Research review paper

Towards the design of 3D multiscale instructive tissue engineering constructs: Current approaches and trends

Sara M. Oliveira, Rui L. Reis, João F. Mano *

3B's Research Group — Biomaterials, Biodegradables and Biomimetics, Dept. of Polymer Engineering, University of Minho, Avepark - Parque de Ciência e Tecnologia, Zona Industrial da Gandra, 4805-017 Barco-Guimarães, Portugal

ICVS/3B's — PT Government Associate Laboratory, Braga/Guimarães 4805-017 Barco-Guimarães, Portugal

A B S T R A C T

The design of 3D constructs with adequate properties to instruct and guide cells both in vitro and in vivo is one of the major focuses of tissue engineering. Successful tissue regeneration depends on the favorable crosstalk between the supporting structure, the cells and the host tissue so that a balanced matrix production and degradation are achieved. Herein, the major occurring events and players in normal and regenerative tissue are overviewed. These have been inspiring the selection or synthesis of instructive cues to include into the 3D constructs. We further highlight the importance of a multiscale perception of the range of features that can be included on the biomimetic structures. Lastly, we focus on the current and developing tissue-engineering approaches for the preparation of such 3D constructs: top-down, bottom-up and integrative. Bottom-up and integrative approaches present a higher potential for the design of tissue engineering devices with multiscale features and higher biochemical control than top-down strategies, and are the main focus of this review.

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* Corresponding author at: ICVS/3B's — PT Government Associate Laboratory, Braga/Guimarães 4805-017 Barco-Guimarães, Portugal.
E-mail address: jmano@dep.uminho.pt (J.F. Mano).

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1. Introduction

Tissue engineering aims to restore the loss of tissue and organ’s functionality resulting from injury, aging or disease (Lanza et al., 2011). Bio-materials, cells and bioactive factors are commonly considered the key elements needed for the preparation of 3D tissue engineered constructs for the regeneration of those damaged tissues (Hench and Polak, 2002; Langer and Vacanti, 1993). Those act primarily as supportive and informative platforms that guide cell behavior. The physicochemical properties of those devices affect cell adhesion, proliferation, differentiation and matrix synthesis. Upon implantation, their properties also dictate host tissue response: inflammatory and immune responses (Franz et al., 2011). Moreover, the cellular and matrix compositions of the injured tissue will influence the cell responses in the construct. The further crosstalk between the construct and the host tissue will thus define the provisional micro/nano-environment, dictating cell behavior and consequently tissue fate: failed healing, repair or regeneration — Fig. 1 (Sephel and Woodward, 2001).

This repairing/regenerative micro/nano-environment is regulated by several cell types, matrix proteins, growth factors and cytokines. Only an adequate balance between new matrix deposition and matrix degradation allows the achievement of a successfully regenerated tissue that is identical to the original one.

Tissue engineering has focused on the design of 3D devices to promote the regeneration of several types of tissues, e.g.: skin (Grober et al., 2011), cartilage (Mano and Reis, 2007), bone (Healy and Guldberg, 2007), tendon (Hampson et al., 2008), and cardiac tissue (Chiou and Radisic, 2013).

The ideal features of the 3D structures have been evolving with increasing understanding of cell–material interactions both in vitro and in vivo (Custudio et al., 2014; Hench and Polak, 2002). The intention of the current paradigm of the construct is to provide instructive cues for cellular activation and guidance. Besides demanding an adequate cell instruction regarding migration, proliferation and differentiation, the ideal construct also requires other generic characteristics, such as suitable nutrients and metabolite diffusion, regulated degradation profile and patient-customization. The design of 3D substrates fulfilling all requirements is still very challenging (Mano, 2015; Oliveira and Mano, 2014). It demands the management and understanding of multiple variables that are distributed along all length-scales and affect cell behavior in vitro and in vivo. Processing techniques are still evolving and promise increasing control on the multiscale and spatial–temporal features of the constructs. Herein, we begin summarizing the major occurring events and features in the normal and regenerative cell environment (niche) as design inspirations for instructive biomaterials. We highlight the importance of a multiscale understanding of the range of properties that can be included on the substrates. Lastly, we overview the current and expected tissue engineering approaches for the preparation of 3D constructs: top-down, bottom-up and integrative approaches.

2. Tissue/cell niches as regenerative inspiration for biomaterials design

The stem cell niches are dynamic and complex structures where diverse biochemical, physical, inflammatory, and cellular-derived cues bi-directionally and reciprocally interact with several local cells and stem cells — Table 1. The niches are responsible for modulating stem cell behavior, which is crucial for the maintenance of tissue homeostasis (Jones and Wagers, 2008; Lane et al., 2014; Li and Xie, 2005; Moore and Lemischka, 2006; Schofield, 1977). The ability of the stem cells to self-renew and differentiate is orchestrated by the spatiotemporal presentation of niche’s cues and the dialogs occurring with several cell types. Both niches and tissues are multiscale and complex systems where smaller units interact originating larger cellular structures. The secreted factors, the physical cues, the extracellular matrix and other cells interact with the cell surface receptors at the nanoscale creating meso-, micro- and macro-complex structures that are hierarchically and spatially organized (Lane et al., 2014; Nakatsuji, 2013). Among the multitude of surface receptors, cells present super-families of integrins, cadherins, receptor tyrosine kinase, selectins, proteoglycans and immunoglobulins. The type, density and stability of the consequently activated/co-activated receptors trigger specific intracellular events defining cell fate at very different extents, such as: survival, motility, polarity, proliferation, cytoskeleton organization, cell–cell interactions and gene-expression (Bernfield et al., 1992; Borghi et al., 2010; Carey, 1997; Choi et al., 2011; Gumbiner, 1996).
The nature of the niche is considered specific for each stem cell type and tissue (Gattazzo et al., 2014). Nevertheless, the categories of the environmental cues influencing cell behavior are transversal for the tissue, cell or culture type (in vivo, ex vivo or in vitro) — Table 1. Therefore, understanding the normal and regenerative (stem) cell niche and tissue can enlighten and inspire the design of cell instructive biomaterials.

