Biomaterials for In Situ Tissue Regeneration: A Review

Saba Abdulghani * and Geoffrey R. Mitchell

Centre for Rapid and Sustainable Product Development, Polytechnic of Leiria, 2430-080 Marinha Grande, Portugal; geoffrey.mitchell@ipleiria.pt
* Correspondence: saba.a.silva@ipleiria.pt

Received: 17 October 2019; Accepted: 17 November 2019; Published: 19 November 2019

Abstract: This review focuses on a somewhat unexplored strand of regenerative medicine, that is in situ tissue engineering. In this approach manufactured scaffolds are implanted in the injured region for regeneration within the patient. The scaffold is designed to attract cells to the required volume of regeneration to subsequently proliferate, differentiate, and as a consequence develop tissue within the scaffold which in time will degrade leaving just the regenerated tissue. This review highlights the wealth of information available from studies of ex-situ tissue engineering about the selection of materials for scaffolds. It is clear that there are great opportunities for the use of additive manufacturing to prepare complex personalized scaffolds and we speculate that by building on this knowledge and technology, the development of in situ tissue engineering could rapidly increase. Ex-situ tissue engineering is handicapped by the need to develop the tissue in a bioreactor where the conditions, however optimized, may not be optimum for accelerated growth and maintenance of the cell function. We identify that in both methodologies the prospect of tissue regeneration has created much promise but delivered little outside the scope of laboratory-based experiments. We propose that the design of the scaffolds and the materials selected remain at the heart of developments in this field and there is a clear need for predictive modelling which can be used in the design and optimization of materials and scaffolds.

Keywords: in situ tissue engineering; biomaterials; natural polymers; synthetic polymers

1. Introduction

Key to the survival of humans is the ability to self-heal when damaged. We are most familiar with this process when our skin self-repairs after an injury, but the same process is happening with many organs. For example, the liver is responsible for removing toxins from the blood and can be easily damaged. Therapies for liver cells create many challenges arising from the complexity of the structure of the liver and its function. The key functional or parenchymal tissue is comprised of hepatocytes which are very sensitive to the environmental architecture and when removed from the body lose both their ability to replicate and provide specific liver functions [1]. To repair the damaged liver tissue, it may be sufficient to eliminate the toxin from the diet, for example if the patient stops drinking alcohol. A developing alternative is the use of cell-based therapies and a number of successes have been reported. If the damage is too great or the toxin cannot be eliminated, then a transplant would be an option. Although such allograft transplants carry risks of rejection and other complications, the major drawback is the limited number of organs available and the suitability of the recipients. The need for alternative approaches has led to the development of the field of tissue engineering (TE). TE focuses on regenerating damaged tissues, rather than replacing them, by developing biological substitutes which restore and improve tissue function [2]. The term ‘tissue engineering’ originated at a National Science Foundation (NSF) workshop in Granlibakken, California and later published...
in the proceedings of the workshop by Skalak in 1988 [3]. This definition was later strengthened by Langer and Vacanti in 1993 [4] and subdivided into three main groups which are focused on (a) isolated cell implantation; (b) introduction of growth factors to the cells to stimulate cell growth and (c) the introduction of cells on or within different scaffolds which aim to mimic the extracellular matrix. The latter is the most commonly used tissue engineering approach that is associated with seeding living cells on natural or synthetic extracellular scaffolds to create an implantable substrate [5], capable of providing a functional 3D structural support system to promote the formation of a new extracellular matrix (ECM) similar to that of the damaged native tissue [6].

We illustrated these different routes to tissue regeneration in Figure 1. Each, as might be expected, has merit in different circumstances. The natural pathway is the medically favored approach in most cases but may not be possible due to the extent of the injury or diseased area. Cell-based therapies for example with respect to spinal cord injuries provide conflicting results and it is now thought that the injection of stem cells does not lead on to differentiation but rather the release of secreting factors, the paracrine effect that promotes the preservation and activity of other cells. The most widely studied tissue engineering technique employs a top-down approach in which the cells are collected, and seeded on or within a porous biodegradable scaffold, with the desired size and shape, and conditioned in an in vitro bioreactor for several weeks where it is exposed to cues including biochemical signals and mechanical stimulation which serve to guide the formation of appropriate tissue type and architecture and ensure fast matrix production [7].

![Figure 1. Pathways for tissue regeneration.](image)

The main advantage of a top-down approach is the prospect of using a variety of biomaterials and the ability to produce porous scaffolds of specific mechanical properties [8]. The use of additive manufacturing [9–11] has proved to be particularly effective in producing scaffolds with the required properties in a shape personalized to the patient and the injury site. However, this approach suffers from some limitations such as positioning cells of multiple types with precision, achieving geometries and cell densities which are tissue specific. An additional limitation is the difficulty of the incorporation of vasculature throughout the 3D space of the tissue constructs [12]. To circumvent these limitations,
‘bottom-up’ strategies have emerged as an alternative for functional tissue engineering, where primitive building blocks composed of cells, materials, and/or bioactive factors are assembled into more complex functional units mimicking the organization and the architecture of native tissues [13]. In this approach, the construction of these primitive building blocks is achieved by different methodologies, including microscale hydrogels (microgels) which encapsulate cells [14], self-assembled cell aggregation [15,16], cell sheets [17], and 3D cell printing [18,19].

2. From In Vitro to In Situ

The environment required for the development of a functional tissue must mimic the native chemical cues from the organism and the biophysical cues to ensure cell survival, migration, proliferation, and differentiation [20]. Even with an appropriate scaffold, TE has a number of limitations such as the availability of a donor tissue biopsy due to restricted anatomical site access and morbidity [21]. Additionally, the tissue derived primary cells require an extensive and complicated cell expansion steps before implantation and these are often heterogeneous and as a consequence difficult to standardize making a successful translation from in vitro to in vivo difficult [19]. In addition to autologous cell source, allogeneic and xenogeneic cell sources can also be used, but they too have their own associated complications such demanding cell manipulation that involves cell isolation and expansion, immunologic responses arising from genetic differences, and the potential transmission of bacteria and viruses from the donor to the host tissue. Stem and progenitor cells can be used as an alternative route, but this is also a complicated ex vivo procedure [22]. The above mentioned limitations makes the conventional tissue engineering approach time consuming and laborious [23] and its commercialization is considered limited due to the combined difficulties of transportation and storage, making it less convenient and in particular less clinically viable [24]. The difficulties and limitations of conventional TE have given rise to a new concept known as “in situ TE” where the body’s own regenerative capability is harnessed to enable it to regenerate and heal [25]. This process is aided through the exploitation of a tissue-specific scaffold that can control the microenvironment at the injured site, that will enable the mobilization of host stem/progenitor cells to these tissues and ensures their subsequent proliferation [19]. In doing so, it eliminates the need for in vitro cell manipulation and offers an alternative therapy path that decreases the time, and resources required to successfully regenerate a tissue or organ [21,26]. It takes advantage of the natural presence of the biochemical and biophysical cues at the injured site of the living tissue by employing the living body as a more effective bioreactor that can regenerate and maintain new tissue at the site of injury in a scalable and cost-effective way [25]. More specifically, in situ TE focuses on using a target specific biomaterial constructs with or without growth factors to recruit host stem or tissue specific progenitor cells to the site of injury (Figure 1), thus allowing the regeneration of the damaged tissue without the need for cell implantation [27,28].

It is worth mentioning that although biomaterials are essential for in situ tissue regeneration, the regenerative capability of the recruited cells is of paramount importance and play a critical part in this process. This regenerative potential is known to vary with age as was clearly shown by Liu et al. [29]. In their study, they showed that in situ bone regeneration with the aid of a biomineralised scaffold was less, by means of lower mineral deposition, in elderly mice. Additionally, it is well known that different tissues regenerate at different rates due to limited blood supply as is the case of cartilage regeneration [30], thus an in situ regeneration using a biomaterial scaffold is expected to follow a similar pattern to that experienced by the native tissue but can be enhanced with the use of additives/surface treatment or specific scaffold design—which is discussed in later sections.

3. Biomaterials: Types and Requirements for In Situ Tissue Regeneration

During in situ tissue regeneration, a tissue-specific biomaterial scaffold, alone or in combination with growth factors, is implanted at the site of the damaged tissue. The biomaterial scaffold is expected to fully occupy the anatomy of the defect site, provide an instructive microenvironment to promote
and recruit host stem cells or progenitor cells in vivo, provide stimulants to trigger native healing cascades, promote differentiation of cells and the proliferation for in situ tissue repair. The tissue also provides structural support until the new tissue is formed without the need for complicated in vitro manipulation [21]. The therapeutic outcome of using the biomaterial scaffold for in situ tissue regeneration thus depends on effective cell homing and control of cell fate [31]. Key to the methodology is the production a scaffold with the required properties tailored to the patient and the shape and volume of the damaged tissue. The properties of the biomaterial scaffold for in situ tissue regeneration may vary depending on the target damaged tissue, but generally will share some common requirements.

3.1. Biocompatibility and Biodegradability

Scaffold biocompatibility and biomimicry is essential for performing and maintaining essential cellular activities which include molecular level and mechanical signaling networks [11]. The incorporation of bioactive features such as biochemical stimuli, mechanical forces, and physiochemical material properties into the chosen biomaterial plays an important role in influencing cell behavior by creating an environment that stimulates a cellular response similar to that of the host tissue [32–34]. Equally important is the biodegradability of a biomaterial, with non-cytotoxic degradation by-products and minimal fibrosis and associated foreign body reaction. The degradation rate is vital for successful in situ tissue regeneration. In this respect, the biomaterial can be designed to provide a dynamic control of the porosity of the scaffold, as well as its mechanical integrity and bioactivity [26]. It has been shown that the degradability rate can influence the mechanical behavior of the scaffold, thus a rapid degradation could lead to early mechanical failure, such as the case of bone tissue regeneration where the scaffold usually endures mechanical loading during the regeneration process. On the other hand, if the degradation process is too slow, complications related to inflammation could also take place [35].

Degradation can occur globally by hydrolysis such as in the case of dextran and poly(lactide-co-glycolide) (PLGA) or by enzymes [36]. Mann et al. showed that photopolymerizable polyethylene glycol (PEG)-based hydrogels are able to be tailored to respond to in situ tissue remodeling by being chemically modified to be degraded by collagenase and elastase, thus permitting cell migration and cell proliferation [37].

3.2. Surface Topography and Chemistry

The scaffolds must possess suitable surface topography and chemistry for the promotion of cell seeding, attachment, and proliferation [27]. It has been shown that surface characteristics play an important role in the extent of cell adhesion and cellular spreading on the surface of biomaterial scaffold [38]. In addition to the type of biomaterial used to fabricate the scaffold, its surface treatment or inclusion of nanoparticles may in fact affect the surface roughness and results in improved scaffold bioactivity [39]. Martins et al. showed that the incorporation of calcium phosphate coatings as hydroxyapatite particles or whiskers, led to an improved bone formation in porous biomaterials [40]. Yavari and co-workers demonstrated that applying a surface treatment to bone scaffolds leads to improved bone mineralization through alterations to the scaffold surface chemistry and roughness [41]. Another interesting example is vascular tissue engineering where it is essential to apply a surface treatment to the fabricated vascular graft as it is considered vital to avoid thrombus formation [42].

