-
umin. Langmuir 2010;26:16625–9.

[50] Knoepfler P, Barash J, Walenski P, Kynk T, Ciach T. Fabrication of in-situ foamed chitosan/lactose-TPC scaffolds for bone tissue engineering application. Mater Lett 2012;85:124–7.

[51] Im O, Li J, Wang M, Zhang LG, Keidar M. Biomimetic three-dimensional nanocrystalline hydroxyapatite and magnetically synthesized single-walled carbon nanotube chitosan nanocomposite for bone regeneration. Int. J. Nanomedicine 2012;7:2087.

[52] Stoppato M, Carletti E, Sidarovich V, Quattrone A, Unger RE, Kirkpatrick CJ. Cui X, Boland T, D’Lima DD, Lotz KM. Thermal inkjet printing in tissue en-
thinking.” Q IEEE Pulse 2013;4:12

[53] Murphy CM, Holmes B, Faucett S, Zhang LG. Three-dimensional printing of biological materials. United States: USPaT Of 2000;21:2529

[54] Nandakumar S, Bandyopadhyay A, Bose S. Calcium phosphate/chitosan nanocomposites: a novel approach for bone regeneration. J. Bioact. Compat. Pol. 2013;28:16

[55] Zhang Y, Xia L, Zhai D, Shi M, Luo Y, Feng C, et al. Mesoporous bioactive glass scaffolded neural tissue engineering. Neurosurgery 2012;71:61”.

[56] Reznova RA, Kasyanov V, Mironov V, da Silva JVL. Organ Printing as an in-
formation technology full of opportunities and challenges. Biodes. Manuf. 2018.https://doi.org/10.1007/s42242-018-0004-3.

[57] Zhan B, Luo Y, Ma L, Gao L, Li L, Y Xue q, et al. 3D bioprinting: an emerging technology full of opportunities and challenges. Biodes. Manuf. 2018.https://doi.org/10.1007/s42242-018-0004-3.

[58] Zhan B, Luo Y, Ma L, Gao L, Li L, Y Xue q, et al. 3D bioprinting: an emerging technology full of opportunities and challenges. Biodes. Manuf. 2018.https://doi.org/10.1007/s42242-018-0004-3.

[59] Zhan B, Luo Y, Ma L, Gao L, Li L, Y Xue q, et al. 3D bioprinting: an emerging technology full of opportunities and challenges. Biodes. Manuf. 2018.https://doi.org/10.1007/s42242-018-0004-3.

[60] Zhan B, Luo Y, Ma L, Gao L, Li L, Y Xue q, et al. 3D bioprinting: an emerging technology full of opportunities and challenges. Biodes. Manuf. 2018.https://doi.org/10.1007/s42242-018-0004-3.

[61] Zhan B, Luo Y, Ma L, Gao L, Li L, Y Xue q, et al. 3D bioprinting: an emerging technology full of opportunities and challenges. Biodes. Manuf. 2018.https://doi.org/10.1007/s42242-018-0004-3.
BMP-2 onto individually formed hydroxyapatite matrices for heterotopic bone induction. Int. J. Oral Max. Surg. 2012;41:1153–60.

Detsch R, Schafer S, Desinger U, Ziegler G, Seitz H, Leukers B. In vitro-osteoclastic activity studies on surfaces of 3D printed calcium phosphate scaffolds. J. Biomater Appl 2011;26:359–80.

Taraferd S, Balla VK, Davies NM, Bandyopadhyay A, Bose S. Microwave-sintered 3D printed tricalcium phosphate scaffolds for bone tissue engineering. J. Tissue Eng. Regen. Med. 2013;7:631–41.

Vorndran E, Klarner M, Klammert U, Grover LM, Patel S, Barralet JE, et al. 3D powder printing of β-tricalcium phosphate ceramics using different strategies. Adv Eng Mater 2008;10:867–71.

Khalyfa A, Vogt S, Weisser J, Grimm G, Rechtenbach A, Meyer W, et al. Development of a new calcium phosphate powder-binder system for the 3D printing of patient specific implants. J Matr Sci: Mater Med 2007;18:909–16.