Upon tissue injury, immune cells and others migrate, triggering the healing cascade or an immune response. The healing cascade consists of four major integrated and overlapping phases: hemostasis; inflammation; proliferation and repair; and tissue remodeling or resolution — Fig. 2 (Enoch and Leaper, 2008; Gosain and DiPietro, 2004; Guo and DiPietro, 2010).

The temporary micro/nano-environment of each stage is regulated by several cell types, matrix proteins, growth factors (GFs) and cytokines — Table 2. Extreme changes are known to happen in the physical, biochemical, cell–cell communication and extracellular matrix polysaccharides or protein patterns, during the normal and efficient healing cascade (Witte and Barbul, 1997). The failure, or prolongation, in one phase might result in a delayed or impaired healing. This may be caused by low stem cell availability, aging, continued activation of inflammatory cells, excessive pro-inflammatory MMP, excessive proliferation and matrix synthesis, among others (Forbes and Rosenthal, 2014; Mehta et al., 2012; Menke et al., 2007; Utz et al., 2010; Witte and Barbul, 1997).

The repair process may only restore some of the structures of the original tissue and involve scar formation. Only an adequate balance between new matrix deposition and matrix degradation, during the healing, allows a successful regenerated tissue identical to the original one to be achieved.

An injury causes vascular endothelium disruption, exposing the collagen layer and other elements that activate platelets. Nowadays, platelets are recognized as major players in the healing cascade, tissue repair and regeneration (Anitua et al., 2004; Gawaz and Vogel, 2013; Stellos et al., 2010). Besides maintaining the blood vessel integrity by adhering, aggregating and forming a pro-coagulant fibrin surface (hemostasis), platelets initiate the healing process. Once activated, the substances released from the platelet’s granules (dense granules, lysosomes and α-granules) are responsible for triggering the healing cascade. For instance, α-granules contain many instructive GFs, such as: vascular endothelial growth factor (VEGF), fibroblast growth factor (FGF), transforming growth factor β (TGFβ), epidermal growth factor (EGF), platelet derived growth factor (PDGF) and insulin-like growth factor (IGF). Besides their role in hemostasis, platelets play important anti and pro-inflammatory roles and their releasates also participate in the other healing stages (Gawaz and Vogel, 2013; Semple et al., 2011; Sephel and Woodward, 2001; Vieira-de-Abreu et al., 2012; Weyrich and Zimmerman, 2004; Zarbock et al., 2007). In the early phase, adherent platelets interact with endothelial cells, monocytes and neutrophils. They activate neutrophils and endothelial cells inducing the production of inflammatory cytokines. Platelets are potent instructive cells, and actually have been inspiring and used as a source of multiple instructive

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Table 1
Categories of physicochemical cues in stem cell niches that affect cell behavior, and some examples of extracellular cue–cell receptor interactions (Humphries et al., 2006; Lane et al., 2014). ICAM, intercellular adhesion molecule; VCAM, vascular cell adhesion molecule.

<table>
<thead>
<tr>
<th>Category</th>
<th>Ligands/cue</th>
<th>Cell receptors/effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Secreted factors</td>
<td>Chemokine/cue</td>
<td>Chemokine receptors</td>
</tr>
<tr>
<td>Extracellular matrix</td>
<td>Fibronectin</td>
<td>Integrins α5β1, α5β3, α5β6, α6β3, αβλβ3, αβγ1, αβδ1, αβε1</td>
</tr>
<tr>
<td>Physical</td>
<td>Topography</td>
<td>Alters formation of focal adhesion points and cytoskeleton contraction</td>
</tr>
<tr>
<td>Hypoxia and metabolic</td>
<td>O2</td>
<td>Enters cells by diffusion and participates in metabolic process.</td>
</tr>
<tr>
<td>Cellar</td>
<td>Tissue specific-cells, stem cells, immune cells, nerve cells, endothelial cell, stromal cell</td>
<td>Cadherins (N, N2, P, E), ICAM, VCAM, integrins (e.g. αv)3 with CD31 in endothelial cell</td>
</tr>
</tbody>
</table>

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**Fig. 2.** a) General phases involved in the healing cascade triggered by a tissue injury (based on Epstein et al., 1999; Mendonca and Coutinho-Netto, 2009; Guo and DiPietro, 2010; Claes et al., 2012). b) Illustration of a cutaneous wound reproduced with permission from Clark et al. (2007): top — crosstalk between the clot and the surrounding tissue in the inflammatory phase usually ending with tissue repair or failed healing. Several bioactive factors have multiple effects and multiple sources. FGF, fibroblast growth factor; IGF, insulin-like growth factor; TGF, transforming growth factor; PDGF, platelet-derived growth factor; VEGF, vascular endothelial growth factor; MMPs, matrix metalloproteinases; t-PA, tissue plasminogen activator; u-PA, urokinase-type plasminogen activator.
proteins for the preparation of new constructs for several tissue-engineering applications (Lima et al., 2015; Oliveira et al., 2015a,b; Santo et al., 2012). Similar to platelets, other cells involved in the healing cascade play important roles by secreting similar or other instructive compounds such as the leukocytes and other granulocytes — Table 2.

The specific nature of the extracellular matrix of the tissue also regulates the GFs and other bioactive protein distribution, type, and stability (Discher et al., 2009; Kim et al., 2011; Schultz and Wysocki, 2009). GFs are presented either in soluble form, or mainly electrostatically bound to the negatively charged glycosaminoglycans (GAGs). GAGs present various molecular arrangements and different sulfation degrees.

Table 2
Major cell types involved in the healing of injured tissues and their respective function and participating phases.
Adapted from Kamath et al. (2001), Sephel and Woodward (2001), Sipe et al. (2004) and Vieira-de-Abreu et al. (2012).