3.3. Scaffold Architecture

The scaffolds must have an appropriate architecture to facilitate the flow of nutrients and the formation of capillary blood vessels through infilling, thus regulating cell growth and infiltration as well as recruitment to the injury location. In addition, the structure of the scaffold can also be used to encapsulate bioactive molecules such as growth factors, genes, and cytokines [21], which are
considered crucial for controlling the in vivo microenvironment and thus optimizing the in situ tissue regeneration process by promoting cell-specific recruitment and growth [43,44].

Thevenot et al. have studied the incorporation of biomolecules into biomaterials to enhance cell recruitment. In one study, PLGA scaffolds incorporated with stromal cell-derived factor 1 (SDF-1α—a potent factor for stem cells recruitment) were found to recruit a greater number of stem cells, in order to increase new blood vessel formation, decrease fibrotic response, as well as regulate the inflammatory cell responses compared with pure PLGA scaffolds [45].

Cell seeding, behavior, and response are also found to be significantly influenced by the architectural properties of the biomaterial scaffold. It has been shown that the scaffolds used in bone tissue regeneration must possess optimum pore size, pore shape, porosity, interconnectivity, and fiber-orientation [38,39]. In a wide-ranging review by Bobbert et al. [39] on the architecture of bone substitute materials, it was shown that scaffolds with small pores (200–300 µm) present a better environment for the seeding of cells, however, limit cell sustainability, proliferation, and differentiation. Additionally, it was shown that scaffolds which adopt “ladder-like”, “rectangular pores”, and those with “large spherical pores” are found to be more inclined to fail than those with uniform small and rounded pore structures [5]. For example, Ti6Al4V structures (Figure 2a) fabricated with different shapes of pores exhibited a variation in the mechanical properties and fatigue strength for the range of unit cells with similar porosity (Figure 2). Bobbert et al. [39] concluded that designing what they referred to as a “tortuous void network” would in fact help control the passage of the cell suspension through the scaffold, thus increasing the surface area and available time for cell attachment.

Figure 2. Examples of different pore sizes, shapes, and biomaterials for scaffolds for tissue engineering. (a) Titanium (Ti6Al4V), (b) Starch poly(ε-caprolactone) (SPCL), (c) poly(lactide-co-glycolide) (PLGA), (d) Bioactive glass (BG), (e) poly(propylene fumarate) (PPF), (f) collagen-apatite, (g) Mesoporous bioactive glass (MBG), and (h) Silk fibroin (SF). Reproduced from [39], CC BY 3.0 license.
Sussman et al. [46] demonstrated that when porous biomaterials with an optimum pore size were subcutaneously implanted in mice, the optimum pore size can control the spatial organization of the M1 (pro-inflammatory) or the M2 (pro-healing) macrophage response. Thus, in situ TE can be used as a powerful tool to control inflammatory response, which significantly impacts tissue regeneration in vivo [47]. Dai et al. [48] investigated the use of poly(lactide-co-glycolide), PLGA scaffold with a radially oriented pore architecture, O-PLGA (Figure 3), in an in situ regeneration of osteochondral defects in rabbits. Their results revealed that the O-PLGA scaffold design was shown to enable cell infiltration and bioactive interflow between neo-host tissues more than a randomly porous R-PLGA scaffold in vivo. These results succeeded in emphasizing the significance of using structured scaffolds to optimize the regeneration of native tissues in vivo. Their histological analysis results demonstrated the clear formation of tide mark in the O-PLGA [Figure 4(a3)] in comparison to the R-PLGA [Figure 4(b3)] and a significant presence of chondrocytes regularly spread in the cartilage layer [Figure 4(a4)]. Furthermore, an observable boundary was seen between the two layers, signifying the simultaneous regeneration of both the cartilage and the subchondral bone layers, which is not observed with the R-PLGA scaffold [Figure 4(c1,d1)].

**Figure 3.** Schematic illustration to show (a) the radially oriented pores scaffold, O-PLGA and (b) its implantation in an osteochondral defect in a rabbit model. The blue and red arrows in (b) refer to the bioactive interflow from neighbor cartilage and subchondral bone layers, respectively. PLGA—poly(lactide-co-glycolide). With permission [48].
Figure 4. Assessment of the osteochondral defects regeneration using (a) O-PLGA and (b) R-PLGA for 12 weeks, respectively. (a1,b1) Gross view of the scaffold in the osteochondral bone, (a2,b2) hematoxylin and eosin staining images, and (b3,b3) periodic acid Schiff staining of glycosaminoglycans. Immunohistochemical staining of collagen Type II (a4,b4), collagen Type I (a5,b5), and collagen Type X (a6,b6), respectively. The black arrows in the images are used to indicate the edges of the defects, while the white arrows highlight debris of in the PLGA scaffold. Safranine O and fast green staining of glycosaminoglycans (red) and collagen (green) of O-PLGA (c1) and R-PLGA (d1) respectively. Reproduced with permission from [48].

Arora et al. [49] also showed that scaffolds with oriented pores are considered more suitable and efficient for cell growth, nutrients and waste exchange, and regular deposition of ECMs. Additionally,
such architecture offers the ideal setting for the critical cell–cell, cell–matrix, and neo-host tissue interaction, and improves the mechanical properties in comparison to randomly porous scaffolds. In general, the tendency towards oriented-porous scaffolds stems from the need to mimic natural tissues such as tendons, cartilage, bone tissue, nerves, etc., all of which naturally possess an oriented structure that helps enhance their biological and mechanical functions [50], hence oriented-structures are more favorable when designing a scaffold for in situ tissue regeneration.

3.4. Mechanical Properties

The scaffold must have equivalent mechanical properties to the native tissue in order to offer effective support during the regeneration process [28]. It has been shown that the modulus of the biomaterial significantly influences cell adhesion, distribution, proliferation, and differentiation as was demonstrated by the work on highly elastic poly-L-lactic acid (PLLA) constructs where chondrogenic differentiation was significantly improved in comparison to the rigid poly(lactic acid) (PLA) and PLGA ones [51]. Bhumiratana et al. [52] incorporated hydroxyapatite in silk sponges in an effort to fabricate a composite scaffold that stimulates the formation of mineralized bone tissue by human mesenchymal stem cells (hMSC). Their results showed enhanced mechanical properties by two mechanisms, one was attributed to the osteoconductive nature of the hydroxyapatite and the second to the fact that it provided nucleation sites for the mineral phase. Additionally, in the case of bone defects, it is reported that the scaffold stiffness has an influence on its integration into the local native tissue especially when there is a significant difference in the stiffness. For example, scaffolds composed of polymers would certainly have a stiffness that is different to those composed of metal or ceramics and as a result of this mechanical difference, stress and deformation profiles would also be different throughout the scaffold [53,54].

3.5. Handling Properties

Biomaterials utilized for in situ tissue regeneration can be designed to be adaptable so as to better serve the necessities of the damaged native tissue such as scaffolds, hydrogels, membranes, tubes, micro and nano spheres [55–59], and use manufacturing methods, such as additive manufacturing, that are cost-effective [10,11]. From a surgical perspective, it is desirable for the biomaterial to be easily manipulated into different forms and sizes to allow in situ tissue regeneration tailored to the individual patient and their injuries as in the case of bone defect treatments [35,60]. When hydrogels are used for in situ tissue regeneration it is important that they display optimum injectability properties relevant to the surgical procedure as was highlighted in a recent study by Steele et al. [61]. In this study, they evaluated in vivo biocompatibility and hemocompatibility using an animal model where they performed subcutaneous and intramuscular injections of hyaluronic acid–nanoparticle (HA–NP) hydrogels and compared the results to a PBS control. Their findings revealed no significant difference in immunological response or the presence of fibrosis. They also examined the hemocompatibility of the hydrogel for cardiovascular applications, including intramyocardial delivery, and reported that intracardiac injection of HA–NP hydrogel in rats resulted in insignificant gross or histological changes in comparison to PBS injected-rats. It may well be that the recent developments in 4D printing are able to contribute to the design of scaffolds to deliver therapies without major surgical intervention [62].

There are three classes of biomaterials used in in situ tissue regeneration: polymers (natural and synthetic), bioceramics, and ECM-based biomaterials.

4. Biomaterials

4.1. Synthetic Polymers

Synthetic biomaterials for use in in situ tissue regeneration are based on degradable synthetic polymers and are fabricated using various assembly approaches in order to produce structures with enhanced physical and mechanical properties such as stiffness, degradation, and porosity [63]. One of
the advantages of using synthetic biomaterials is the ability to produce patient-specific scaffolds to match the target anatomy as well as fit the required physical and chemical properties of the injured tissue [64]. In this way, synthetic polymers have no immunological concerns [65] and a higher degree of processing flexibility, thus it is easier to program their biodegradation rate, mechanical properties and microstructure [66]. Talacua and co-workers investigated the use of resorbable poly(ε-caprolactone) (PCL) grafts as capillary blood vessels, which are filled with a fibrin gel and the monocyte chemoattractant protein 1, in a rat model of abdominal aortas. After 3 months in situ, their study revealed higher structured systems with a layer on the inside formed from elastin fibres coated with endothelium cells, as well as a layer of smooth muscle cells at the centre of the structure [67]. Moreover, synthetic biomaterials can be designed to have functional sites by coupling with biomolecules [68] and there is less biosafety concerns in particular with regards to the host tissue response and disease transfer in comparison to natural polymers [69]. Of the interesting groups of biodegradable synthetic polymers used in tissue engineering are aliphatic polyesters such as PLA, PCL, polyglycolide (PGA), and their copolymer PLGA [70]. These synthetic polymers undergo degradation, involving the hydrolysis of the ester groups in their backbone chains, into non-cytotoxic products [71]. More importantly, the properties of this class of biomaterials can be tailored by selecting the molecular weight distribution, the nature of the porous architecture, and the selection of the degradation rate for a specific tissue regeneration application by variation of the monomer composition of copolymers [72]. These factors amongst others will determine the resultant mechanical properties. Two important polymer chemistry tools in this respect are the use of copolymerization and blending to obtain tailored degradation rates. These polymers are among the few synthetic polymers already approved by the US Food and Drug Administration (FDA) for clinical applications such as surgical sutures and some implantable devices [68]. One of the most common synthetic biomaterials is PLGA where it is extensively used in bone and cartilage tissue regeneration [73]. However, conventional synthetic polymers have disadvantages due to the lack of functional groups resulting in their reduced capability to combine with bioactive moieties to strengthen their cell affinity [12]. Functional synthetic polymers have gained popularity due to their facile design and modification which enables them to provide bioactive signals to enhance the spatiotemporal cell-biomaterial interaction [74]. Other examples of synthetic biomaterials and their applications are shown in Table 1.

| Application | Tissue | Biomaterial * | Animal Model | Reference |
|-------------|--------|---------------|--------------|-----------|
| Bone        | PLA    | Rat calvarial bone defect | Gómez et al. [75] |
|             | P(HEMA) | Rabbit femoral defect | Mabilleau et al. [76] |
|             | PEUU   | Rat myocardial infraction model | Fujimoto et al. [77] |
| Cartilage   | PGA    | Sheep cartilage defect | Erggelet et al. [78] |
|             | PLGA   | Rabbit articular osteochondral defect | Dai et al. [48] |
| Heart Valve | Polycarbonate bis-urea (PC-BU) | Sheep pulmonary valve | Kluin et al. [23] |
|             | Polyester-urethane | Sheep aortic valve | Yosuke et al. [79] |
| Periodontal Tissue | PLGA | Canine periodontal | Herberg et al. [80] |
| Blood Vessels | PLLA/PCL | Rat abdominal aorta | Jiang et al. [81] |

* PLGA—poly(lactide-co-glycolide), PEUU—polyester urethane urea, PCL—poly(ε-caprolactone), PGA—poly(glycolic acid), PLGA—poly(lactic acid), P(HEMA)—poly(2-hydroxyethyl methacrylate), PLLA—poly(l-lactic acid).