Klammert U, Gbureck U, Vorndran E, Rödiger J, Meyer-Marcquot P, Küler B. 3D powder printed calcium phosphate implants for reconstruction of cranial and maxillofacial defects. J. Cranio. Maxill. Surg. 2010;38:565–70.

Suwanprateeb J, Thammarakcharoen F, Wongsuvan V, Chokevivat W. Development of porous powder printed high density polyethylene for personalized bone implants. J. Porous. Mat. 2012;19:623–32.

Giordano RA, Wu BM, Borland SW, Cima LG, Sachs EM, Cima MJ. Mechanical properties of dense polyactic acid structures fabricated by three dimensional printing. J. Biomat. Sci-Polym. E 1997;8:63–75.

Kao CT, Lin CC, Chen YW, Yeh CH, Fang HY, Shey MY. Poly (dopamine) coating of 3D printed poly (lactic acid) scaffolds for bone tissue engineering. Mater Sci Eng C 2015;56:165–73.

Wang MO, Piard CM, Melchiorri A, Dreher ML, Fisher JP. Evaluating changes in structure and cytotoxicity during in vitro degradation of three-dimensional printed scaffolds. Tissue Eng. Part A 2015;21:1642–53.

Lam CFX, Mo XM, Teoh SH, Hutmacher DW. Scaffold development using 3D printing with a starch-based polymer. Mater Sci Eng C 2002;20:49–56.

Suwanprateeb J, Suwannapruk W, Wasoontararat K, Leelapatranurak K, Wanumkarong N, Suvannapruk W. Preparation and comparative study of a new porous polyethylene oculum implant using powder printing technology. J. Bioact. Compat. Pol. 2011;26:317–31.

Ganey T. Cell proliferation and vitality determination of osteoblasts on different materials and surface characteristics; Interpretation of laboratory data. Confidential OPM Report-March., 2011.

Kim BS, Yang SS, Park H, Lee SH, Cho YS, Lee J. Improvement of mechanical properties of β-tricalcium phosphate/polycaprolactone 3-dimensional printed scaffolds by ε-polycarbonate coating. J Biomater Sci Polym Ed 2017;28:1256–70. https://doi.org/10.1080/02698282.2017.1312059.

Yao Q, Wei B, Guo Y, Jin C, Du X, Yan C, et al. Design, construction and mechanical testing of digital 3D anatomical data-based PCL–HA bone tissue engineering scaffold. J. Mater. Sci-Mater. M. 2005;16:807–16.

Pati F, Song TH, Rijal G, Jang J, Kim SW, Cho DW. Ornamenting 3D printed scaffolds to biomimic extracellular matrix. J. Mater. Sci-Mater. M. 2015;26:51.

Bose S, Tarafder S. Calcium phosphate ceramic systems in growth factor and methasone delivery. Artif Cells Nanomed Biotechnol 2019;47:4020–6.

Kalita SJ, Bose S, Hosick HT, Bandyopadhyay A. Development of controlled porosity polymer-ceramic composite scaffolds via fused deposition modeling. Mater Sci Eng C 2003;23:61–20.

Langer R, Tirrell DA. Designing materials for biology and medicine. Nature 2004;428:87–92.

Tappa K, Jammalamadaka U. Novel biomaterials used in medical 3D printing techniques: Review. J Funct Biomater 2018;9:1–16.

Rhee S, Puetzer JL, Mason BN, Reinhart-King CA, Bonassar LJ. 3D Bioprinting of spatially heterogeneous collagen constructs for cartilage tissue engineering. ACS Biomater Sci Eng 2016;2:1800–5.

Ren Z, Parisi C, di Silvio I, Dini D, Forte AE. Cryogenic 3D printing of super soft hydrogels. Sci Rep 2017;7:16293. https://doi.org/10.1038/s41598-017-16668-9.

Turnbull G, Clarke J, Picard F, Riches P, Jia L, Han F, et al. 3D bioactive composite scaffolds for bone tissue engineering. Bioact. Mater. 2018;3:278–314.

Harris B. Engineering composite materials. London: The Institute of Materials; 1999. retrieved from, https://pdfs.semanticscholar.org/6818/f30049385b7dd0959435087cf52eb8493e61.pdf. [Accessed 11 December 2018].