<table>
<thead>
<tr>
<th>Cell Type</th>
<th>Phase</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platelets</td>
<td>Hemostasis</td>
<td>Major players in hemostasis. Granules release cytokines, chemokines,</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>GFS, clotting agents, proteases and inflammation mediators. Facilitate adhesion,</td>
</tr>
<tr>
<td></td>
<td>Repair</td>
<td>coagulation, vasoconstriction, repair and clot resorption. Their activation attracts and activates several cells</td>
</tr>
<tr>
<td></td>
<td>Remodeling</td>
<td>Granulocytes (neutrophils, basophils, eosinophils) and agranulocytes</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>(lymphocytes monocytes and macrophages) are immune system cells</td>
</tr>
<tr>
<td></td>
<td></td>
<td>involved in the foreign-body reaction. The type of leukocytes recruited</td>
</tr>
<tr>
<td></td>
<td></td>
<td>is controlled by the type and concentration of the released chemokines in</td>
</tr>
<tr>
<td></td>
<td></td>
<td>the injury place. Granulocytes are recruited from bone marrow within the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>first day and are responsible for the degradation of unviable tissue</td>
</tr>
<tr>
<td></td>
<td></td>
<td>releasing their granule content. For instance, basophils, e.g., release</td>
</tr>
<tr>
<td></td>
<td></td>
<td>heparin, histamine, proteases, and chondroitin. Neutrophils can secrete</td>
</tr>
<tr>
<td></td>
<td></td>
<td>products that stimulate monocytes and macrophages.</td>
</tr>
<tr>
<td>Leukocytes</td>
<td>Hemostasis</td>
<td>Resident cells that contain granules rich in histamine and heparin</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>Tissue resident cells such as resident macrophages and stem cell release</td>
</tr>
<tr>
<td></td>
<td>Repair</td>
<td>healing mediators in the early and long-term response</td>
</tr>
<tr>
<td>Mast cells</td>
<td>Hemostasis</td>
<td>Macrophages arrive at the injury site shortly after neutrophils where they</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>persist for days or longer. Release cytokines, chemokines and GFS.</td>
</tr>
<tr>
<td></td>
<td>Repair</td>
<td>Participate in the phagocytosis of debris and on the development of granulation tissue</td>
</tr>
<tr>
<td>Resident cells</td>
<td>Hemostasis</td>
<td>These cells are recruited by locally released GFs and extracellular matrix</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>degradation products. In the case of skin, these cells are responsible for</td>
</tr>
<tr>
<td></td>
<td>Repair</td>
<td>matrix synthesis, wound strength and contraction and tissue remodeling.</td>
</tr>
<tr>
<td></td>
<td>Remodeling</td>
<td>This cells form capillaries upon GF instruction, which are essential for</td>
</tr>
<tr>
<td></td>
<td></td>
<td>nutrients, gas, metabolite diffusion and for the influx of inflammatory</td>
</tr>
<tr>
<td>Macrophages</td>
<td>Hemostasis</td>
<td>Major players in hemostasis. Granules release cytokines, chemokines,</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>GFS, clotting agents, proteases and inflammation mediators. Facilitate adhesion,</td>
</tr>
<tr>
<td>Fibroblast, pericytes,</td>
<td>Repair</td>
<td>coagulation, vasoconstriction, repair and clot resorption. Their activation attracts and activates several cells</td>
</tr>
<tr>
<td>smooth muscles cells</td>
<td>Remodeling</td>
<td>Granulocytes (neutrophils, basophils, eosinophils) and agranulocytes</td>
</tr>
<tr>
<td>Endothelial cells</td>
<td>Hemostasis</td>
<td>(lymphocytes monocytes and macrophages) are immune system cells</td>
</tr>
<tr>
<td></td>
<td>Inflammation</td>
<td>involved in the foreign-body reaction. The type of leukocytes recruited</td>
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<td>releasing their granule content. For instance, basophils, e.g., release</td>
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<td></td>
<td></td>
<td>heparin, histamine, proteases, and chondroitin. Neutrophils can secrete</td>
</tr>
<tr>
<td></td>
<td></td>
<td>products that stimulate monocytes and macrophages.</td>
</tr>
</tbody>
</table>

Table 3
Tissue expression of extracellular matrix molecules adapted from Sephel and Woodward (2001) and other extracellular characteristics. According to the type, the structural function of the collagen varies: fibrillar collagens (I, II, III, V, XI), network-forming collagen (IV, VIII, X), non-fibrillar collagens (VI, IX, XII, XV, XVIII) and anchoring (VII, XVII).