Synthetic polymers enable scaffolds to be manufactured with precision and with controlled architecture and target mechanical properties. There are a limited number of polymers which are FDA approved for clinical applications.
4.2. Natural Polymers

Natural polymers, derived from polysaccharides and proteins, are types of biomaterials that have excellent biodegradable and biocompatible properties as well as other features mimicking the extracellular matrix (ECM) making them a very attractive choice for TE applications [31]. Popular polysaccharide-based natural polymers that have been widely used for in situ tissue regeneration include fibrin, alginate, hyaluronic acid and chitosan [82]. Chitosan, which is derived from chitin, is the second most abundant biosynthesized biomaterial [83]. Chitosan exhibits a cationic nature which makes it an ideal system for the delivery of anionic glycosaminoglycans, growth factors, cytokines, and genes [84]. Another example is hyaluronic acid (HA), which is found in the extracellular matrix of many tissues, is widely used in laboratory studies of cartilage repair due to its immune and biological responses including proliferation, morphogenesis, and wound repair [85]. It can be easily modified to form hydrogels that are known to promote better cartilage regeneration than hydrogels based on polyethyleneglycol PEG hydrogels [86] and used to deliver growth factors such as VEGF and bone morphologic protein (BMP) [87,88].

Proteins, such as collagen, fibrin, gelatin and silk, form the other class of natural-based polymers [89, 90]. Collagen, the most abundant type of proteins in humans, is an FDA-approved biomaterial due to its ease of processing and minimal inflammatory and immune responses [20], thus frequently used in biomedical applications such as wound dressings and artificial skin [91]. Shi and co-workers [92] demonstrated that the conjugation of collagen scaffolds with antibodies specific to stem cells could capture stem cells at a wound site and promote cardiomyocyte regeneration in a mouse model. Other examples of protein-based natural polymers are shown in Table 2.

More recently, Wu and co-workers fabricated an electrospun silk fibroin scaffold and used Polydopamine (PDA) to modify the surfaces of the scaffolds which were further modified using E7, a newly discovered peptide with specific affinity for bone marrow mesenchymal stem cells (BMSCs) in a rat calvarial bone defect model [93]. Their findings showed that these enhanced electrospun scaffolds had led to improved hydrophilicity, enabling cell proliferation and adhesion, and furthermore boosted the osteogenic differentiation of BMSCs by creating favorable osteoinduction conditions under the synergistic effects of PDA and E7 both in vitro and in vivo.

However, there are some shortcomings to the use of natural biomaterials such as relatively poor mechanical properties limiting their applications in anatomical sites with demanding mechanical loading such as hard tissue regeneration [94]. Thus, in order to improve such properties, natural polymers are often combined with synthetic ones to produce hybrid biomaterials that enjoy the advantages of both classes of polymers without compromise (Table 3).
Table 2. Recent examples of natural polymer biomaterials used for in situ tissue regeneration.

| Application Tissue | Biomaterials | Animal Model | Reference                |
|--------------------|--------------|--------------|--------------------------|
| Bone               | Silk fibroin | Rat critical size calvarial bone defect | Wu et al. [93]          |
|                    | Chitosan/Silk-Fibrin | Rat calvarial bone defect model | Wu et al. [95]        |
|                    | Fibrin       | Rat cranial defect | Woodruff et al. [96]  |
|                    | Gelatin      | Mouse maxillae | Kodama et al. [97]    |
|                    | Gelatin      | Rat calvarial bone defect model | Feng et al. [98] |
| Kidney             | Collagen     | Renal ischemia/reperfusion rat model | Lee et al. [99] |
| Heart and vessel   | Alginate     | Rat myocardial infraction model | Landa et al. [100] |
| Cartilage          | Collagen     | Rabbit articular cartilage | Kubo et al. [101] |
|                    | Fibrin       | New Zealand white rabbit full thickness cartilage defect | Dai et al. [102] |
|                    | Alginate     | Rabbit cartilage defect | Ma et al. [103] |
| Muscle             | Collagen     | Rabbit muscle | Kin et al. [104] |
|                    | Gelatin      | Rat muscle | Ju et al. [21] |
|                    | Collagen     | Rat diaphragm defect | Brouwer et al. [105] |
| Periodontal tissue | Collagen     | Canine periodontal | Nakahara et al. [106] |
| Skin               | Chitosan     | Porcine burned skin | Boucard et al. [107] |
|                    | Hyaluronic acid-HA | Mouse cutaneous wound model | Wang et al. [108] |

Table 3. Hybrid biomaterials used for in situ tissue regeneration.

| Application Tissue | Biomaterial * | Animal Model | Reference            |
|--------------------|---------------|--------------|----------------------|
| Bone               | Fibrin/PLGA   | Rat calvarial bone defect | Chung et al. [109] |
|                    | Chitosan/β-sodium glycerol phosphate (CS/GP) | Rat critical size calvarial bone defect | Wu et al. [110] |
|                    | PCL-PDA-HAp   | Mouse critical size calvarial bone defect | Lee et al. [111] |
| Cartilage          | HA-GelMa      | Sheep model | Di Bella et al. [112] |
|                    | Collagen-HAP  | Human osteochondral defect | Perdisa et al. [113] |
|                    | HA-MA/PLGA    | Rabbit full thickness cartilage defect | Dai et al. [114] |
| Blood Vessels      | PCL/fibrin    | Rat model aorta | Talacua et al. [67] |
| Heart Valve Leaflets | P4HB/Gelatin | Ovine model pulmonary valve | Capulli et al. [115] |
| Stomach            | Collagen/PGA  | Canine stomach | Hori et al. [116] |
| Spine              | PGA/HA        | Rabbit disc defect | Abbushi et al. [117] |

* ECM—extracellular matrix, PLGA—poly(lactide-co-glycolide), HA—hyaluronic acid, PEUU—polyester urethane urea, PCL—poly(ε-caprolactone), PGA—poly(glycolic acid), PLA—poly(lactic acid), HEMA2—hydroxyethyl methacrylate gelatin methacrylamide (GelMa), HA-GelMa—gelatin methacrylamide (GelMa) and hyaluronic acid methacrylate (HAMA), Hap—hydroxyapatite, P4HB—poly-4-hydroxybutyrate.

Natural polymers offer the advantage that they may come closer to mimicking the ECM, but suffer from the difficulty of processing the material into the required shapes whilst maintaining the biological function.

4.3. ECM

ECM-based biomaterials are derived from decellularized ECM (dECM) tissues and these have been extensively explored for in situ tissue regeneration due to the fact that they provide an environment that
is very similar to that of the native tissue ECM especially where cellular behavior is concerned [118]. Decellularized extracellular matrix is naturally the closest scaffold to nature, retaining its unique micro- and macro-structural design features as well as its complex composition [119]. The decellularization process preserves the structural integrity and thus ensures an intact vasculature [120].

Decellularized urinary bladder matrix and small intestinal submucosa are used as a source tissue for ECM-based biomaterials [121]. They are reported to maintain the necessary bioactive features to stimulate positive remodeling effects when implanted in vivo. These bioactivities include recruitment/differentiation of endogenous stem/progenitor cells and the modulation of the host immune response. One example was demonstrated with a study conducted by Dziki and co-workers [122] in patients \( (n = 13) \) who were treated for a volumetric muscle loss with porcine-derived ECM sheets harvested from either the urinary bladder, the small intestine, or the dermis. These sheets of ECM materials were placed in contact with the healthy tissue for subsequent in situ remodeling. Six months after implantation, and regardless of the type of ECM employed, their findings demonstrated that patients improved their strength by an average of \( \sim 37\% \) and showed \( \sim 27\% \) improvement in range-of-motion tasks in comparison with pre-operative performance. The results further showed that these ECM-sheets promoted the local recruitment of perivascular stem cells with the downstream formation of new functional skeletal muscle. More examples are shown in Table 4. The main disadvantage of dECM is donor shortage. However, ECM-based biomaterials can also be derived from in vitro cultured cells [123] and these have already been successfully used clinically for bone and cartilage regeneration [124]. In another study, Lih et al. fabricated biomimetic PLGA/ECM of decellularized porcine kidney scaffolds for kidney tissue regeneration in a mouse model [125]. Their results showed an increased number of kidney-related cells in the PLGA/ECM group as compared to the PLGA control, and the scaffold degradation rate was also found to be faster in PLGA/ECM scaffolds than in the control PLGA scaffolds. They further demonstrated that ECM-derived bioactive molecules in PLGA/ECM scaffolds have encouraged cell growth and metabolic acids. The hydrolytic degradation of PLGA and the enzymatic degradation of the ECM were both accelerated through the actions of enzymes secreted by active living cells. Therefore, in comparison to the control PLGA scaffold, the rapidly biodegradable PLGA/ECM scaffold is thought to be a more favorable microenvironment for cell proliferation, migration, and recruitment.

Some of the reported problems identified with the use of decellularized tissue scaffolds is the accelerated degeneration with lack of cell repopulation and remodeling evidence [126]. Recently, Dai et al. fabricated a scaffold of a porous matrix metalloproteinase degradable poly (ethylene glycol) hydrogel and decellularized porcine aortic valve in a rat subdermal model [127]. The hydrogel was loaded with stromal cell-derived factor-1α (SDF-1α) which, consequently, served to turn the decellularized scaffold biologically active. Such bioactivity caused by the SDF-1α is thought to stimulate the in vivo tissue recellularization, by attracting progenitor cells from the bloodstream, modulating immune responses and being pivotal in tissue repair, remodeling and regeneration [128]. Another study also demonstrated that SDF-1α could control valve cell phenotype and is involved in scaffold recellularization and remodeling by stimulating the attraction of stem cells [129]. Dai et al. results demonstrated that their hydrogel surface layers provided a niche for cell activities and helped protect the decellularized scaffold from rapid degeneration, inflammation, and calcification resulting in an improved recellularization and remodeling processes of the implanted decellularized heart valves [127].
Table 4. Examples of decellularized extracellular matrix (dECM)-based scaffolds for in situ tissue regeneration

| Application | Tissue | Biomaterials | Animal Model | Reference |
|-------------|--------|--------------|--------------|-----------|
| Esophagus   | Urinary Bladder Matrix—UBM | Rat Abdominal Esophagus | Keane et al. [130] |
| Esophagus   | Porcine acellular dermal matrix (PADM) | In humans | Katerinakiet al. [131] |
| Abdominal wall | Porcine-derived tissue matrix | Primates | Connoret al. [132] |
| Kidney     | PLGA/dECM of porcine kidney tissue | Mouse | Li et al. [125] |
| Skin       | Porcine subcutaneous adipose tissue | Mouse subcutaneous model | Tan et al. [133] |
| Hemilarynx | Porcine UBM | Canine model | Kitamura et al. [134] |
| Heart Valve | PEG/decellularized porcine aortic valve | Rat subdermal model | Dai et al. [127] |
|            | PCL/decellularized porcine aortic valve | Rat subcutaneous model | Zhou et al. [135] |

ECM and dECM based materials naturally reflect closely the ECM of the native tissue, but as with natural polymers they face challenges in manufacturing scaffolds with a defined macro-architecture.