<table>
<thead>
<tr>
<th>Tissue or fluid</th>
<th>Primary mesoderm cell</th>
<th>Prominent collagen types</th>
<th>Noncollagen proteins</th>
<th>GAGs and PGs</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td>–</td>
<td>–</td>
<td>Albumin, fibrinogen, globulins Fibronecin, elastin, fibrin fibrillin</td>
<td>Hyaluronic acid, sulfated GAGs Red and white blood cells, platelets, plasma</td>
<td>Stratified and vascularized tissue</td>
</tr>
<tr>
<td>Dermis</td>
<td>Fibroblast</td>
<td>I, III, V, VI, XII</td>
<td>Fibronecin, elastin, fibrillin</td>
<td>Hyaluronic acid, decorin, biglycan, fibromodulin</td>
<td>Soft tissue with oriented actin and myosin filaments</td>
</tr>
<tr>
<td>Muscle</td>
<td>Muscle cell/fibroblasts</td>
<td>I, III, V, VI, VIII, XII</td>
<td>Fibronecin, elastin, fibrillin</td>
<td>Aggreican, biglycan, decorin, fibromodulin</td>
<td>Bone–muscle connection. Mostly parallel array of collagen fibers closely packed</td>
</tr>
<tr>
<td>Tendon</td>
<td>Fibroblast</td>
<td>I, III, V, VI, XII</td>
<td>Fibronecin, tenacin fibrillin (myotendon junction), elastin, fibrillin</td>
<td>Decorin, biglycan, fibromodulin, lumican, versican</td>
<td>Bone–muscle connection. Usually parallel array of collagen fibers closely packed</td>
</tr>
<tr>
<td>Ligament</td>
<td>Fibroblast</td>
<td>I, III, V</td>
<td>Fibronecin, elastin fibrillin</td>
<td>Decorin, biglycan, versican</td>
<td>Avascular and transparent stratified tissue</td>
</tr>
<tr>
<td>Cornea</td>
<td>Fibroblast</td>
<td>I, III, V, VI, XII</td>
<td>–</td>
<td>Lumican, keratan, mimecan, biglycan, decorin</td>
<td>Flexible, avascular and ECM rich in PGs and elastin</td>
</tr>
<tr>
<td>Cartilage</td>
<td>Chondrocyte</td>
<td>II, IX, VI, VIII, XI, X (hypertrophic chondrocytes)</td>
<td>Anchornor CII, fibronecin, tenasin</td>
<td>Hyaluronic acid, aggrecan, biglycan, decorin, fibromodulin, lumican, perlecan (minor)</td>
<td>Rigid and vascularized tissue containing 50–70% of calcium phosphate, 20–40% organic matrix, 5–10% of water and &lt;3% of lipids. Long bones contain medullar cavity. Nano-stratified membrane surrounding cells or cellular structures</td>
</tr>
<tr>
<td>Bone</td>
<td>Osteocyte, osteoblasts, osteoclasts</td>
<td>I, V</td>
<td>Osteocalcin, osteopontin, bone sialoprotein, osteonecin</td>
<td>Decorin, fibromodulin, biglycan</td>
<td>Avascular and transparent stratified tissue</td>
</tr>
<tr>
<td>Basement membranes</td>
<td>Epithelial, endothelial, adipocytes, Schwann cell, muscle cells, pericyte, fibroblasts</td>
<td>IV, XV, XVIII</td>
<td>Laminin, nidogen/entactin Heparan sulfate, proteoglycans, perlecan</td>
<td>Hyaluronic acid, sulfated GAGs Red and white blood cells, platelets, plasma</td>
<td>Stratified and vascularized tissue</td>
</tr>
</tbody>
</table>
and are usually linked to a small protein core, forming proteoglycans (Kreuger et al., 2006; Nigam and Bush, 2014; Schultz and Wysocki, 2009). Furthermore, the nature of the major cell and protein types forming the extracellular matrix of the tissues shows certain specificity, highlighted in Table 3.

Both the specific nature of the extracellular matrix and the healing cascade have been inspiring the development of cell instructive tissue engineered constructs (Custodio et al., 2014a). Those relevant cues have been incorporated into biomaterials by: i) the incorporation of human or recombinant GFs, or analogues moieties; ii) the incorporation of ECM-tissue specific compounds, or analogues moieties; iii) the use of co/multi-cultures or cell–cell contact analogues; and iv) the use of immunomodulatory biomaterials. The presentation and incorporation of GFs into biomaterials is commonly controlled by recourse to release systems or surface modification techniques. More translational developments in this area have been hindered by the high cost of recombinant GFs. Thereby, platelet derivatives have attracted immense attention as a human source of multiple GFs and other bioactive molecules demonstrating huge mitogenic, immunomodulatory and differentiation potential (Andia and Maffulli, 2013; Anitua et al., 2004; Cinotti et al., 2013; Kassolis et al., 2000; Kim et al., 2014a; Marx et al., 1998; Oliveira et al., 2015a,b; Semple et al., 2011; Stellos et al., 2010; Vieira-de-Abreu et al., 2012; Weibrich et al., 2002; Weyrich and Zimmerman, 2004).

Engraftment of small peptides mimicking the bioactive epitopes of natural ECM molecules has also shown promising results (Nichol and Khademhosseini, 2009). Frequently, those moieties play important roles during the tissue morphogenesis. For example, the peptide with the sequence of Asp-Gly-Glu-Ala (DGEA), which is derived from collagen type I, has been shown to induce an early-commitment of human mesenchymal stem cells towards the osteogenic lineage (Anderson et al., 2009; Mizuno and Kuboki, 2001). The peptide Glu-Glu-Glu (EEE), which is inspired in the acidic residues of non-collagenous matrix proteins such as osteocalcin and osteopontin, has been shown to promote a more mature osteogenic differentiation than the DGEA (Ceylan et al., 2014).

The combination of different cell types is another approach to obtain a closer representation of the complex cross-talk that occurs in the natural tissue (Battiston et al., 2014). The addition of another cell-type producer of bioactive factors allows a different scheme of interactions, promoting paracrine, autocrine routes and cell–contact dependent effects. Several studies have shown synergistic effects upon the use of co-culture systems and the capability to induce stem cell differentiation (Cooke et al., 2011; J. Wang et al., 2011; Schneider et al., 2011).

Biomaterials for immunomodulation are one of the keys to instruct cell behavior upon implantation and promote the long-term functionality of the device (Forbes and Rosenthal, 2014; Franz et al., 2011; Mokarram and Bellamkonda, 2014). The surgical procedure always initiates an inflammatory response and may elicit an adaptive immune reaction towards the construct surface that should be controlled. Immunomodulatory biomaterials allow the control of the tissue response at the implant site regarding primarily cell adhesion and/or cellular activations. An indirect modulation controls immune cell adhesion and activation, consequently indirectly inducing specific GF secretion. On the other hand, on a direct approach, specific singling molecules are included in the biomaterial (Franz et al., 2011). Controlling hydrophilicity (Jones et al., 2007; Song and Mano, 2013) and fouling properties...
3. Multiscale construct design features

Designing the construct features for in vitro cell fate control and, simultaneously, adequate in vivo performance, is one of the main focuses in tissue engineering (Alves et al., 2010; Lutolf et al., 2009; Marklein and Burdick, 2010). According to the processing approach used for the structure preparation, the best scale working range varies: top-down approaches are more adequate for macro–micro-scale and bottom-up approaches allow higher micro–nano-scale control, while full scale range might be controlled using integrative approaches. Thereby, different approaches offer different control degrees on the final properties of the scaffolds. Nonetheless, multiple variables must be considered, independent of the processing approach. Besides the mechanical properties and the immune reaction, the type and spatial dependence of the processing approach will guide and instruct cell behavior, namely: cell adhesion, cell viability, proliferation, differentiation, matrix production and degradation. According to their major function and scale range, the general scaffold design features can be arbitrarily categorized as: i) geometry; ii) cell-anchorage; iii) release-system; iv) cellularity; v) topographical cues; and vi) biochemical cues — Fig. 3. The scale definition considered is: macro (>10 mm), sub-macro (100 μm–10 mm), micro (1 μm–100 μm), sub-micro or meso (100 nm–1 μm), and nano (<100 nm).