5. Bioceramics

Bioceramics are a class of ceramic materials specifically used for the repair and reconstruction of damaged tissues of the body. Bioceramics are categorized into the following groups: nearly inert (based on alumina and zirconia), bioactive (based on bioactive glass), and resorbable ceramics (based on $\beta-$ and $\alpha-$tricalcium phosphate [136]. The most commonly employed bioceramics for in situ TE are those used for bone tissue regeneration such as hydroxyapatite (HAp), calcium phosphates (CP), and tri-calcium phosphate (TCP) with different calcium to phosphate ratio and crystallographic structure, which in turn influences the degree of their solubility [137]. Such bioceramics clearly demonstrate good biocompatibility, bioactivity, osteoconductivity [12], and osteoinductivity that are essential for the initiation of in situ bone regeneration even without the use of inductive factors [138]. Additionally, it has been reported that Ha and TCP present no immunogenicity or toxic side effects [139].

In bone tissue regeneration, Samavedi et al. showed that certain CP ceramics are osteoinductive and that their osteoinduction property depends on the surface chemistry, which can consequently influence protein adsorption and promote cell differentiation through cell–ECM interactions [140]. However, present CP ceramics still face many challenges to meet the requirements of regenerative medicine, which include the relatively low bioactivity of the ceramic in comparison to the natural bone tissue, degradation kinetics which do not match new bone formation, and exhibiting poor machining performance to process them into specific shapes [141]. Additionally, a critical problem which restricts their wide clinical application is their poor mechanical property and brittleness. Their use to date is limited to either non-loadbearing implants, such as HAPEX as a middle ear implant [142] and bone defect filling material in the oral cavity and skeleton, or as coating on dental and orthopedic metallic implants [143]. This mechanical inferiority can be overcome by creating polymer and ceramic composites in a variety of forms such as nanofibers, foams, hydrogels, and 3D printed scaffolds for tissue-specific regeneration [144,145]. The combination of polymers and ceramics not only improves the mechanical property of the final construct but also offers biological benefits as was demonstrated by Meka et al. who showed that the inclusion of bioactive ceramic nanoparticles in the PCL composite scaffolds could possibly substitute the use of expensive bone stimulating growth factors [146]. Song and co-workers implanted biphasic calcium phosphate (BCP) granules in the dorsal muscle of lumbar region in a canine model to demonstrate how bone-marrow-derived mesenchymal cells (BMSCs)
migrated from bone marrow to where new bone formation was induced without the introduction of growth factors [147]. In another study, Wang and co-workers [148] investigated the effect of chemical composition on protein (BMP-2) adsorption and osteoinductive potential in four types of bioceramics: hydroxyapatite (HAp), β-TCP, biphasic calcium phosphate-1(BCP-1), and biphasic calcium phosphate-2(BCP-2) (the ratios of HAp to β-TCP in BCP-1 and BCP-2 were about 70%/30% and 30%/70%) in a Balb/c mouse model. Histological and histomorphometric analyses of their results revealed that porous BCP-2 demonstrated a stronger osteoinductive capability than the other three groups of ceramics. The highest expression of BMP-2 and osteocalcin (OCN) was also achieved in the BCP-2 group. The amount of BMP-2 present in the local microenvironment of the implant is thought to influence the osteoinduction of porous CaP ceramics (Figure 5). Recently, Li et al. fabricated nanocrystalline of biphasic Calcium Phosphate (BCP-N) granules by combining alginate gelatinization with microwave hybrid sintering methods, and investigated the in vivo osteogenesis potential of the resultant materials in a rabbit mandible critical-size bone defect [149]. Their results showed that the nanotopography in BCP-N might be responsible for the stronger osteoinductivity and bone regenerative ability than exhibited by microcrystalline biphasic CP (BCP-G) and a commercially available type (BAM®, National Engineering Research Center for Biomaterials of Sichuan University, China).

Figure 5. Photomicrographs of hematoxylin and eosin staining for the decalcified sections of porous Calcium Phosphate-based ceramics implanted into the thigh muscles of mice for 45 and 90 days. Yellow markers indicate the newly formed bone. Bar, 200 µm, reproduced with permission from [148].
Resorbable ceramics such as β- and α-TCP have the advantage of degrading gradually over a period of time while the natural host tissue regenerates itself and replaces the degrading scaffold [150]. On the other hand, hydroxyapatite (HAp) possess a porosity similar to that of native bone tissue thus promoting bone growth within the pores of the scaffold [151]. Macroporous HAp scaffolds with interconnected oval shaped pores have been found to exhibit enhanced cellular functionality and supports osteoclast differentiation in comparison to the results found for dense HAp scaffolds [152]. Additionally, it has been reported that during the degradation process of HAp, calcium and phosphate ions are released which help to induce osteogenic response and contribute to the osteoinductivity of the HAp scaffold [153]. However, its use as a scaffold material is limited because of its low mechanical properties and particularly slow rate of resorption [154]. Table 5 shows further examples of bioceramics.

| Application Tissue | Biomaterials * | Animal Model | Reference |
|--------------------|----------------|--------------|-----------|
| Sinus Mucosa       | CP             | Rabbit sinus lift model | Trbakovic et al. [155] |
| Bone               | BCP-N          | Rabbit mandible critical size defect model | Li et al. [149] |
| DNA-loaded nano-calcium phosphate | New Zealand white rabbits, critical size bone defect model | Schickewei et al. [156] |
| PLA/CP            | Mouse subcutaneous model | Oliveira et al. [157] |
| Calcium-silicate   | Rabbit mandibular alveolar bone defect model | Shao et al. [158] |
| Alginate* CSi-Sr₄ and CaP | New Zealand rabbits, distal femur defect | Fu et al. [159] |

* CP (calcium phosphate), BCP-N (nanocrystalline biphasic calcium phosphate), PLA (poly(lactic acid)), CSi-Sr₄ (strontium-substituted calcium silicate), CaP (beta-tricalcium phosphate).

Bioceramics bring a substantial history of applications in medicine and it has been clearly demonstrated that it is possible to generate scaffolds with precise complex macro-architectures. Clearly in some areas such as in bone regeneration they offer advantages of similarity in terms of chemical composition and some of the key properties of bone such as conductivity and surface topography.

The choice of material for a scaffold is largely determined by the requirement of tissue equivalent. Bioceramics can present a valuable option in the case of bone regeneration with the potential for producing a hierarchal structure at the macro, micro, and nano-levels, thus mimicking the native tissue and ensuring osteoinductivity. However, an extension of this to other tissue types is clearly inappropriate. By combining bioceramics with polymers there is a greater flexibility to achieve tissue equivalence as well as improving the overall mechanical properties. This approach has been explored by several researchers [160].

6. Future Prospects

In situ tissue engineering extends the family of therapies available for regenerative medicine in the field of regenerative medicine. The in situ approach avoids the necessity to remove cells from a patient and to place them in environments such as a bioreactor where the physiological environment may lead to a loss of function or the delivery of inappropriate signals. An often stated problem with the ex-situ TE process is the long delays between cell sampling and implantation especially in, for example, coronary interventions with the need for the development of small vascular grafts [161].

This review has shown that much of what has been learnt in terms of biomaterials for the development of scaffold-based ex-situ tissue engineering can in part be transferred to in situ tissue engineering and so the development of new therapies may be expedited. It is also the case that new developments in additive manufacturing will be particularly helpful in the development of realistic scaffolds for use with in situ tissue engineering which reflect the actual complexity of the extracellular matrix and the environments required for regeneration. One of the emerging developments in additive
manufacturing is 4D printing [62], in which a 3D object is manufactured with a specific form and using an external stimulus, such as temperature, pH, electric and magnetic fields, which is used to initiate a transformation of the material such that the form of the object changes to an alternative design. We can foresee such materials and concepts of particular value in the design of therapies which do not need major surgical intervention to deliver in situ tissue engineering.

In all scaffold-based tissue engineering, much is made of degradable or resorbable materials especially polymers, and although such concepts are central to regenerative medicine, there appears to be little work report on how to design such materials of scaffolds to meet the personalized needs of specific patients and injuries. We see that much work is focused on biopolymers and other natural materials and although these may have advantages in biocompatibility and with degradation, they present many challenges in precisely shaping the scaffold to the particular site of the required therapy and its personalization to the patient. The hybrid nature of bone means that bioceramics present interesting opportunities linked to the need for mineralization and the generation of a structure which mimics bone. This is particular the case for bone generation with therapies to cope with bone disease, especially osteoporosis.

Most publications in this area highlight the possibility of tailoring the properties of the scaffolds to match the tissue environment but there appears to be limited work which is focused on predictive modelling which can be used to support material innovation and scaffold design.

As a number of authors identify, regeneration medicine has created much promise but delivered little outside the scope of laboratory-based experiments (EUROPEAN INITIATIVE FOR REGENERATIVE MEDICINE. Final report 2016 A Webster: https://ec.europa.eu/futurium/en/content/european-initiative-regenerative-medicine downloaded September 2019). The development of in situ tissue engineering therapies which do not require significant surgical intervention using 4d printed scaffolds may provide the much needed stimulus to this field.

7. Conclusions

In situ tissue engineering extends the family of therapies available for regenerative medicine in the field of bone regeneration. Much of the understanding of scaffolds and scaffold materials on scaffold-based ex situ tissue engineering may be in part transferred to the in situ methodology, perhaps enabling a faster development cycle. New developments in 4d printing provide specific opportunities for the development of therapies without major surgical intervention. The design of the scaffolds and the materials selected remain at the heart of developments in this field and there is a clear need for predictive modelling which can be used to optimize the design of materials and scaffolds.

Funding: This research was funded by FCT (Portugal), grant number POCI-01-0145-FEDER-031146.

Acknowledgments: We would like to thank Mr. Renato Batista at CDRSP for his assistance with some of the artwork.

Conflicts of Interest: The authors declare no conflict of interest.