3.1. Geometry — macro, sub-macro

The geometry of the scaffold has, generally, to be tailored according to the implantation site request, in order to fill the tissue’s void space and support the site’s mechanical demands. Additive (Hutmacher et al., 2004; Melchels et al., 2012; Tsang et al., 2007), ablation (Meng et al., 2009) and molding (C.H. Chen et al., 2014; Correlo et al., 2009; Zhang et al., 2014) technologies, based on previous three dimensional patient data acquisition (e.g., by X-ray microtomography or magnetic resonance imaging), allow the preparation of geometry-specific 3D structures. On the other hand, some applications do not demand specific geometries or high mechanical strength. In such cases, combinations of materials, cells and bioactive factors are usually injected filling the void spaces (Custodio et al., 2015; Custódio et al., 2014; Koshy et al., 2014; Kretlow et al., 2007; Larsson and Hannink, 2011). Shaping the frequently affects the bulk mechanical properties of the construct. For instance, scaffolds prepared by rapid prototyping show different compression moduli according to the alignment (Hutmacher et al., 2001) or the distance between struts (Sobral et al., 2011). Nonetheless, computer assisted design technologies offer the possibility to simulate and optimize the inner geometry (strut alignment, pore size, pore geometry and interconnectivity) in order to enhance the mechanical performance, by using finite element analysis and CAD design tools (Lacroix et al., 2006; Lin et al., 2004; Sandino et al., 2006). Those micro-features, pore geometry (Bidan et al., 2013; Rumpler et al., 2008), pore size (Oh et al., 2007; Woodfield et al., 2005; Zeltinger et al., 2001) and porosity (Karageorgiou and Kaplan, 2005), also affect oxygen perfusion, the removal of cell metabolites, cell seeding efficiency and tissue growth rate, making them very important features regarding cell viability and growth. Additionally, those micro-properties also influence the invasion of inflammatory and immune cells and the ingrowth of blood vessels or other cells from the host tissue (Karageorgiou and Kaplan, 2005; Karp et al., 2004; Kasten et al., 2008).

3.2. Cell-anchorage — sub-macro, micro, sub-micro, nano

The cell-anchorage defines the physical elements supporting the adhesion of the cells at scale smaller than the geometry of the scaffold. Depending on the technique used to shape the materials or, e.g., depending on the subsequent/simultaneous incorporation of gels (Correa et al., 2012; Prabaharan et al., 2007b), sponges (Chen et al., 2003; Mano et al., 2008), micro/nano-fibers or fibrillar structures (S.M. Oliveira et al., 2013b), the anchorage points and the micro-properties of the scaffold can be tailored. Introducing anchorage points within the pores is also a way to increase cell surface growth area/volume and alter the spatial distribution of the tissue growth. Moreover, their physical properties (e.g. stiffness/elasticity and adhesion area) will affect cell morphology by altering the cytoskeletal organization and contractibility, which can guide stem cell differentiation into specific lineages (Engler et al., 2006; Guilak et al., 2009).

3.3. Release systems — micro, sub-micro, nano

The incorporation of release systems in the substrates is considered an essential tool for the temporal or spatial–temporal controlled delivery of instructive or pharmaceutical compounds (Lima et al., 2012; Madduri and Gander, 2012; Oliveira and Mano, 2011; Santo et al., 2012c, 2013). Some examples of release systems of interest in tissue engineering relate to the delivery of one or multiple compounds such as GFs (Johnson and Wang, 2013; Kim et al., 2013; Richardson et al., 2001; Tabata, 2003), dexamethasone (Duarte et al., 2009a; Wang et al., 2010, 2014), gentamicin (Gao et al., 2011; Li and Chang, 2005), ketoprofen (Prabaharan and Mano, 2005; Prabaharan et al., 2007a) and platelet derivatives (Coppening et al., 2004; Kawase et al., 2003; Santo et al., 2012a; Y.K. Kim et al., 2014). Typically, a release-system can be prepared by the sole mixing of the compounds with the materials prior to the 3D processing (Perets et al., 2003; Santo et al., 2012a), or by its incorporation afterwards, using surface modification techniques (e.g., by layer-by-layer assembling (Gilde et al., 2012) or adsorption (Santo et al., 2012b)) or impregnation (e.g. supercritical fluid CO2 (Cabezas et al., 2014; Biondi et al., 2008)). In combination with stimuli responsive materials (Mano, 2008), these systems can be triggered in a controlled manner, for instance by temperature (Costa et al., 2009; Shi et al., 2008), pH (Costa et al., 2009; Santos et al., 2010) and light stimuli (Borges et al., 2014; Mano, 2008; Stuart et al., 2010). Besides being interesting for the development of smart release systems, stimuli responsive materials have been shown to provide variations in other important properties, such as wettability, affecting cell adhesion (da Silva et al., 2007) or mineralization (Cole et al., 2009; Shi et al., 2007).

3.4. Cellularity — micro

For cellular tissue engineered approaches, the type of cells seeded on the 3D structure has major repercussions on new tissue formation. According to the source tissue or donor, stem cells possess different differentiation potentials (Bianco et al., 2001; Chamberlain et al., 2007; Kern et al., 2006; Odorico et al., 2001). Moreover, the seeding of more than one cell type (i.e. co/multiple cell culture) and the ratio between those cells allows different crosstalk to occur. The presence of other cells will affect the nano/micro-environment, e.g., by creating a new profile of cell-secreted cytokines and GFs (Cooke et al., 2011; Schneider et al.,...
3.5. Topographical cues — micro, sub-micro, nano

The nano/micro-topographical cues are one of the primary mediators of the adsorption of proteins, from the surrounding media (e.g., serum media or plasma), that further mediate the cell adhesion process (Lord et al., 2010; Raffaini and Ganazzoli, 2013; Roach et al., 2006; S.M. Oliveira et al., 2014; Song and Mano, 2013). Controlling these features can synergistically improve cell behavior. As it has been shown, designing specific surface grooves and patterns can induce specific cell shapes (Chou et al., 1995; McBeath et al., 2004; Singhvi et al., 1994) or alignments (Hosseini et al., 2014; Luckier et al., 2014; McCloskey, 2013) that alter cytoskeleton contractibility guiding cell differentiation. Well-controlled topographical cues, patterns and surface biochemistries can be easily prepared in 2D. However, the transposition of those features to 3D is still a major challenge. Surface properties design and surface engineering techniques have been extensively reviewed elsewhere (Alves et al., 2010; Braffman, 2013; Costa and Mano, 2014; Discher et al., 2009; Dvir et al., 2011; Han et al., 2014; Lutolf and Hubbell, 2005; Lutolf et al., 2009; Marklein and Burdick, 2010).

3.6. Biochemical cues — nano

At the nanoscale, the surface biochemistry instructs cell behavior either by directly mediating the adsorption of surrounding proteins to the construct, or by binding/activating cell surface receptors — Table 1. The biochemical cues, such as GfS (Gitay-Goren et al., 1992; King and Krebsbach, 2011; Saik et al., 2011), extracellular matrix proteins (Custodio et al., 2010; Sayyar et al., 2014; van den Dolder et al., 2003; Zhang et al., 2005) (e.g., collagen and fibronection), functional groups (Amorim et al., 2013; Arima and Iwata, 2007; da Costa et al., 2012) (e.g. sulfation (S.M. Oliveira et al., 2013a)), small peptides (Hersel et al., 2003; Massia and Hubbell, 1990, 1991) (e.g., –RGD), and cell contacts (Chen et al., 2004; Lambert et al., 2000) (e.g., cell–cell interactions or functional groups mimicking cells contact) will trigger specific signaling pathways in the cell and define their fate. A controlled design of the biochemistry is commonly made recurring to nanotechnologies and surface modification methods, such as: layer-by-layer assembling (Jan and Kotov, 2007; Macdonald et al., 2011; Martins et al., 2010; Neto et al., 2014; S.M. Oliveira et al., 2013a) and antibody mediated binding (Custodio et al., 2014b; Custódio et al., 2014).

Despite most features might be roughly allocated to a defined range of length scale, their properties can be intimately related with other features (S.M. Oliveira et al., 2014). A classical example is the case of roughness—wettability (Neto et al., 2011; S.M. Oliveira et al., 2011, 2014). Introducing nano/micro-roughness in a substrate in the first instance may represent a higher surface area available for cell growth. Nevertheless, the altered roughness might also change the surface wettability into values that are not as favorable for cell adhesion/growth (Alves et al., 2009; Song et al., 2009). Whether the ability of a surface to be wet is altered, its profile of protein adsorption will also change in terms of conformation and density of proteins. Consequently, the communication between cell and the support can be significantly altered at the cell receptor level (Gittens et al., 2014).

4. Differentiation biases and aging

Regenerative success is intricately dependent on all the interactions that cells are subjected to upon the in vitro culture, the interaction with the construct and the in vivo implantation. Besides the crucial roles of the biomaterials’ properties considered in most of the research conducted in tissue engineering explored in the last section, there are others affecting mostly the cellular phenotype and genotype quality. Those features concern issues such as cell senescence, aging and differentiation biases.

Most of the commonly used in vitro expansion platforms are not adequate and damage cell telomeres, causing cell senescence, aging, reduction of differentiation potential and differentiation biases (Haik et al., 2000; Oh et al., 2014; Salehinejad et al., 2013). The conditions used for stem cell expansion can alter the lineage fate. Moreover, it has been reported that stem cells have mechanical memory of previous culture conditions (Yang et al., 2014). Also, it has been shown that the chemistry of the surface used to expand stem cells does affect their in vivo regeneration capability (Huang et al., 2014), as well as the tendency for a certain lineage differentiation (Hsu and Huang, 2013). For instance, the biases lineage-commitment tendency of mesenchymal stem cells to undergo aging, telomere shortening, and spontaneous low quality osteogenic differentiation upon regular culture expansion, as well as decreased multilineage differentiation quality has been reported (Li et al., 2011; Yang et al., 2014). To overcome those issues, geometrical, topographical and physical cues can be used to alter cytoskeleton contractibility, guiding stem cell lineage specification, and reducing the biases as well as telomerase activator can be incorporated in order to delay telomeres shortening (Engler et al., 2006; Guilak et al., 2009; Kilian et al., 2010; Reilly and Engler, 2010). There is still an urgent need to develop and spread anti-aging, anti-biases procedures, and new 2D/3D expansion substrates to improve specificity, cellular performance and tissue regeneration.

5. Approaches for 3D construct fabrication

3D constructs can be prepared using top-down, bottom-up or integrative methodologies. Those approaches allow varied length scale degrees of control of the building and final properties. Using top-down approaches, well-defined scaffolds, where multiscale control is not a demand, can be prepared, whereas bottom-up stands for a higher control at the nano/micro scale. On the other hand, integrative approaches emerge as a combination of both top-down and bottom-up approaches, promising scaffolds with properties and features controlled over the entire scale range. Bottom-up and integrative approaches present higher potential for the design of supportive biomaterials with multiscale features and high biochemistry control than top-down, being more focused in this review.