References

1. Kung, J.W.C.; Currie, I.S.; Forbes, S.J.; Ross, J.A. Liver Development, Regeneration, and Carcinogenesis. J. Biomed. Biotechnol. 2010, 2010, 8. [CrossRef]
2. Atala, A. Tissue engineering and regenerative medicine: Concepts for clinical application. Rejuvenation Res. 2004, 7, 15–31. [CrossRef] [PubMed]
3. Skalak, R.; Fox, C.F. Tissue Engineering; Alan, R., Ed.; Liss. Inc.: New York, NY, USA, 1988.
4. Vacanti, J.P.; Langer, R. Tissue engineering: The design and fabrication of living replacement devices for surgical reconstruction and transplantation. The Lancet 1999, 354, 32–34. [CrossRef]
5. Mi, H.-Y.; Jing, X.; Salick, M.R.; Cordie, T.M.; Peng, X.-F.; Turng, L.-S. Morphology, mechanical properties, and mineralization of rigid thermoplastic polyurethane/hydroxyapatite scaffolds for bone tissue applications: Effects of fabrication approaches and hydroxyapatite size. J. Mater. Sci. 2014, 49, 2324–2337. [CrossRef]
6. O’Brien, F.J. Biomaterials & scaffolds for tissue engineering. Mater. Today 2011, 14, 88–95.
7. Ebrahimi, M. 13-Biomimetic principle for development of nanocomposite biomaterials in tissue engineering. In Applications of Nanocomposite Materials in Orthopedics Inamuddin; Asiri, A.M., Mohammad, A., Eds.; Woodhead Publishing: Sawston, Cambridge, CA, USA, 2019; pp. 287–306.
8. Khademhosseini, A.; Langer, R. A decade of progress in tissue engineering. Nat. Protoc. 2016, 11, 1775. [CrossRef] 
9. Lin, K.; Sheikh, R.; Romanazzo, S.; Roohani, I. 3D Printing of Bioceramic Scaffolds-Barriers to the Clinical Translation: From Promise to Reality, and Future Perspectives. Materials 2019, 12, 2660. [CrossRef] 
10. Miar, S.; Shafiee, A.; Guda, T.; Narayan, R. Additive Manufacturing for Tissue Engineering. In 3D Printing and Biofabrication; Ovsianikov, A., Yoo, J., Mironov, V., Eds.; Springer International Publishing: Cham, Switzerland, 2018; pp. 3–54.
11. Shick, T.M.; Abdul Kadir, A.Z.; Ngadiman, N.H.A.; Ma’aram, A. A review of biomaterials scaffold fabrication in additive manufacturing for tissue engineering. J. Bioact. Compat. Polym. 2019. [CrossRef] 
12. Cheng, C.W.; Solorio, L.D.; Alsberg, E. Decellularized tissue and cell-derived extracellular matrices as scaffolds for orthopaedic tissue engineering. Biotechnol. Adv. 2014, 32, 462–484. [CrossRef] 
13. Jeon, O.; Lee, Y.B.; Hinton, T.J.; Feinberg, A.W.; Alsberg, E. Cryopreserved cell-laden alginate microgel bioink for 3D bioprinting of living tissues. Mater. Today Chem. 2019, 12, 61–70. [CrossRef] 
14. Newsom, J.P.; Payne, K.A.; Krebs, M.D. Microgels: Modular, tunable constructs for tissue regeneration. Acta Biomater. 2019, 88, 32–41. [CrossRef] [PubMed] 
15. Fernando, W.A.; Papantoniou, I.; Mendes, L.F.; Hall, G.N.; Bosmans, K.; Tam, W.L.; Teixeira, L.M.; Moos, M., Jr.; Geris, L.; Luymen, F.P. Limb derived cells as a paradigm for engineering self-assembling skeletal tissues. J. Tissue Eng. Regen. Med. 2018, 12, 794–807. [CrossRef] [PubMed] 
16. Patel, A.; Xue, Y.; Hartley, R.; Sant, V.; Eley, J.R.; Cui, X.T.; Stolz, D.B.; Sant, S. Hierarchically aligned fibrous hydrogel films through microfluidic self-assembly of graphene and polysaccharides. Biotechnol. Bioeng. 2018, 115, 2654–2667. [CrossRef] [PubMed] 
17. Kobayashi, J.; Akiyama, Y.; Yamato, M.; Shimizu, T.; Okano, T. Design of Temperature-Responsive Cell Culture Surfaces for Cell Sheet-Based Regenerative Therapy and 3D Tissue Fabrication. In Cutting-Edge Enabling Technologies for Regenerative Medicine; Chun, H.J., Park, C.H., Kwon, I.K., Khang, G., Eds.; Springer Singapore: Singapore, 2018; pp. 371–393.
18. Nagarajan, N.; Dupret-Bories, A.; Karabulut, E.; Zorlutuna, P.; Vrana, N.E. Enabling personalized implant and controllable biosystem development through 3D printing. Biotechnol. Adv. 2018, 36, 521–533. [CrossRef] [PubMed] 
19. Whitley, M.; Cereceres, S.; Dhavalikar, P.; Salhador, K.; Wilems, T.; Smith, B.; Mikos, A.; Cosgriff-Hernandez, E. Improved in situ seeding of 3D printed scaffolds using cell-releasing hydrogels. Biomaterials 2018, 185, 194–204. [CrossRef] 
20. Ko, I.K.; Lee, S.J.; Atala, A.; Yoo, J.J. In situ tissue regeneration through host stem cell recruitment. Exp. Amp. Mol. Med. 2013, 45, 57. [CrossRef] 
21. Ju, Y.M.; Atala, A.; Yoo, J.J.; Lee, S.J. In situ regeneration of skeletal muscle tissue through host cell recruitment. Acta Biomater. 2014, 10, 4332–4339. [CrossRef] 
22. Ma, P.X. Biomimetic materials for tissue engineering. Adv. Drug Deliv. Rev. 2008, 60, 184–198. [CrossRef] 
23. Klun, J.; Talacua, H.; Smits, A.I.P.M.; Emmert, M.Y.; Brugmans, M.C.P.; Fioretta, E.S.; Dijkman, P.E.; Söntjens, S.H.M.; Duijvelshoff, R.; Dekker, S.; et al. In situ heart valve tissue engineering using a bioresorbable elastomeric implant – From material design to 12 months follow-up in sheep. Biomaterials 2017, 125, 101–117. [CrossRef] 
24. Augustinste, R.; Dalvi, Y.B.; Dan, P.; George, N.; Helle, D.; Varghese, R.; Thomas, S.; Menu, P.; Sandhyaranir, N. Nanoceria Can Act as the Cues for Angiogenesis in Tissue-Engineering Scaffolds: Toward Next-Generation in Situ Tissue Engineering. ACS Biomater. Sci. Eng. 2018, 4, 4338–4353. [CrossRef] 
25. Cipitria, A.; Boettcher, K.; Schoenhals, S.; Garske, D.S.; Schmidt-Bleeke, K.; Ellinghaus, A.; Dienelt, A.; Peters, A.; Mehta, M.; Madl, C.M.; et al. In-situ tissue regeneration through SDF-1α driven cell recruitment and stiffness-mediated bone regeneration in a critical-sized segmental femoral defect. Acta Biomater. 2017, 60, 50–63. [CrossRef] [PubMed] 
26. Lee, S.J.; Yoo, J.J.; Atala, A. Fundamentals of In Situ Tissue Regeneration. In Situ Tissue Regeneration; Elsevier B.V.: Amsterdam, The Netherlands, 2016.
27. Li, Q.; Ma, L.; Gao, C. Biomaterials for in situ tissue regeneration: Development and perspectives. J. Mater. Chem. B. 2015, 3, 8921–8938. [CrossRef]
28. Kim, B.S.; Baez, C.E.; Atala, A. Biomaterials for tissue engineering. World J. Urol. 2000, 18, 2–9. [CrossRef] [PubMed]
29. Liu, M.; Nakasaki, M.; Shih, Y.-R.V.; Varghese, S. Effect of age on biomaterial-mediated in situ bone tissue regeneration. Acta Biomater. 2018, 78, 329–340. [CrossRef] [PubMed]
30. Abdulghani, S.; Morouço, P.G. Biofabrication for osteochondral tissue regeneration: Bioink printability requirements. J. Mater. Sci. Mater. Med. 2019, 30, 20. [CrossRef] [PubMed]
31. Xia, H.; Li, X.; Gao, W.; Fu, X.; Fang, R.H.; Zhang, L.; Zhang, K. Tissue repair and regeneration with endogenous stem cells. Nat. Rev. Mater. 2018, 3, 174–193. [CrossRef]
32. Williams, D.F. On the mechanisms of biocompatibility. Biomaterials 2008, 29, 2941–2953. [CrossRef]
33. Stevens, M.M.; George, J.H. Exploring and Engineering the Cell Surface Interface. Science 2005, 310, 1135. [CrossRef]
34. Crowder, S.W.; Leonardo, V.; Whittaker, T.; Papathanasiou, P.; Stevens, M.M. Material Cues as Potent Regulators of Epigenetics and Stem Cell Function. Cell Stem Cell 2016, 18, 39–52. [CrossRef]
35. Turnbull, G.; Clarke, J.; Picard, F.; Riches, P.; Jia, L.; Han, F.; Li, B.; Shu, W. 3D bioactive composite scaffolds for bone tissue engineering. Bioact. Mater. 2018, 3, 278–314. [CrossRef]
36. Sengupta, D.; Waldman, S.D.; Li, S. From In Vitro to In Situ Tissue Engineering. Ann. Biomed. Eng. 2014, 42, 1537–1545. [CrossRef] [PubMed]
37. Mann, B.K.; Gobin, A.S.; Tsai, A.T.; Schmedlen, R.H.; West, J.L. Smooth muscle cell growth in photopolymerized hydrogels with cell adhesive and proteolytically degradable domains: Synthetic ECM analogs for tissue engineering. Biomaterials 2001, 22, 3045–3051. [CrossRef]
38. Guijak, F.; Cohen, D.M.; Estes, B.T.; Gimble, J.M.; Liedtke, W.; Chen, C.S. Control of Stem Cell Fate by Physical Interactions with the Extracellular Matrix. Cell Stem Cell 2009, 5, 17–26. [CrossRef] [PubMed]
39. Bobbert, F.S.L.; Zadpoor, A.A. Effects of bone substitute architecture and surface properties on cell response, angiogenesis, and structure of new bone. J. Mater. Chem. B. 2015, 3, 6172–6192. [CrossRef]
40. Martins, A.M.; Alvés, C.M.; Reis, R.L.; Mikos, A.G.; Kasper, F.K. Toward Osteogenic Differentiation of Marrow Stromal Cells and In Vitro Production of Mineralized Extracellular Matrix onto Natural Scaffolds. In Biological Interactions on Materials Surfaces: Understanding and Controlling Protein, Cell, and Tissue Responses; Puleo, D.A., Bizios, R., Eds.; Springer US: New York, NY, USA, 2009; pp. 263–281.
41. Amin Yavari, S.; van der Stok, J.; Chai, Y.C.; Wauthle, R.; Tahmasebi Birgani, Z.; Habibovic, P.; Mulier, M.; Schrooten, J.; Weinans, H.; Zadpoor, A.A. Bone regeneration performance of surface-treated porous titanium. Biomaterials 2014, 35, 6172–6181. [CrossRef]
42. Ravi, S.; Chaikof, E.L. Biomaterials for vascular tissue engineering. Regen. Med. 2009, 5, 107–120. [CrossRef]
43. Thevenot, P.T.; Nair, A.M.; Shen, J.; Lotfi, P.; Ko, C.-Y.; Tang, L. The effect of incorporation of SDF-1α into PLGA scaffolds on stem cell recruitment and the inflammatory response. Biomaterials 2010, 31, 3997–4008. [CrossRef]
44. Bai, X.; Gao, M.; Syed, S.; Zhuang, J.; Xu, X.; Zhang, X.-Q. Bioactive hydrogels for bone regeneration. Bioact. Mater. 2018, 3, 401–417. [CrossRef]
45. Agarwal, R.; García, A.J. Biomaterial strategies for engineering implants for enhanced osseointegration and bone repair. Adv. Drug Deliv. Rev. 2015, 94, 53–62. [CrossRef]
46. Sussman, E.M.; Halpin, M.C.; Muster, J.; Moon, R.T.; Ratner, B.D. Porous Implants Modulate Healing and Induce Shifts in Local Macrophage Polarization in the Foreign Body Reaction. Ann. Biomed. Eng. 2014, 42, 1508–1516. [CrossRef]
47. Brown, B.N.; Londono, R.; Tottey, S.; Zhang, L.; Kukla, K.A.; Wolf, M.T.; Daly, K.A.; Reing, J.E.; Badyak, S.F. Macrophage phenotype as a predictor of constructive remodeling following the implantation of biologically derived surgical mesh materials. Acta Biomater. 2012, 8, 978–987. [CrossRef] [PubMed]
48. Dai, Y.; Shen, T.; Ma, L.; Wang, D.; Gao, C. Regeneration of osteochondral defects in vivo by a cell-free cylindrical poly(lactide-co-glycolide) scaffold with a radially oriented microstructure. J. Tissue Eng. Regen. Med. 2018, 12, 1647–1661. [CrossRef] [PubMed]
49. Arora, A.; Kothari, A.; Katti, D.S. Pore orientation mediated control of mechanical behavior of scaffolds and its application in cartilage-mimetic scaffold design. J. Mech. Behav. Biomed. Mater. 2015, 51, 169–183. [CrossRef] [PubMed]
50. Fu, Q.; Rahaman, M.N.; Bal, B.S.; Brown, R.F. Preparation and in vitro evaluation of bioactive glass (13–93) scaffolds with oriented microstructures for repair and regeneration of load-bearing bones. *J. Biomed. Mater. Res. Part A* **2010**, *93A*, 1380–1390. [CrossRef] [PubMed]