5.1. Top-down approach

Top-down approaches provide considerable higher control over the macro/micro scale properties/features rather than the nano/sub-micro ones. A bulk material is scaled down into small compartments or nano-scale details are incorporated, indirectly controlling the processing parameters of the fabrication technique (Yan et al., 2011). Top-down scaffolds have been produced recurring to several processing techniques, such as: freeze-drying (Correia et al., 2011; Ho et al., 2004; Wu et al., 2010), solvent-casting (Mota et al., 2012; S.S. Silva et al., 2013; Sin et al., 2010; Thadavirul et al., 2014), electrospinning (Park et al., 2008; Yoshimoto et al., 2003; Zong et al., 2005), rapid prototyping (Landers et al., 2002; Silva et al., 2009; Yang et al., 2002) and supercritical fluids (Duarte et al., 2009b, 2012). They present pores and mechanical properties that can be controlled to some extent, for instance, by controlling the inner architecture and the nature of the materials. In particular, rapid prototyping allows a precise and computer assisted design of the inner geometry and, simultaneous, control of the external shape. All different kinds of materials and blends, including natural and synthetic polymers and proteins, ceramics and metals, have been used to prepare 3D constructs. With exception of extracellular derived compounds or moieties that not always are obtained from cost-
effective sources, those constructs frequently lack cell instructive cues (Silva et al., 2010). To improve cell behavior on such structures, usually the scaffolds are prepared combining different top-down techniques (W. Wang et al., 2011; Yeo and Kim, 2014) or are subjected to further functionalization requiring integrative approaches (Macdonald et al., 2011; S.M. Oliveira et al., 2013b; Yeo and Kim, 2011). For instance, electrospinning and 3D printing can be combined to prepare hierarchical scaffolds with dual scale fibers: the printed micro-struts combined with electrospun nano/microfibers (Park et al., 2008; Santos et al., 2008; Tuzlakoglu et al., 2005). The electrospun fibers increase cell-anchorage points and seeding efficiency (Park et al., 2008) and can promote cell alignment (Park et al., 2014).

5.2. Bottom-up approach

Bottom-up methodologies are those applying chemical or physical forces leading to the assembling of micro/nanoscale-defined building blocks into larger scale structures. The chance to control the construct over several length-scales makes this approach more attractive for matrices’ engineering than the top-down approach, namely in finding new opportunities to mimic a tissue microstructure and the spatial organization of multiple cell types (Nichol and Khademhosseini, 2009). The most commonly used building blocks comprise cells, cell aggregates/spheroids, nano/microparticles, tubes, layers, fibers and gels. The development of methodologies to produce, functionalize, and assemble those small units is the major challenge — Fig. 4. Currently, there are three general alternatives to assemble the units: i) randomly; ii) mediated by an external force; and iii) specifically assembled — see Fig. 4.

5.2.1. Random assembling

Layer-by-layer assembling (LbL) is a simple and random bottom-up assembling method based on the alternating deposition of polyelectrolytes with different charges (Borges and Mano, 2014; S.M. Oliveira et al., 2013a; Tang et al., 2006). Using LbL, multilayered structures can be created around the units, binding and stabilizing the entire set and forming a scaffold (Miranda et al., 2011; Sher et al., 2015a,b), temporary templates to produce high porous foams (J.M. Silva et al., 2013; Sher et al., 2010) or hierarchical capsules (Correia et al., 2013; Costa et al., 2013). Such extension of the LbL methodology towards the fabrication of 3D structures is an example of the application of this technology in the biomedical field (Costa and Mano, 2014).

It has also been shown that the building blocks can be unspecifically assembled by cells which colonize their surfaces, consolidating the structure (García Cruz et al., 2008; M. B. Oliveira et al., 2011; McGuigan and Sefton, 2006). The presentation of cell-recognizable moieties on the surface can accelerate this cell-based assembling. For instance, chitosan microparticles functionalized with a mitogenic GF (PDGF) have shown a stable and faster assembling, within 12 h, than in the absence of PDGF (Custódio et al., 2014).

Scaffold free technologies represent another derivation of the bottom-up approach. Cells grown on thermoresponsive culture dishes are retrieved as a confluent monolayer, containing the extracellular matrix proteins, by decreasing the local temperature (da Silva et al., 2007; Yang et al., 2005, 2007). These cell sheets can be consecutively stacked on top of each other to produce thicker and more complex tissues. A reported alternative is based on consecutive cell seeding on top of confluent cell layers previously coated with proteins (Chetprayoon et al., 2013; Matsusaki, 2012). The preparation of thick cell-constructs and their low load-bearing capability are the major challenges in using such technology in tissue engineering.

5.2.2. Mediated assembling

The assembling of small units can be guided by external stimuli, allowing the control over their spatial organization to a certain extent.
For example, the successful attachment of gel surfaces has been demonstrated using silica particles (Abe et al., 2014a,b; Tamagawa and Takahashi, 2008). Through electrostatic interactions at the particle-gel interface and cohesion forces in the particle layer, two gel-surfaces can be attached. The adhesion force, mediated by the particles, can reach 20 kPa as opposed to 1.5 kPa when using an oppositely charged polyelectrolyte solution (Abe et al., 2014a,b; Tamagawa and Takahashi, 2008). Hydrogel units composed of materials presenting opposite charges can also be assembled by electrostatic attractions (Katayama et al., 2013). However, to promote the electrostatic binding, the distance among the units has to be decreased, e.g. by direct positioning (Abe et al., 2014b), by shaking (Katayama et al., 2013) or using acoustic waves (Xu et al., 2011a). Alternatively, it has been shown that template gels can be successively dipped onto oppositely charged micro-gel colloids to form multilayered-like constructs (Han et al., 2013). Moreover, the use of shape complementary building blocks allows the formation of lock-and-key shaped assemblies mediated by electrostatic interaction or surface tension (Du et al., 2008; Katayama et al., 2013). In alternative, magnetic nanoparticles can be incorporated into cells (Castro and Mano, 2013; Souza et al., 2010) and biomaterial units (Gil and Mano, 2014; Liu et al., 2014; Xu et al., 2011b) and further 3D organized recurring to an external magnetic field. However, until now a perfect control of the position in the space of the different unit has been limited.

5.2.3. Specific assembling

While the alternatives for the unspecific 3D assembling of building block are many, the current available methods to design 3D constructs with specific spatial organizations are yet limited and more complex.

Microfluidic technologies have been proposing promising tools for the assembly, for example, of small gel units mostly by mediated assembling. By controlling the inlets, outlets and flow, some microfluidic devices can actually guide the assembling of small units into 1D, 2D and 3D structures (Chung et al., 2011). Moreover, it has been shown that using the so-called multilayer-microfluidic devices, mosaic hydrogels with a well-defined spatial composition of materials, cells or pores can be prepared (Leng et al., 2012).