51. Jung, Y.; Park, M.S.; Lee, J.W.; Kim, Y.H.; Kim, S.-H.; Kim, S.H. Cartilage regeneration with highly-elastic three-dimensional scaffolds prepared from biodegradable poly(l-lactide-co-ε-caprolactone). *Biomaterials* **2008**, *29*, 4630–4636. [CrossRef]

52. Bhumiratana, S.; Grayson, W.L.; Castaneda, A.; Rockwood, D.N.; Gil, E.S.; Kaplan, D.L.; Vunjak-Novakovic, G. Nucleation and growth of mineralized bone matrix on silk-hydroxyapatite composite scaffolds. *Biomaterials* **2011**, *32*, 2812–2820. [CrossRef] [PubMed]

53. Wu, C.; Zhang, Y.; Zhu, Y.; Friis, T.; Xiao, Y. Structure–property relationships of silk-modified mesoporous bio glass scaffolds. *Biomaterials* **2010**, *31*, 3429–3438. [CrossRef]

54. Nauman, E.A.; Fong, K.E.; Keaveny, T.M. Dependence of Intertrabecular Permeability on Flow Direction and Anatomic Site. *Ann. Biomed. Eng.* **1999**, *27*, 517–524. [CrossRef]

55. Hollister, S.J. Porous scaffold design for tissue engineering. *Nat. Mater.* **2005**, *4*, 518–524. [CrossRef]

56. Slaughter, B.B.V.; Slaughter, B.B.V.; Khurshid, S.S.; Khurshid, S.S.; Fisher, O.Z.; Fisher, O.Z.; Khademhosseini, A.; Khademhosseini, A.; Peppas, N.A.; Peppas, N.A. Hydrogels in Regenerative Medicine. *Adv. Mater.* **2009**, *4*, 3307–3329. [CrossRef]

57. Chen, B.; Jones, R.R.; Mi, S.; Foster, J.; Alcock, S.G.; Hamley, I.W.; Connon, C.J. The mechanical properties of amniotic membrane influence its effect as a biomaterial for ocular surface repair. *Soft Matter* **2012**, *8*, 8379–8387. [CrossRef]

58. Wei, G.; Jin, Q.; Giannobile, W.V.; Ma, P.X. The enhancement of osteogenesis by nano-fibrous scaffolds incorporating rhBM-P-7 nanospheres. *Biomaterials* **2007**, *28*, 2087–2096. [CrossRef] [PubMed]

59. Freier, T.; Montenegro, R.; Koh, H.S.; Shoichet, M.S. Chitin-based tubes for tissue engineering in the nervous system. *Biomaterials* **2005**, *26*, 4624–4632. [CrossRef] [PubMed]

60. Webber, M.J.; Khan, O.F.; Sydlik, S.A.; Tang, B.C.; Langer, R. A Perspective on the Clinical Translation of Scaffolds for Tissue Engineering. *Ann. Biomed. Eng.* **2015**, *43*, 641–656. [CrossRef] [PubMed]

61. Steele, A.N.; Stapleton, L.M.; Farr, J.M.; Lucian, H.J.; Paulsen, M.J.; Eskandari, A.; Hironaka, C.E.; Thakore, A.D.; Wang, H.; Yu, A.C.; et al. A Biocompatible Therapeutic Catheter-Deliverable Hydrogel for In Situ Tissue Engineering. *Adv. Healthc. Mater.* **2019**, *8*, 1801147. [CrossRef]

62. Momeni, F.; Mehdi, M.; Hassani, N.S.; Liu, X.; Ni, J. A review of 4D printing. *Mater. Des.* **2017**, *122*, 42–79. [CrossRef]

63. Iqbal, N.; Khan, A.S.; Asif, A.; Yar, M.; Haycock, J.W.; Rehman, I.U. Recent concepts in biodegradable polymers for tissue engineering paradigms: A critical review. *Int. Mater. Rev.* **2019**, *64*, 91–126. [CrossRef]

64. Murdoch, M.H.; Badylok, S.F. Biomaterials-based In Situ Tissue Engineering. *Curr. Opin. Biomed. Eng.* **2017**, *1*, 4–7. [CrossRef]

65. Magnusson, J.P.; Saeed, A.O.; Fernández-Trillo, F.; Alexander, C. Synthetic polymers for biopharmaceutical delivery. *Polym. Chem.* **2011**, *2*, 48–59. [CrossRef]

66. Rezwan, K.; Chen, Q.Z.; Blaker, J.J.; Boccaccini, A.R. Biodegradable and bioactive porous polymer/inorganic composite scaffolds for bone tissue engineering. *Biomaterials* **2006**, *27*, 3431–3431. [CrossRef]

67. Talacua, H.; Smits, A.I.P.M.; Muylaert, D.E.P.; van Rijswijk, J.W.; Vink, A.; Verhaar, M.C.; Driessen-Mol, A.; van Herwerden, L.A.; Bouten, C.V.C.; Kluij, J.; et al. In Situ Tissue Engineering of Functional Small-Diameter Blood Vessels by Host Circulating Cells Only. *Tissue Eng. Part A* **2015**, *21*, 2583–2594. [CrossRef] [PubMed]

68. Guo, B.; Ma, P.X. Synthetic biodegradable functional polymers for tissue engineering: A brief review. *Sci. China Chem.* **2014**, *57*, 490–500. [CrossRef] [PubMed]

69. Pérez, R.A.; Won, J.E.; Knowles, J.C.; Kim, H.W. Naturally and synthetic smart composite biomaterials for tissue regeneration. *Adv. Drug Deliv. Rev.* **2013**, *65*, 471–496. [CrossRef] [PubMed]

70. Cameron, D.J.A.; Shaver, M.P. Aliphatic polypester polymer stars: Synthesis, properties and applications in biomedicine and nanotechnology. *Chem. Soc. Rev.* **2011**, *40*, 1761–1776. [CrossRef] [PubMed]

71. Hacker, M.C.; Krieghoff, J.; Mikos, A.G. *Synthetic Polymers*, 3rd ed.; Academic Press: Boston, MA, USA, 2019.

72. Hakkarainen, M.; Albertsson, A.-C. Degradation Products of Aliphatic and Aliphatic–Aromatic Polymers. In *Chromatography for Sustainable Polymeric Materials: Renewable, Degradable and Recyclable*; Albertsson, A.-C., Hakkarainen, M., Eds.; Springer: Heidelberg/Berlin, Germany, 2008; pp. 85–116.
73. Pan, Z.; Ding, J. Poly(lactide-co-glycolide) porous scaffolds for tissue engineering and regenerative medicine. Interface Focus 2012, 2, 366–377. [CrossRef] [PubMed]

74. Bi, X.; You, Z.; Gao, J.; Fan, X.; Wang, Y. A functional polyester carrying free hydroxyl groups promotes the mineralization of osteoblast and human mesenchymal stem cell extracellular matrix. Acta Biomater. 2014, 10, 2814–2823. [CrossRef]

75. Gómez, G.; Korkiakoski, S.; González, M.M.; Länssmark, S.; Ellä, V.; Salo, T.; Kellomäki, M.; Ashammakhi, N.; Arnaud, E. Effect of FGF and poly(lactide) scaffolds on calvarial bone healing with growth factor on biodegradable polymer scaffolds. J. Craniofacial Surg. 2006, 17, 935–942. [CrossRef]

76. Mabileau, G.; Aguado, E.; Stancu, I.C.; Cincu, C.; Baslé, M.F.; Chappard, D. Effects of FGF-2 release from a hydrogel polymer on bone mass and microarchitecture. Biomaterials 2008, 29, 1593–1600. [CrossRef]

77. Fujimoto, K.L.; Guan, J.; Oshima, H.; Sakai, T.; Wagner, W.R. In Vivo Evaluation of a Porous, Elastic, Biodegradable Patch for Reconstructive Cardiac Procedures. Ann. Thorac. Surg. 2007, 83, 648–654. [CrossRef]

78. Erggelet, C.; Endres, M.; Neumann, K.; Morawietz, L.; Haberstroh, K.; Sittinger, M.; Kaps, C. Formation of cartilage repair tissue in articular cartilage defects pretreated with microfracture and covered with cell-free polymer-based implants. J. Orthop. Res. 2009, 27, 1353–1360. [CrossRef]

79. Yoshinobu, O.; Patrick, W.S. Acute performance of a novel restorative transcatheter aortic valve: Preclinical results. EuroIntervention 2017, 13, 1410–1417.

80. Herberg, S.; Siedler, M.; Pippig, S.; Schuetz, A.; Kim, C.K.; Wikesjö, U.M.E. Development of an injectable composite as a carrier for growth factor-enhanced periodontal regeneration. J. Clin. Periodontol. 2008, 35, 976–984. [CrossRef] [PubMed]

81. Lee, K.Y.; Jeong, L.; Kang, Y.O.; Lee, S.J.; Park, W.H. Electrospinning of polysaccharides for regenerative medicine. Adv. Drug Deliv. Rev. 2009, 61, 1020–1032. [CrossRef] [PubMed]

82. Khor, E.; Lim, L.Y. Implantable applications of chitin and chitosan. Biomaterials 2003, 24, 2339–2349. [CrossRef]

83. Croisier, F.; Jérôme, C. Chitosan-based biomaterials for tissue engineering. Eur. Polym. J. 2013, 49, 780–792. [CrossRef]

84. Burdick, J.A.; Prestwich, G.D. Hyaluronic Acid Hydrogels for Biomedical Applications. Adv. Mater. 2011, 23, 41–56. [CrossRef] [PubMed]

85. Kim, I.L.; Khetan, S.; Baker, B.M.; Chen, C.S.; Burdick, J.A. Fibrous hyaluronic acid hydrogels that direct MSC chondrogenesis through mechanical and adhesive cues. Biomaterials 2013, 34, 5571–5580. [CrossRef]

86. Elia, R.; Fueggy, P.W.; VanDelden, A.; Firpo, M.A.; Prestwich, G.D.; Peatle, R.A. Stimulation of in vivo angiogenesis by in situ crosslinked, dual growth factor-loaded, glycosaminoglycan hydrogels. Biomaterials 2010, 31, 4630–4638. [CrossRef]

87. Bhakta, G.; Rai, B.; Lim, Z.X.H.; Hui, J.H.; Stein, G.S.; van Wijnen, A.J.; Nurcombe, V.; Prestwich, G.D.; Cool, S.M. Hyaluronic acid-based hydrogels functionalized with heparin that support controlled release of bioactive BMP-2. Biomaterials 2012, 33, 6113–6122. [CrossRef]

88. Dinjaski, N.; Kaplan, D.L. Recombinant protein blends: Silk beyond natural design. Curr. Opin. Biotechnol. 2016, 39, 1–7. [CrossRef] [PubMed]

89. Sheikh, Z.; Najeeb, S.; Khurshid, Z.; Verma, V.; Rashid, H.; Glogauer, M. Biodegradable Materials for Bone Repair and Tissue Engineering Applications. Materials 2015, 8, 5744–5794. [CrossRef] [PubMed]

90. Cen, L.; Liu, W.; Cui, L.; Zhang, W.; Cao, Y. Collagen Tissue Engineering: Development of Novel Biomaterials and Applications. Pediatric Res. 2008, 63, 492. [CrossRef] [PubMed]

91. Shi, C.; Li, Q.; Zhao, Y.; Chen, W.; Chen, B.; Xiao, Z.; Lin, H.; Nie, L.; Wang, D.; Dai, J. Stem-cell-capturing collagen scaffold promotes cardiac tissue regeneration. Biomaterials 2011, 32, 2508–2515. [CrossRef] [PubMed]

92. Wu, J.; Cao, L.; Liu, Y.; Zheng, A.; Jiao, D.; Zeng, D.; Wang, X.; Kaplan, D.L.; Jiang, X. Functionalization of Silk Fibroin Electrospun Scaffolds via BMSC Affinity Peptide Grafting through Oxidative Self-Polymerization of Dopamine for Bone Regeneration. ACS Appl. Mater. Interfaces 2019, 11, 8878–8895. [CrossRef] [PubMed]

93. Haugen, H.J.; Lyngstadaas, S.P.; Rossi, F.; Perale, G. Bone grafts: Which is the ideal biomaterial? J. Clin. Periodontal. 2019, 46, 92–102. [CrossRef]
95. Wu, J.; Zheng, K.; Huang, X.; Liu, J.; Liu, H.; Boccaccini, A.R.; Wan, Y.; Guo, X.; Shao, Z. Thermally triggered injectable chitosan/silk fibroin/bioactive glass nanoparticle hydrogels for in-situ bone formation in rat calvarial bone defects. *Acta Biomater.* 2019, 91, 60–71. [CrossRef]

96. Woodruff, M.A.; Rath, S.N.; Susanto, E.; Haupt, L.M.; Hutmacher, D.W.; Nurcombe, V.; Cool, S.M. Sustained release and osteogenic potential of heparan sulfate-doped fibrin glue scaffolds within a rat cranial model. *J. Mol. Histol.* 2007, 38, 425–433. [CrossRef]

97. Kodama, N.; Nagata, M.; Tabata, Y.; Ozeki, M.; Ninomiya, T.; Takagi, R. A local bone anabolic effect of rhFGF2-impregnated gelatin hydrogel by promoting cell proliferation and coordinating osteoblastic differentiation. *Bone* 2009, 44, 699–707. [CrossRef]

98. Feng, Q.; Wei, K.; Lin, S.; Xu, Z.; Sun, Y.; Shi, P.; Li, G.; Bian, L. Mechanically resilient, injectable, and bioadhesive supramolecular gelatin hydrogels crosslinked by weak host-guest interactions assist cell infiltration and in situ tissue regeneration. *Biomaterials* 2016, 101, 217–228. [CrossRef]

99. Lee, S.J.; Wang, H.J.; Kim, T.H.; Choi, J.S.; Kulkarni, G.; Jackson, J.D.; Atala, A.; Yoo, J.J. In Situ Tissue Regeneration of Renal Tissue Induced by Collagen Hydrogel Injection. *Stem Cells Transl. Med.* 2018, 7, 241–250. [CrossRef] [PubMed]

100. Landa, N.; Miller, L.; Feinberg, M.S.; Holbova, R.; Shachar, M.; Freeman, I.; Cohen, S.; Leor, J. Effect of injectable alginate implant on cardiac remodeling and function after recent and old infarcts in rats. *Circulation* 2008, 117, 1388–1396. [CrossRef] [PubMed]

101. Kubo, M.; Imai, S.; Fujiyama, M.; Isoya, E.; Ando, K.; Mimura, T.; Matsusue, Y. Exogenous collagen-enhanced recruitment of mesenchymal stem cells during rabbit articular cartilage repair. *Acta Orthop.* 2007, 78, 845–855. [CrossRef] [PubMed]

102. Dai, Y.; Liu, G.; Ma, L.; Wang, D.; Gao, C. Cell-free macro-porous fibrin scaffolds for in situ inductive regeneration of full-thickness cartilage defects. *J. Mater. Chem. B* 2016, 4, 4410–4419. [CrossRef]

103. Ma, F.; Ge, Y.; Liu, N.; Pang, X.; Shen, X.; Tang, B. In situ fabrication of a composite hydrogel with tunable mechanical properties for cartilage tissue engineering. *J. Mater. Chem. B* 2019, 7, 2463–2473. [CrossRef]

104. Kin, S.; Hagiwara, A.; Nakase, Y.; Kuriu, Y.; Nakashima, S.; Yoshikawa, T.; Sakakura, C.; Otsui, E.; Nakamura, T.; Yamagishi, H. Regeneration of skeletal muscle using in situ tissue engineering on an acellular collagen sponge scaffold in a rabbit model. *Asaio J.* 2007, 53, 506–513. [CrossRef]

105. Brouwer, K.M.; Daamen, W.F.; van Lochem, N.; Reijnen, D.; Wijnen, R.M.H.; van Kuppevelt, T.H. Construction and in vivo evaluation of a dual layered collagenous scaffold with a radial pore structure for repair of the diaphragm. *Acta Biomater.* 2013, 9, 6844–6851. [CrossRef]

106. Nakahara, T.; Nakamura, T.; Kobayashi, E.; Inoue, M.; Shigeno, K.; Tabata, Y.; Eto, K.; Shimizu, Y. Novel approach to regeneration of periodontal tissues based on in situ tissue engineering: Effects of controlled release of basic fibroblast growth factor from a sandwich membrane. *Tissue Eng.* 2003, 9, 153–162. [CrossRef]

107. Boucard, N.; Viton, C.; Agay, D.; Mari, E.; Roger, T.; Chancerelle, Y.; Domard, A. The use of physical hydrogels of chitosan for skin regeneration following third-degree burns. *Biomaterials* 2007, 28, 3478–3488. [CrossRef]

108. Wang, S.Y.; Kim, H.; Kwak, G.; Yoon, H.Y.; Jo, S.D.; Lee, J.E.; Cho, D.; Kwon, I.C.; Kim, S.H. Development of Biocompatible HA Hydrogels Embedded with a New Synthetic Peptide Promoting Cellular Migration for Advanced Wound Care Management. *Adv. Sci.* 2018, 5, 1800852. [CrossRef]

109. Chung, Y.I.; Ahn, K.M.; Jeon, S.H.; Lee, S.Y.; Lee, J.H.; Tae, G. Enhanced bone regeneration with BMP-2 loaded functional nanoparticles-hydrogel complex. *J. Control. Release* 2007, 121, 91–99. [CrossRef] [PubMed]

110. Wu, G.; Feng, C.; Quan, J.; Wang, Z.; Wei, W.; Zang, S.; Kang, S.; Hui, G.; Chen, X.; Wang, Q. In situ controlled release of streptomycin-cell derived factor-1α and antimîR-138 for on-demand cranial bone regeneration. *Carbohydr. Polym.* 2018, 182, 215–224. [CrossRef] [PubMed]

111. Lee, J.S.; Jin, Y.; Park, H.-J.; Yang, K.; Lee, M.S.; Yang, H.S.; Cho, S.-W. In Situ Bone Tissue Engineering With an Endogenous Stem Cell Mobilizer and Osteoinductive Nanofibrous Polymeric Scaffolds. *Biotechnol. J.* 2017, 12, 1700602. [CrossRef] [PubMed]

112. Di Bella, C.; Duchi, S.; O’Connell, C.D.; Blanchard, R.; Augustine, C.; Yue, Z.; Thompson, F.; Richards, C.; Beirne, S.; Onorillo, C.; et al. In situ handheld three-dimensional bioprinting for cartilage regeneration. *J. Tissue Eng. Regen. Med.* 2018, 12, 611–621. [CrossRef] [PubMed]

113. Perdisa, F.; Fialdari, G.; Sessa, A.; Busacca, M.; Zaffagnini, S.; Maracci, M.; Kon, E. One-Step Treatment for Patellar Cartilage Defects With a Cell-Free Osteochondral Scaffold: A Prospective Clinical and MRI Evaluation. *Am. J. Sports Med.* 2017, 45, 1581–1588. [CrossRef]
114. Dai, Y.; Gao, Z.; Ma, L.; Wang, D.; Gao, C. Cell-Free HA-MA/PLGA Scaffolds with Radially Oriented Pores for In Situ Inductive Regeneration of Full Thickness Cartilage Defects. *Macromol. Biosci.* 2016, 16, 1632–1642. [CrossRef]

115. Capulli, A.K.; Emmert, M.Y.; Pasqualini, F.S.; Kehl, D.; Caliskan, E.; Lind, J.U.; Sheehy, S.P.; Park, S.J.; Ahn, S.; Weber, B.; et al. JetValve: Rapid manufacturing of biohybrid scaffolds for biomimetic heart valve replacement. *Biomaterials* 2017, 133, 229–241. [CrossRef]

116. Hori, Y.; Nakamura, T.; Matsumoto, K.; Kurokawa, Y.; Satomi, S.; Shimizu, Y. Experimental study on In Situ tissue engineering of the stomach by an acellular collagen sponge scaffold graft. *Asaio J.* 2001, 47, 206–210. [CrossRef]

117. Abbushi, A.; Endres, M.; Cabrera, M.; Kroppenstedt, S.N.; Thomale, U.; Sittig, M.; Hegewald, A.A.; Morawietz, L.; Lemke, A.J.; Bansemer, V.G.t.; et al. Regeneration of intervertebral disc tissue by resorbable cell-free polyglycolic acid-based implants in a rabbit model of disc degeneration. *Spine* 2008, 33, 1527–1532. [CrossRef]

118. Badylak, S.F.; Freytes, D.O.; Gilbert, T.W. Reprint of: Extracellular matrix as a biological scaffold material: Structure and function. *Acta Biomater.* 2015, 23, 17–26. [CrossRef] [PubMed]

119. Taylor, D.A.; Sampaio, L.C.; Ferdous, Z.; Gobin, A.S.; Taiie, R.J. Decellularized matrices in regenerative medicine. *Acta Biomater.* 2018, 74, 74–89. [CrossRef] [PubMed]

120. Agmon, G.; Christman, K.L. Controlling stem cell behavior with decellularized extracellular matrix scaffolds. *Curr. Opin. Solid State Mater. Sci.* 2016, 20, 193–201. [CrossRef] [PubMed]

121. Hussey, G.S.; Cramer, M.C.; Badylak, S.F. Extracellular Matrix BioScaffolds for Building Gastrointestinal Tissue. *Cell. Mol. Gastroenterol. Hepatol.* 2018, 5, 1–13. [CrossRef] [PubMed]