Based on the oligonucleotide pairs complementarity principle, gel interfaces can be specifically assembled (Deschner et al., 2014; Qi et al., 2013). Setting the spatial/surface presentation of the complementary sequences in the different units allows the formation of controlled assemblies. Although being assemble-specific, this approach might be scale-up limitative. Other alternative methods, for both materials and cell controlled assembling, are based on bioprinting technologies. Bioprinting is a particular type of rapid prototyping technique that deposits small units of multiple materials and cells, instead of continuous filaments (Schuurman et al., 2011). Highly complex organization with cellular vascular networks, multiple cell types and extracellular matrices were already shown to be achievable (Kolesky et al., 2014).

Even though bottom-up approaches seem ideal to mimic a tissue organization, major developments and new concepts to improve the assembling and spatial organization of the assembling units still necessary. Limitations such as the weak control on the final 3D shape, the lack of scalability and the lack of understanding on the short and long-term interactions among the units and cell behavior are yet to be further investigated.

5.3. Integrative approach

The combination of the top-down and bottom-up approaches constitutes a valuable strategy to increase the functionality of the devices for tissue engineering and achieve a multiscale control over the 3D substrate properties.

According to the integration between the bottom-up and top-down approaches, three sub-categories of integrative methods can be defined: i) sequential integration; ii) combination; and iii) technical integration – Fig. 5.

Employing an integrative approach could allow the increment of one or more feature/property in the scaffold without impairing the previous structure properties (Hollister and Murphy, 2011). Thereby, fully customizable constructs with multiple scales of complexity could be designed. With such ideal control and the increment of measurable properties, in silico modulation could contribute greatly for the understanding of the relationship between multiscale properties and cell behavior. Those resulting models would allow the mathematical prediction of cell behavior, contributing to the device optimization and eventually to the reduction of in vitro/in vivo experimentation costs (Walpole et al., 2013).

5.3.1. Sequential integration

3D constructs may be prepared using different techniques in a stepwise manner; i.e., by sequential integration. In these cases, the 3D structures are step-by-step prepared using at least one top-down and one bottom-up method. This category is the most used to overcome the lack of bioactivity and cell instruction of the traditional 3D scaffolds prepared by top-down approaches. Modification techniques such as adsorption, layer-by-layer assembling, self-assembling and grafting can be used to modify 3D scaffolds without impairing their initial features. Those methods have been used to include instructive cues such as adhesive proteins (Assmann et al., 2013; Choi et al., 2013; Ragetly et al., 2010; Truong et al., 2012), GFs (Crouzier et al., 2011; Macdonald et al., 2011; Shah et al., 2014; Tang et al., 2013), modular peptides (Lee et al., 2009) and bioactive calcium phosphate particles (Zhou et al., 2014). Controlling the biochemical environment and the incorporation
of release system is manageable using sequential approaches; however, it is yet challenging to control the topographical cues on 3D top-down structures. The use of microfabrication techniques for the preparation of micro-layers of the substrate is an appealing solution. Several features might be controlled in each layer (biochemistry, topography, release system and cells), which are further assembled or stacked (Kolewe et al., 2013; Lima et al., 2014b). However, assembling and stabilizing those layers are still challenging as in the case of the bottom-up approaches.

5.3.2. Combination
Contrary to the other categories, in the integration by combination, the building blocks are incorporated prior to the processing and the top-down method is performed as if in absence of such units. Thereby, this approach allows the incorporation of new features, such as release systems in the bulk structure of the constructs. For instance, supercritical CO2 fluid foaming was used to prepare scaffolds of poly-ε-lactic acid with nanoparticles carrying platelet lysate (Santo et al., 2012a). The incorporation of those bioactive nanoparticles enhanced the osteogenic differentiation of human adipose derived stem cells on the foams. Polymeric microcarriers, which allow a controlled expansion and formation of cellular aggregates, were successfully combined with gelatin methacrylame–gellan gum bio-inks (Levato et al., 2014). With this combination, 3D scaffolds could be achieved with simultaneous high cell density, viability and improved mechanical performance.

5.3.3. Technical integration
The technical integration refers to the development of 3D constructs recurring to specialized equipment for simultaneous top-down and bottom-up processing. With such type of equipment’s, topography, release system, biochemistry, shape and cells could be ideally controlled. A foreseen example employing this approach could be the combination of rapid prototyping and spraying of small-units containing instructive cues to include in the 3D structures. Recurring to a software-assisted deposition of both, shape controlled scaffolds with spatially defined topography, release systems and biochemistry could be fabricated. Though very appealing, such sophisticated technologies have yet to be developed.

6. Conclusions and perspectives
The major nano/micro-environment properties and events occurring in the normal and regenerative tissue have been inspiring the selection or synthesis of instructive cues to include in the 3D structures. However, there is still a need to understand which are the most effective spatial—temporal cues for each tissue. Besides the biochemistry, other scaffold features, e.g. topography, cells, cell anchorage and geometry, play major roles for the construct success both in vitro and in vivo. High-content screening of 3D scaffold properties and features promises a huge contribute in deciphering which cues, materials and methods will be more effective for the regeneration of each tissue (Hook et al., 2010; M.B. Oliveira et al., 2013, 2014; Simon and Lin-Gibson, 2011).

The control of all such features calls for processing approaches that allow a multiple scale control, i.e. integrative approaches. According to the combinations between the top-down and bottom-up techniques, different integrative approaches were defined: sequential, combination and technical integration. The assembling of building blocks in a single bottom-up approach is yet in its infancy, in regard to tissue engineering applications or the development of 3D tissue-models. However, their integration with top-down approaches, e.g. 3D printing, may accelerate their development and the preparation of a highly controlled multiscale 3D tissue engineering devices. The increase in complexity of the fabricated devices should also be balanced with the regulatory issues and commercial viability so that their realistic employment in the clinics could be envisaged.

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