122. Dzik, J.; Badylak, S.; Yabroudi, M.; Scart, B.; Ambrosio, F.; Stearns, K.; Turner, N.; Wyse, A.; Boninger, M.L.; Brown, E.H.P.; et al. An acellular biologic scaffold treatment for volumetric muscle loss: Results of a 13-patient cohort study. *Npj Regen. Med.* 2016, 1, 16008. [CrossRef] [PubMed]

123. Cai, R.; Nakamoto, T.; Kawazoe, N.; Chen, G. Influence of stepwise chondrogenesis-mimicking 3D extracellular matrix on chondrogenic differentiation of mesenchymal stem cells. *Biomaterials* 2015, 52, 199–207. [CrossRef] [PubMed]

124. Benders, K.E.M.; Weeren, P.R.V.; Badylak, S.F.; Saris, D.B.F.; Dhert, W.J.A.; Malda, J. Extracellular matrix scaffolds for cartilage and bone regeneration. *Trends Biotechnol.* 2013, 31, 169–176. [CrossRef]

125. Lié, E.; Park, K.W.; Chun, S.Y.; Kim, H.; Kwon, T.G.; Joug, Y.K.; Han, D.K. Biomimetic Porous PLGA Scaffolds Incorporating Decellularized Extracellular Matrix for Kidney Tissue Regeneration. *ACS Appl. Mater. Interfaces* 2016, 8, 21145–21154. [CrossRef]

126. Woo, J.S.; Fishbein, M.C.; Reemtsen, B. Histologic examination of decellularized porcine intestinal submucosal extracellular matrix (CorMatrix) in pediatric congenital heart surgery. *Cardiovasc. Pathol.* 2016, 25, 12–17. [CrossRef]

127. Dai, J.; Qiao, W.; Shi, J.; Liu, C.; Hu, X.; Dong, N. Modifying decellularized aortic valve scaffolds with stromal cell-derived factor-1α loaded proteolytically degradable hydrogel for recellularization and remodeling. *Acta Biomater.* 2019, 88, 280–292. [CrossRef]

128. Levato, R.; Planell, J.A.; Mateos-Timoneda, M.A.; Engel, E. Role of ECM/peptide coatings on SDF-1α triggered mesenchymal stromal cell migration from microcarriers for cell therapy. *Acta Biomater.* 2015, 18, 59–67. [CrossRef]

129. Muylaert, D.E.P.; van Almen, G.C.; Talacu, H.; Fledderus, J.O.; Kluin, J.; Hendriks, S.S.; van Dongen, J.L.J.; Sijbesma, E.; Bosman, A.W.; Mes, T.; et al. Early in-situ cellularization of a supramolecular vascular graft is modified by synthetic cell-derived factor-1α derived peptides. *Biomaterials* 2016, 76, 187–195. [CrossRef] [PubMed]

130. Keane, T.J.; DeWard, A.; Londono, R.; Saldin, L.T.; Castleton, A.A.; Carey, L.; Nieponice, A.; Lagasse, E.; Badylak, S.F. Tissue-Specific Effects of Esophageal Extracellular Matrix. *Tissue Eng. Part. A* 2015, 21, 2293–2300. [CrossRef] [PubMed]

131. Katerinaki, E.; Zanetto, U.; Sterne, G.D. Histological appearance of Stratticetm tissue matrix used in breast reconstruction. *J. Plast. Reconstr. Aesthetic Surg.* 2010, 63, 840–841. [CrossRef] [PubMed]

132. Connor, J.; McQuillan, D.; Sandor, M.; Wan, H.; Lombardi, J.; Bachrach, N.; Harper, J.; Xu, H. Retention of structural and biochemical integrity in a biological mesh supports tissue remodeling in a primate abdominal wall model. *Regen. Med.* 2009, 4, 185–195. [CrossRef] [PubMed]
133. Tan, Q.-W.; Zhang, Y.; Luo, J.-C.; Zhang, D.; Xiong, B.-J.; Yang, J.-Q.; Xie, H.-Q.; Lv, Q. Hydrogel derived from decellularized porcine adipose tissue as a promising biomaterial for soft tissue augmentation. *J. Biomed. Mater. Res. Part. A* 2017, 105, 1756–1764. [CrossRef]

134. Kitamura, M.; Hirano, S.; Kanemaru, S.-i.; Kitani, Y.; Ohno, S.; Kojima, T.; Nakamura, T.; Ito, J.; Rosen, C.A.; Gilbert, T.W. Glottic regeneration with a tissue-engineering technique, using acellular extracellular matrix scaffold in a canine model. *J. Tissue Eng. Regen. Med.* 2016, 10, 825–832. [CrossRef]

135. Zhou, J.; Ding, J.; Zhu, Z.; Xu, J.; Yi, Y.; Li, Y.; Fan, H.; Bai, S.; Yang, J.; Tang, Y.; et al. Surface biofunctionalization of the decellularized porcine aortic valve with VEGF-loaded nanoparticles for accelerating endothelialization. *Mater. Sci. Eng. A* 2019, 79, 632–643. [CrossRef]

136. Cao, W.; Hench, L.L. Bioactive Materials. *Ceram. Int.* 1996, 22, 493–507. [CrossRef]

137. Dorozhkin, S.V. Amorphous calcium (orthophosphates. *Acta Biomater.* 2010, 6, 4457–4475. [CrossRef]

138. Okamoto, M.; John, B. Synthetic biopolymer nanocomposites for tissue engineering scaffolds. *Prog. Polym. Sci.* 2013, 38, 548–581. [CrossRef]

139. Nyberg, E.; Holmes, C.; Witham, T.; Grayson, W.L. Growth factor-eluting technologies for bone tissue engineering. *Drug Deliv. Transl. Res.* 2016, 6, 184–194. [CrossRef] [PubMed]

140. Samavedi, S.; Whittington, A.R.; Goldstein, A.S. Calcium phosphate ceramics in bone tissue engineering: A review of properties and their influence on cell behavior. *Acta Biomater.* 2013, 9, 8037–8045. [CrossRef] [PubMed]

141. Changchun, Z.; Xiangfeng, L.; Junqiu, C.; Hongsong, F.; Xingdong, Z. Bioactive Ceramics and Metals for Regenerative Engineering. In *Regenerative Engineering*; CRC Press: Boca Raton, FL, USA, 2018.

142. Tanner, K.E. 3-Hard tissue applications of biocomposites. In *FishAbility: Inherent Biocomposites in Fish and Fisheries*; Ambrosio, L., Ed.; Woodhead Publishing: Swaston, Cambridge, UK, 2017.

143. Habraken, W.; Habibovic, P.; Eppe, M.; Böhmer, M. Calcium phosphates in biomedical applications: Materials for the future? *Mater. Today*, 2016, 19, 69–87. [CrossRef]

144. Leite, A.J.; Sarker, B.; Zehnder, T.; Silva, R.; Mano, J.F.; Boccaccini, A.R. Bioplotting of a bioactive alginate dialdehyde-gelatin composite hydrogel containing bioactive glass nanoparticles. *Biofabrication* 2016, 8, 035005. [CrossRef] [PubMed]

145. Meka, S.R.K.; Jain, S.; Chatterjee, K. Strontium eluting nanofibers augment stem cell osteogenesis for bone tissue regeneration. *Colloids Surf. B Biointerfaces* 2016, 146, 649–656. [CrossRef] [PubMed]

146. Fu, Q.; Saiz, E.; Rahaman, M.N.; Tomsia, A.P. Bioactive glass scaffolds for bone tissue engineering: State of the art and future perspectives. *Mater. Sci. Eng. C* 2011, 31, 1245–1256. [CrossRef]

147. Song, G.; Habibovic, P.; Bao, C.; Hu, J.; van Blitterswijk, C.A.; Yuan, H.; Chen, W.; Xu, H.H.K. The homing of bone marrow MSCs to non-osseous sites for ectopic bone formation induced by osteoinductive calcium phosphate. *Biomaterials* 2013, 34, 2167–2176. [CrossRef]

148. Wang, J.; Chen, Y.; Zhu, X.; Yuan, T.; Tan, Y.; Fan, Y.; Zhang, X. Effect of phase composition on protein adsorption and osteoinduction of porous calcium phosphate ceramics in mice. *J. Biomed. Mater. Res. Part. A* 2014, 102, 4234–4243. [CrossRef]

149. Li, X.; Song, T.; Chen, X.; Wang, M.; Yang, X.; Xiao, Y.; Zhang, X. Osteoinductivity of Porous Biphasic Calcium Phosphate Ceramic Spheres with Nanocrystalline and Their Efficacy in Guiding Bone Regeneration. *ACS Appl. Mater. Interfaces* 2019, 11, 3722–3736. [CrossRef]

150. Amini, A.R.; Laurencin, C.T.; Nakavarapu, S.P. Bone tissue engineering: Recent advances and challenges. *Crit. Rev. Biomed. Eng.* 2012, 40, 363–408. [CrossRef] [PubMed]

151. Barrière, F.; van Blitterswijk, C.A.; de Groot, K. Bone regeneration: Molecular and cellular interactions with calcium phosphate ceramics. *Int. J. Nanomed.* 2006, 1, 317–332.
155. Trbakovic, A.; Hedenqvist, P.; Mellgren, T.; Ley, C.; Hilborn, J.; Ossipov, D.; Ekman, S.; Johansson, C.B.; Jensen-Waern, M.; Thor, A. A new synthetic granular calcium phosphate compound induces new bone in a sinus lift rabbit model. J. Dent. 2018, 70, 31–39. [CrossRef] [PubMed]

156. Schlickewei, C.; Klatte, T.O.; Wildermuth, Y.; Laaff, G.; Rüeger, J.M.; Ruesing, J.; Chernousova, S.; Lehmann, W.; Epple, M. A bioactive nano-calcium phosphate paste for in-situ transfection of BMP-7 and VEGF-A in a rabbit critical-size bone defect: Results of an in vivo study. J. Mater. Sci. Mater. Med. 2019, 30, 15. [CrossRef]

157. Oliveira, H.; Catros, S.; Boiziau, C.; Siadous, R.; Marti-Munoz, J.; Bareille, R.; Rey, S.; Castano, O.; Planell, J.; Amédée, J.; et al. The proangiogenic potential of a novel calcium releasing biomaterial: Impact on cell recruitment. Acta Biomater. 2016, 29, 435–445. [CrossRef]

158. Shao, H.; Sun, M.; Zhang, F.; Liu, A.; He, Y.; Fu, J.; Yang, X.; Wang, H.; Gou, Z. Custom Repair of Mandibular Bone Defects with 3D Printed Bioceramic Scaffolds. J. Dent. Res. 2018, 97, 68–76. [CrossRef]

159. Fu, J.; Zhuang, C.; Qiu, J.; Ke, X.; Yang, X.; Jin, Z.; Zhang, L.; Yang, G.; Xie, L.; Xu, S.; et al. Core–Shell Biphasic Microspheres with Tunable Density of Shell Micropores Providing Tailorable Bone Regeneration. Tissue Eng. Part. A 2018, 25, 588–602. [CrossRef]

160. Ma, H.; Feng, C.; Chang, J.; Wu, C. 3D-printed bioceramic scaffolds: From bone tissue engineering to tumor therapy. Acta Biomater. 2018, 79, 37–59. [CrossRef] [PubMed]

161. Carrabba, M.; Maddeddu, P. Current Strategies for the Manufacture of Small Size Tissue Engineering Vascular Grafts. Front. Bioeng. Biotechnol. 2018, 6, 1–12. [CrossRef] [PubMed]